National Institute for Health and Care Excellence

Final version

Low back pain and sciatica in over 16s: assessment and management

Low back pain and sciatica in over 16s

NICE guideline NG59

Appendices I-J

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Appendices

Appendix I: Economic evidence tables

I.1 Clinical Examination

None.

I.2 Risk assessment tools/stratification

Table 1: Apeldoorn 2012^{2,3}

Apeldoorn AT, Bosmans JE, Ostelo RW, de Vet HC, van Tulder MW. Cost-effectiveness of a classification-based system for sub-acute and chronic low back pain. European Spine Journal. 2012; 21(7):1290-1300. (Guideline Ref ID APELDOORN2012)

Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness
Economic analysis: CUA (health outcome: QALYs) Study design: Within-trial (RCT, associated clinical paper Apeldoorn2012A) Approach to analysis: EQ-5D data collected at baseline and 1 year follow-up to calculate QALYs. Within-trial reported resource use, including primary and secondary care utilisation, unit costs applied.	Population: Adults with low back pain (with or without sciatica) Cohort settings: Start age: 42.6 years Male: 42.9% Intervention 1: (n=82) Usual physical therapy care based on Dutch physical therapy low back pain guidelines. Intervention 2: (n=74)	Total costs (mean per patient): Intervention 1: £574 Intervention 2: £505 Incremental (2–1): saves £69 (95% CI: -£312 to £226; p=NR) Currency & cost year: 2009 Dutch Euros (presented here as 2009 UK pounds (a)) Cost components incorporated:	QALYs (mean per patient): Intervention 1: 0.80 Intervention 2: 0.82 Incremental (2–1): 0.02 (95% CI: -0.03 to 0.08; p=NR)	ICER (Intervention 2 versus Intervention 1): Intervention 2 dominates intervention 1 (lower costs and higher QALYs) (da) 95% CI: NR Probability Intervention 2 cost-effective (£20K/30K threshold): NR Analysis of uncertainty: Bootstrapping of ICER conducted but only from a societal perspective not a health care provider perspective. Therefore this is not reported here. Bootstrapping of costs conducted and confidence intervals are presented here. Additional sensitivity analyses were conducted (including using a per-protocol analysis and complete cases only) however

Perspective: Dutch healthcare payer perspective Follow-up: 1 year Discounting: Costs: n/a; Outcomes: n/a	Hicks/Delitto classification based interventions: spinal manipulation, stabilisation exercises or direction specific exercises for a minimum of 4 weeks.	Primary care utilisation including: GP contacts, physical and manual therapy, psychologist and professional home care. Secondary care utilisation including: X-ray, MRI scan, outpatient specialist visit, hospitalisation, herniated nucleus pulposus surgery, outpatient rehabilitation, epidural injection and facet denervation.		these were all from a societal perspective and so are not reported here.
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Data sources

Health outcomes: Within-trial analysis (RCT, Apeldoorn 2012A)^{3,4}. Health outcomes included patient reported EQ-5D collected baseline and 1 year follow-up. **Quality-of-life weights:** Dutch EQ-5D tariff. **Cost sources:** Patient-reported resource use based on cost diaries completed at 8, 26, 39 and 52 weeks. Unit costs based on Dutch guidelines for costs studies and Dutch national medication costs.

Comments

Source of funding: Netherlands Organisation for Health Research and Development. **Limitations:** Dutch resource use data (2008-2010) and unit costs (2009) may not reflect current NHS context. Dutch EQ-5D tariff used. Not all risk stratification tools from the review protocol are included in this study. Within-trial analysis and so may not reflect full body of evidence for this comparison; Apeldoorn 2012A is 1 of 2 studies in the clinical review for risk stratification comparing Hicks/Delitto.

Bootstrapping of ICER from NHS and PSS perspective not undertaken. **Other:** none.

Overall applicability(b): Partially applicable Overall quality(c): Potentially serious limitations

Abbreviations: 95% CI: 95% confidence interval; CUA: cost—utility analysis; da: deterministic analysis; EQ-5D: Euroqol 5 dimensions (scale: 0.0 [death] to 1.0 [full health], negative values mean worse than death); ICER: incremental cost-effectiveness ratio; NR: not reported; QALYs: quality-adjusted life years

- (a) Converted using 2009 purchasing power parities⁴²
- (b) Directly applicable / Partially applicable / Not applicable
- (c) Minor limitations / Potentially serious limitations / Very serious limitations

Table 2: Whitehurst 2012^{66,67}/Hill 2011^{20,21}

Whitehurst DGT, Bryan S, Lewis M, Hill J, Hay EM. Exploring the cost-utility of stratified primary care management for low back pain compared with current best practice within risk-defined subgroups. Annals of Rheumatic Diseases. 2012; 71(11):1796-1802. (Guideline Ref ID WHITEHURST2012) Hill JC, Whitehurst DG, Lewis M, Bryan S, Dunn KM, Foster NE et al. Comparison of stratified primary care management for low back pain with current best practice (STarT Back): a randomised controlled trial. Lancet. 2011; 378(9802):1560-1571. (Guideline Ref ID HILL2011)

Study details	Population & interventions	Costs (a)	Health outcomes	Cost effectiveness
Economic analysis: CUA (health outcome: QALYs) Study design: Within-trial (RCT, associated clinical paper Hill 2011) Approach to analysis: EQ-5D data collected at baseline, 4 and 12 months follow-up. QALYs constructed through area under the curve method. Number of study-related physiotherapy sessions reported via case report forms and audit of clinical notes. All other healthcare resource use collected at 12-months follow-up via self-report questionnaires. Unit costs applied. Perspective: UK NHS Follow-up: 1 year Discounting: Costs: n/a; Outcomes: n/a	Population: Adults with low back pain (with or without sciatica) Cohort settings: Start age: 49.8 years Male: 41.2% Intervention 1: (n=283) Current best practice: STarT Back stratification followed by physiotherapist assessment lasting 30 minutes which included initial treatment advice and exercise with the option for onward referral for further physiotherapy, based on physiotherapist clinical judgement. Intervention 2: (n=568) STarT Back stratification followed by one of three treatment pathways based on risk. Physiotherapist assessment lasting 30	Total costs (mean per patient) Intervention 1: £243.52 Intervention 2: £212.88 Incremental (2–1): saves £30.64 (95% CI: NR; p=NR) Intervention costs (mean per patient): Intervention 1: £92.77 Intervention 2: £107.50 Incremental (2–1): £14.73 (95% CI: NR; p=NR) Currency & cost year: 2008/2009 UK pounds Cost components incorporated: Intervention cost; primary care utilisation including: GP and nurse contacts; secondary care utilisation including: consultant contacts, X-ray, MRI scan, CT scan, blood tests epidural	QALYs (mean per patient): Intervention 1: NR Intervention 2: NR Incremental (2–1): 0.039 (95% CI: 0.01 to 0.07; p=0.01)	Overall ICER (Intervention 2 versus Intervention 1): Intervention 2 dominates intervention 1 (lower costs and higher QALYs) (da) 95% CI: NR Probability Intervention 2 cost-effective (£20K threshold): NR Analysis of uncertainty: Bootstrapping of ICER undertaken however this included private healthcare costs as well as NHS costs. Therefore this is not reported here. Sensitivity analyses were conducted using the complete case analysis rather than the primary imputed analysis. Intervention 2 remained dominant (lower costs and higher QALYs).

minutes, including initial treatment with advice on promoting appropriate levels of activity, return to work and a pamphlet about local exercise venues and self-help groups. All were shown a 15-minute educational video and given the Back Book.

injections; other healthcare professional contacts including additional physiotherapy and prescribed medication.

Low risk group only received above initial session.

Medium risk group referred for standardised physiotherapy sessions to address symptoms and function.

High risk group referred for psychologically-informed physiotherapy sessions to address symptoms and function and also psychosocial obstacles to recovery.

Data sources

Health outcomes: Within-trial analysis (RCT, Hill 2011) ^{20,21}. Health outcomes included patient reported EQ-5D collected baseline and 12 months follow-up. QALYs were calculated using the area under the curve approach adjusted for baseline utility. **Quality-of-life weights:** EQ-5D UK tariff. **Cost sources:** Number of study-related physiotherapy sessions reported via case report forms and audit of clinical notes. All other healthcare resource use collected at 12-months follow-up via self-report questionnaires. Unit costs form UK published sources including PSSRU, BNF and NHS reference costs.

Comments

Source of funding: Arthritis Research UK. **Limitations:** Not all risk stratification tools from the review protocol are included in this study. Within-trial analysis: Hill 2011 is 1 of 2 studies included in the clinical review for risk stratification comparing STarT Back. Bootstrapping of ICER from NHS and PSS perspective not undertaken. **Other:** None

Overall applicability(b): Directly applicable Overall quality(c): Potentially serious limitations

Abbreviations: 95% CI: 95% confidence interval; CUA: cost—utility analysis; EQ-5D: Euroqol 5 dimensions (scale: 0.0 [death] to 1.0 [full health], negative values mean worse than death); ICER: incremental cost-effectiveness ratio; NR: not reported; pa: probabilistic analysis; QALYs: quality-adjusted life years

- (a) Hill 2011 presented total healthcare costs that included both NHS and private healthcare resource use, these were recalculated and costs presented here are for NHS only healthcare resource use only.
- (b) Directly applicable / Partially applicable / Not applicable
- (c) Minor limitations / Potentially serious limitations / Very serious limitations

Table 3: Whitehurst 2015 11,12,65,67

Whitehurst DG, Bryan S, Lewis M, Hay EM, Mullis R, Foster NE. Implementing Stratified Primary care Management for low Back Pain: Cost Utility Analysis alongside a Prospective, Population-based, Sequential Comparison Study. Spine. 2015; Epublication. (Guideline Ref ID WHITEHURST2015)

Foster NE, Mullis R, Hill JC, Lewis M, Whitehurst DGT, Doyle C et al. Effect of stratified care for low back pain in family practice (IMPaCT Back): a prospective population-based sequential comparison. Annals of Family Medicine. 2014; 12(2):102-111 (Guideline Ref ID FOSTER2014)

Study details	Population & interventions	Costs (a)	Health outcomes	Cost effectiveness
Economic analysis: CUA (health outcome: QALYs) Study design: Within-trial (cohort study, associated clinical paper Foster 2014) Approach to analysis: EQ-5D data collected at baseline, 2 and 6 months follow-up. QALYs constructed through area under the curve method. Healthcare resource use collected at 6-months follow-up via self-report questionnaires. Unit costs applied. Perspective: UK NHS Follow-up: 6 months	Population: Adults with low back pain (with or without sciatica) Cohort settings: Start age: 48.7 years Male: 44.7% Intervention 1: (n=630) Usual care: Family physician management involving assessment, advice, medication, sickness certification and referral for investigations or further treatment as appropriate, based on clinical judgement. Community based physical therapists managed patients using clinical judgement to	Total costs (mean per patient) Intervention 1: £169.43 Intervention 2: £164.54 Incremental (2–1): saves £4.89 (95% CI: NR; p=NR) Currency & cost year: 2008/2009 UK pounds Cost components incorporated: Primary care utilisation including: GP and nurse contacts; physiotherapy service; secondary care utilisation including: consultant contacts, admissions, radiograph, MRI	QALYs (mean per patient): Intervention 1: NR Intervention 2: NR Incremental (2–1): 0.003 (95% CI: -0.01 to 0.02; p=NR)	Overall ICER (Intervention 2 versus Intervention 1): Intervention 2 dominates intervention 1 (lower costs and higher QALYs) (da) 95% CI: NR Probability Intervention 2 cost-effective (£20K threshold): NR Analysis of uncertainty: Bootstrapping of ICER undertaken however this included private healthcare costs as well as NHS costs and was done by risk group only. Therefore this is not reported here. Sensitivity analyses were conducted using the complete case analysis rather than the primary imputed analysis. Intervention 2 remained dominant (lower costs and higher QALYs).

Discounting: Costs: n/a;

Outcomes: n/a

determine content and number of treatment sessions.

Intervention 2: (n=1,017) STarT Back stratification followed by one of three treatment pathways based on risk.

Low risk group: family physician provided written information on selfmanagement and advice to keep active, prescription of reassurance regarding good included a minimal package scan, CT scan, blood tests epidural injections; other healthcare professional contacts including acupuncture and osteopathy; and prescribed medication.

Medium risk group: Family therapy and address their highlighted by stratification reducing pain and disability using activity, exercise and

early return to work.		
High risk group: Family		
physician encouraged to		
refer patients to physical		
therapy and address their		
back-related concerns		
highlighted by stratification		
tool. Psychologically		
informed physical therapy		
provided.		

Data sources

Health outcomes: Within-trial analysis (cohort study, Foster 2014)¹². Health outcomes included patient reported EQ-5D collected baseline, 2 and 6 months follow-up. QALYs were calculated using the area under the curve approach adjusted for baseline utility. **Quality-of-life weights:** EQ-5D UK tariff. **Cost sources:** Healthcare resource use collected at 6-months follow-up via self-report questionnaires. Unit cost sources not reported.

Comments

Source of funding: The Health Foundation. **Limitations:** Not all risk stratification tools from the protocol are included in study. A longer time horizon may be preferable if effects may persist beyond 6 months. Source of unit costs not reported. Within-trial analysis and so does not reflect full body of available evidence for this comparison; Foster 2014 is 1 of 2 studies included in risk stratification review comparing STarTBack to usual care. Appropriate bootstrapping of ICER not undertaken. **Other:** None

Overall applicability(b): Directly applicable Overall quality(c): Potentially serious limitations

Abbreviations: 95% CI: 95% confidence interval; CUA: cost—utility analysis; EQ-5D: Euroqol 5 dimensions (scale: 0.0 [death] to 1.0 [full health], negative values mean worse than death); ICER: incremental cost-effectiveness ratio; NR: not reported; pa: probabilistic analysis; QALYs: quality-adjusted life years

- (a) Foster 2014 presented total healthcare costs that included both NHS and private healthcare resource use, these were recalculated and costs presented here are for NHS only healthcare resource use only.
- (b) Directly applicable / Partially applicable / Not applicable
- (c) Minor limitations / Potentially serious limitations / Very serious limitations

I.3 Imaging

Table 4: Gilbert 2004^{14,15}

Gilbert FJ, Grant AM, Gillan MG, Vale LD, Campbell MK, Scott NW et al. Low back pain: influence of early MR imaging or CT on treatment and outcome. Multicenter randomized trial. Radiology. 2004; 231(2):343-351. (Guideline Ref ID GILBERT2004)

Gilbert FJ, Grant AM, Gillan MGC, Vale L, Scott NW, Campbell MK. Does early magnetic resonance imaging influence management or improve outcome in patients referred to secondary care with low back pain? A pragmatic randomised controlled trial. Health Technology Assessment. England 2004; 8(17):1-144. (Guideline Ref ID GILBERT2004A)

Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness
Economic analysis: CUA (health outcome: QALYs)	Population:	*Total costs (mean per patient):	*QALYs (mean per patient):	ICER (Intervention 2 versus Intervention 1):
(ilealtii outcoille. QAL13)	Adults with low back pain (with or without sciatica)	Intervention 1: £427.21	Intervention 1: 1.03	£1527 per QALY gained (pa) 95% CI: NR
Study design: Within trial	Dationt shousehouistics.	Intervention 2: £488.28	Intervention 2: 1.07	Probability Intervention 2 cost-effective (20K
Approach to analysis: The main measure for assessing the effects on health was the EQ-5D (EuroQol-5 dimensions). The utility scores obtained at baseline, 8 months and 24 months for each participant were used to estimate QALYs. This was done by estimating the area under the lines that link the utility scores, obtained at the three time points. The Aberdeen Low Back Pain (ALBP) score, and the SF-36 (Short Form with 36 Items) were also	Patient characteristics: Mean age (intervention 1): 42.8 years Mean age (intervention 2): 43.9 years Male (intervention 1): 48.8% Male (intervention 2): 49.1% Intervention 1 (n =389): Delayed, selective imaging (no imaging unless a clear clinical indication developed) Intervention 2 (n=393): Early imaging (MRI or CT as	Incremental (2–1): £61.07 (95% CI: –25.24, 147.36; p< 0.001) *Based on imputed costs because of missing questionnaire data Currency & cost year: 2000-01 UK Pounds Cost components incorporated: The areas of treatment considered were related to hospital based services (outpatient consultation; imaging; physiotherapy; hospital admission; surgery;	Incremental (2–1): 0.04 (95% CI: –0.015, 0.10; p= 0.01) *Based on adjusted estimates taking into account differences at baseline.	Analysis of uncertainty: Bootstrapping of ICER (using adjusted QALYs) was conducted from a health care payer perspective. The results are presented above. Additional sensitivity analyses were conducted to show the effect on cost per QALY gained from changing the estimated cost of imaging. This found as the cost of imaging increases, the likelihood that 'early imaging' would be cost-effective decreases. Bootstrapping was also conducted using unadjusted QALYs. This resulted in approximately a 98% probability that early imaging was cost-effective.

reported but not used in the analysis For some areas of resource only one source of data (participant completed questionnaires or case notes) was deemed appropriate. However, for other areas of resource use the choice was informed on by the results of a small study that investigated the similarities between different methods of data collection.	soon as practicable)	injection; provision of back supports, corsets, or braces), primary care services (general practitioner visits, use of prescription and nonprescription medicines), and other tests (blood and urine tests) and devices.	
Perspective: UK NHS Follow-up: 2 years Discounting: Costs: 6%; Outcomes: 0%			

Data sources

Health outcomes: Within-trial analysis (RCT, same paper). Health outcomes included patient reported EQ-5D collected at baseline, 8 months, and 24 months follow-up. QALYs were calculated by using the area under the curve approach obtained at the three time points.

Quality-of-life weights: EQ-5D,

UK tariff. The SF-36 and Aberdeen Low Back Pain (ALBP) score were also reported, but not used to estimate QALYs. **Cost sources:** Within-trial analysis of resource use was captured alongside clinical trial via self-completed questionnaires performed at 8 and 24 months. Resource use came from either data abstraction of patients' medical notes, patient questionnaire, or patient time and travel questionnaire. In general, resource use data came from case notes to provide estimates of care in secondary care and questionnaires were used as the source of data for primary care. Costing sources were the British National Formulary and Scottish Health Service Costs. In some case, bottom-up costing was conducted, expert opinion was sought, and in one case (GP consultations) another paper was referenced.

Comments

Source of funding: Scottish Executive Health Department. **Limitations:** Discounting only applied to costs at a rate of 6%, as opposed to 3.5% for both costs and effects (NICE reference case). Within-trial analysis (same paper): Gilbert 2004 is one of a number of studies included in the clinical review for this question and may not reflect the fully body of evidence. In addition, Because of some missing questionnaire data, some resource use areas required imputation. **Other:** None.

Overall applicability^a: Partially applicable Overall quality^b(b): Potentially serious limitations

Abbreviations: 95% CI: 95% confidence interval; CUA: cost—utility analysis; EQ-5D: Euroqol 5 dimensions (scale: 0.0 [death] to 1.0 [full health], negative values mean worse than death); ICER: incremental cost-effectiveness ratio; NR: not reported; pa: probabilistic analysis; QALYs: quality-adjusted life years.

- (a) Directly applicable / Partially applicable / Not applicable
- (b) Minor limitations / Potentially serious limitations / Very serious limitations

I.4 Self-management

Table 5: Hollinghurst 2008²³

Hollinghurst S, Sharp D, Ballard K, Barnett J, Beattie A, Evans M et al. Randomised controlled trial of Alexander technique lessons, exercise, and massage (ATEAM) for chronic and recurrent back pain: economic evaluation. Spine. United Kingdom 2008; 337:a2656. (Guideline Ref ID HOLLINGHURST2008)

Study details	Population & interventions	Costs	Health outcomes	Cost	effective	ness					
Economic analysis: CUA (health outcome:	Population: People with chronic or	Total costs (mean per patient):	QALYs (mean per patient):	Full incremental analysis(a):with strategies ranked by ascending order of effectiveness							
QALYs) NB CEA also but not presented in this table. Study design: Within-	recurrent low back pain Intercruited from primary care (without sciatica). Inte	Intervention 1: £54 Intervention 2: £258 Intervention 3: £218 Intervention 4: £610 Intervention 5: £154	NR Incremental versus usual care: Intervention 1: 0	Int	Inc Cost vs base- line	Inc QALY vs base- line	Inc cost ^(b)	Inc QALY (b)	ICER ^(b)		
trial analysis (ATEAM RCT – associated clinical paper Little 2008 ^{35,36})	N: 579 Mean age: 45 (SD 11) Male: 31%	Intervention 5: £154 Intervention 6: £267 Intervention 7: £240 Intervention 8: £661 Cost breakdown Intervention cost/other cost:	Intervention 7: £240 In	Intervention 3: £134 Intervention 6: £267 Intervention 6: £267 Intervention 7: £240 Intervention 3: 0.03	Intervention 2: -0.01 Intervention 3: 0.03	2 1 3	£204 £0 £163	-0.01 0 0.03	Dominated Baseline Dominated		
Approach to analysis: Analysis of individual level data for EQ-5D and resource use. Unit costs applied.	Intervention 1: Usual care (UC) Intervention 2: Massage (6 sessions)		Intervention 5: 0.04 Intervention 6: 0.06 Intervention 7: 0.06 Intervention 8: 0.09	5 4 6 7	£100 £556 £213 £185	0.04 0.05 0.06 0.06		Dominated Dominated			
Perspective: UK NHS	Intervention 3: £159/£59			8 Proba	£607 ability cos	0.09 st effectiv	£421 ve not repo	0.03 orted for	£14,042 full		

(participant and societal perspectives also analysed but not presented here)

Follow-up: 12 months

Discounting: Costs:

n/a; Outcomes: n/a

Alexander technique (6 lessons)

Intervention 4:

Alexander technique (24 lessons)

Intervention 5:

UC + exercise prescription*

Intervention 6:

Exercise prescription* + massage (6 sessions)

Intervention 7:

Exercise prescription * + Alexander technique (6 lessons)

Intervention 8:

Exercise prescription * + Alexander technique (24 lessons)

*Exercise prescription in the study was a prescription from a doctor for home-based general exercise and a practice nurse's behavioural counselling. Intervention 4: £560/£50 Intervention 5: £30/£124 Intervention 6: £189/£79 Intervention 7: £198/£42 Intervention 8: £596/£65

Currency & cost year:

2005 UK pounds

Cost components incorporated:

Interventions (teaching and equipment), primary care contacts, outpatient appointments, inpatient hospital stays and medication.

increr	mental ar	nalyses.						
Alexa only(a		hnique stra	ategies and us	sual care				
Int (a)	Inc cost ^(b)	Inc QALY ^(b)	ICER ^(b)	Prob. CE				
Witho	out exerc	ise prescrip	otion					
1	Baselin	e						
3	£163	0.03	£5,899					
4	£392	0.02	£20,993					
With	exercise	prescriptio	n					
5	Baselin	e						
7	£86	0.02	£5,332					
8	£421 0.03 £13,914							
With	or withou	ut exercise	prescription					
1/5	Baselin	e						
3/7	£124	0.022	£5,704	NR				
4/8	£407	0.023	£17,454	NR				
Mass	age and	usual care	only(a):					
Int (a)	Inc cost ^(b)	Inc QALY ^(b)	ICER ^(b)	Prob. CE				
Witho	out exerc	ise prescrip	otion					
1	Baselin	e						
2	£204	-0.01	Dominated	~30% (£5K threshold)				
With	exercise	prescriptio	n					
5	Baselin	e						
6	£113	0.02	>90% (£5k £5,304 threshold)					
With	or withou	ut exercise	prescription					

incremental analyses

1/5	Ва	Baseline						
2/6	£1	58	0.01	.5	£10	,793	N	R
Unsu	oerv	vised	exer	ise a	nd us	ual care	onl	y(a):
Int ^(a)		Inc cost	(b)	Inc QAL	.Y ^(b)	ICER ^(b)		Prob. CE
Witho	out r	nassa	age o	r AT				
1		Base	eline					
5		£10	0	0.04		£2847		>95% (£5K threshold)
With	With or without massage or AT							
1/2/3	/4	Bas	Baseline					
5/6/7	/8	£44	44 0.04 £1096 NR				NR	

Analysis of uncertainty:

Sensitivity analyses looked at the impact of:

- 100% adherence to the interventions on cost results mostly did not change. In the AT only comparison without exercise prescription, 24 sessions now had an ICER of £26,550.(a)
- 2. The exclusion of inpatient stay costs (3 hospital stays during the trial 2 in the exercise prescription only group and 1 in the massage plus exercise group). Overall conclusions were not impacted. Although massage and exercise now dominated AT 6 lessons and exercise prescription instead of the other way round.
- 3. Using complete cases only for analysis of QALYs. The overall conclusion that 24 AT lessons were cost effective. Normal care with exercise prescription, massage or 6 Alexander technique lessons had fewer QALYs than normal care alone and higher costs and so were all dominated.
- 4. Using complete case only for analysis of personal

costs was under taken but is not reported here.

Data sources

Health outcomes: QALYs were calculated using patient-level utility data collected at baseline, 3 months and 1 year and the area under the curve approach adjusted for baseline difference across the groups. Missing data was imputed (38%). Quality-of-life weights: EQ-5D UK tariff. Cost sources: Resource use: within-trial analysis of prospectively collected data. Intervention costs based on number of attended session. Unit costs: Mostly UK national sources with some data from published sources or trial participants.

Comments

Source of funding: Medical Research Council. Limitations: Study does not include all available non-invasive treatment options; resource use data (2002-2004) and unit costs (2005) may not reflect current NHS context. Time horizon may not be sufficient to capture all benefits and costs - authors suggest that the effects of Alexander technique lessons may be longer lasting than massage or an exercise prescription. Within-trial analysis and so does not reflect full body of available evidence for all comparators. Uncertainty has not been quantified for the full incremental analysis. Usual care not described and unclear if this is was provided also in the massage and AT groups.

Overall applicability(c): partially applicable **Overall quality(d):** AT = minor limitations; massage = potentially serious limitations; exercise prescription = potential serious limitations; overall analysis = potentially serious limitations

Abbreviations: CEA: cost-effectiveness analysis; 95% CI: 95% confidence interval; CUA: cost-utility analysis; EQ-5D: Euroqol 5 dimensions (scale: 0.0 [death] to 1.0 [full health], negative values mean worse than death); ICER: incremental cost-effectiveness ratio; NR: not reported; QALYs: quality-adjusted life years

- (a) Calculated by NGC
- (b) Incremental cost/QALYs/cost effectiveness ratio compared to next most effect treatment option that is not ruled out by dominance or extended dominance. An option is ruled out by dominance when another option has higher QALYs and lower costs. An option is ruled out by extended dominance when it has a higher ICER than the next, more effective, option and so this option can never be the most cost effective.
- (c) Directly applicable/Partially applicable/Not applicable
- (d) Minor limitations/Potentially serious limitations/Very serious limitations

1.5 Exercise

Table 6: Beam 2004⁶⁰

UK BEAM Trial Team. United Kingdom back pain exercise and manipulation (UK BEAM) randomised trial: cost-effectiveness of physical treatments for back pain in primary care. Spine. 2004; 329:1381-1385:1381-1385. (Guideline Ref ID BEAM2004)

Study details	Population & interventions	Costs	Health outcomes	Cost e	effective	ness			
Economic analysis: CUA (health outcome:	Population: Low back pain mixed population (with or	Total costs (mean per patient):	QALYs (mean per patient):	•		ntion 2 ve	ersus Inte sis ^(a) :	rvention	L):
QALYs)	without sciatica).	Intervention 1: £346	Intervention 1: 0.618	Int	Cost	QALY	Inc	Inc	ICER (c)

2016

Study design: Withintrial analysis (UK BEAM RCT – associated clinical paper Underwood 2004^{61,61})

Approach to analysis: Analysis of individual level data for EQ-5D (adjusted for baseline differences) and resource use. Unit costs applied.

Perspective: UK NHS Follow-up: 1 year **Discounting:** Costs: n/a; Outcomes: n/a

Adults 18-65 years with low back pain who had experienced pain: 1) every day for the 28 days before randomisation; or for 21 out of 28 days and also 21 out of the 28 days before that. Those complaining mainly of pain below the knee were excluded.

Subgroup of full UK BEAM trial with sufficient data for economic analysis (97%).

Patient characteristics:

N = 1297

Mean age: NR (SD: NR)

Male: NR

Intervention 1: Best care (self management [SM] programme & advice to stay active)

Intervention 2: Best care + 'Back to fitness programme' (SM + biomechanical exercise) (initial assessment and up to 9 classes over 12 weeks)

Intervention 3: Best care + spinal manipulation therapy (SM + mixed modality manual therapy) (8 sessions over 12 weeks)

Intervention 4: Best care +

Intervention 2: £486 Intervention 3: £541 Intervention 4: £471 For incremental analysis see cost effectiveness column

Subanalysis exercise not available (n=623): Intervention 1: £346 Intervention 3: £541 Incremental (2-1): £195 (95% CI NR; p=NR)

not available (n=668): Intervention 1: £346 Intervention 2: £486 Incremental (2-1): £140 (95% CI NR; p=NR)

Subanalysis manipulation

Cost breakdown

Intervention cost/other costs:

Intervention 1: £0/£346 Intervention 2: £41/£445 Intervention 3: £147/£394 Intervention 4: £152/£319

Currency & cost year: 2000/1 UK pounds Cost components

Intervention 2: 0.635 Intervention 3: 0.659 Intervention 4: 0.651 For incremental analysis see cost effectiveness column

Subanalysis exercise not available (n=623): Intervention 1: 0.622 Intervention 3: 0.663 Incremental (2-1):

0.041 (95% CI NR; p=NR)

Subanalysis

manipulation not available (n=668): Intervention 1: 0.610 Intervention 2: 0.627 Incremental (2–1):

0.017 (95% CI NR; p=NR)

£346	0.618	Baseline				
2 £486 0.635 Dominated by 4						
£471	0.651	.651 £126 0.033 £3,800				
3 £541 0.659 £70 0.008 £8,700						
Probability cost-effective (£20K/30K threshold) ^(d) :						
	£486 £471 £541	£4860.635£4710.651£5410.659	£486 0.635 Dominat £471 0.651 £126 £541 0.659 £70	£486 0.635 Dominated by 4 £471 0.651 £126 0.033 £541 0.659 £70 0.008		

cost (c)

QALY

(c)

Intervention 1: 0%/0%

Intervention 2: <10%/<10% Intervention 3: >50%/>55%

Intervention 4: ~39%/~37%

Subanalysis exercise not available (n=623):

3 vs 1: £4,800 per QALY gained

95% CI: NR

(a)

(b)

Probability intervention 3 cost-effective (£20K/30K threshold)(d): >95%/100%

Subanalysis manipulation not available (n=668):

2 vs 1: £8,300 per QALY gained

95% CI: NR

Probability intervention 3 cost-effective (£20K/30K

threshold)(d): ~60%/~70%

Analysis of uncertainty: Bivariate multilevel analysis was used to quantify uncertainty due to sampling variation. Three sensitivity analyses relating to costs were undertaken:

• Exclusion of high cost outliers (>£2000): interventions 2 and 4 become ruled out by extended dominance by 3. The ICER for 3 versus 1 is £3000 per QALY gained. In subgroup analysis where manipulation is not

'Back to fitness programme'+ spinal manipulation therapy (SM + biomechanical exercise + mixed modality manual therapy) (same as above except 6 weeks of manipulation followed by 6 weeks of CPP)

incorporated:

Interventions, primary care contacts (GP, practice nurse, physiotherapist, other), secondary care contacts (hospital admissions and outpatient appointments).

- available the ICER for intervention 2 versus 1 was £4100.
- Costing assuming NHS buys all manipulation from private sector: ICERs increased to £8600 (4 versus 1) and £10,600 (3 versus 4)
- Costing assuming NHS buys some manipulation from private sector (as per trial rates): ICERs increased to £6600 (4 versus 1) and £8700 (3 versus 4)

Data sources

Health outcomes: QALYs were calculated using patient-level utility data collected at baseline, 3 and 12 months and the area under the curve approach adjusted for baseline differences across the groups. Quality-of-life weights: Within-RCT analysis: EQ-5D UK tariff. Resource use: Within-RCT analysis. Intervention cost was based on the number of attended sessions. Cost sources: UK national sources for NHS provided care and a major insurance provider for privately provided care. Base case analysis costs all manipulation as provided by NHS irrespective of how provided in trial (explored in sensitivity analysis).

Comments

Source of funding: Medical Research Council & NHS **Limitations:** Study does not include all non-invasive treatment options. Resource use data (1999-2002) and unit costs (2000/01) may not reflect the current NHS context. A longer time horizon may be preferable given than interventions continued to show benefit at 12 months. Within-trial analysis and so does not reflect full body of available evidence for this intervention; Underwood 2004 is 1 of 8 studies included in the clinical review for mixed manual therapy – although the only one compared to usual care and with EQ5D data. **Other:**

Overall applicability^(e): Partially applicable Overall quality^(f): mixed MT = Minor limitations

Abbreviations: 95% CI: 95% confidence interval; CUA: cost—utility analysis; EQ-5D: Euroqol 5 dimensions (scale: 0.0 [death] to 1.0 [full health], negative values mean worse than death); ICER: incremental cost-effectiveness ratio; NR: not reported; QALYs: quality-adjusted life years

- (a) Intervention number in order of least to most effective in terms of QALYs
- (b) Total cost/QALYs
- (c) Incremental cost/QALYs/cost effectiveness ratio compared to next most effect treatment option that is not ruled out by dominance or extended dominance. An option is ruled out by dominance when another option has higher QALYs and lower costs. An option is ruled out by extended dominance when it has a higher ICER than the next, more effective, option and so this option can never be the most cost effective.
- (d) Estimated from graph
- (e) Directly applicable / Partially applicable / Not applicable
- (f) Minor limitations / Potentially serious limitations / Very serious limitations

Table 7: Chuang 2012^{7,8}

Chuang LH, Soares MO, Tilbrook H, Cox H, Hewitt CE, Aplin J et al. A pragmatic multicentered randomized controlled trial of yoga for chronic low back pain: economic evaluation. Spine. 2012; 37(18):1593-1601. (Guideline Ref ID CHUANG2012)

Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness
Study details Economic analysis: CUA (health outcome: QALYs) Study design: Withintrial analysis (RCT – associated clinical paper Tilbrook 2011 ^{57,57}) Approach to analysis: Analysis of individual level data for EQ-5D and resource use with missing data imputed and adjusted for baseline differences. Unit costs applied. Perspective: UK NHS (societal also analysed but not presented here) Follow-up: 12 months Discounting: Costs: n/a; Outcomes: n/a	Population & interventions Population: People 18-65 that had consulted their GP for low back pain in past 18 months. Patient characteristics: N: 313 Mean age: 46 years (SD 11) Male: 30% Intervention 1: Usual care including The Back Book, and one yoga class after the final follow-up. Intervention 2: Yoga (75 minute weekly group class [maximum 15 participants] for 12 weeks, relaxation CD, yoga manual, yoga mat; participants were encouraged to practice at home for 30 minutes daily or at least 2 times per week and use the relaxation CD) plus usual care including The Back Book.	Total costs (mean per patient): Intervention 1: NR Intervention 2: NR Incremental (2–1): £507 (95% CI £159 to £855); p=NR) Cost breakdown (unadjusted and without imputation) Intervention cost/NHS costs: Intervention 1: £0/£530 Intervention 2: £293/£762 Currency & cost year: 2008/9 UK pounds Cost components incorporated: Intervention, primary care contacts (GP, practice nurse, physiotherapist and other) and secondary care contacts (emergency service, outpatient appointments, inpatient hospital stays,	Health outcomes QALYs (mean per patient): Intervention 1: NR Intervention 2: NR Incremental (2–1): 0.037 (95% CI 0.006 to 0.069; p=NR)	ICER (Intervention 2 versus Intervention 1): £13,606 per QALY gained 95% CI: NR Probability intervention 2 cost-effective (£20K/30K threshold): 72%/~87% Analysis of uncertainty: Method for estimating probability cost effective was not stated. As an alternative to using results based on imputing missing data, complete case analysis was undertaken: ICER: £9,266 per QALY gained The impact of the cost of yoga was explored. While the value of the ICER did change, yoga remained cost effective even when a higher cost of £486 (based on the cost of cardiac rehabilitation) was used.

Data sources

Health outcomes: QALYs were calculated using patient level utility data collected at baseline, 3, 6 and 12 months and the area under the curve approach adjusted for baseline differences across the groups. Missing data was imputed (usual care 23%; Yoga 28%). Quality-of-life weights: Within-RCT analysis: EQ-5D, tariff used is not stated although as this is a UK study it is judged likely to be the UK tariff. Resource use: within-trial analysis of prospectively collected data adjusted for baseline differences across the groups. Missing data was imputed (usual care 18%; yoga 26%). Intervention cost was the average cost per patient based on total cost of classes

and equipment and total number of patients. Unit costs: Mostly UK national sources with some data from published sources or trial participants.

Comments

Source of funding: Arthritis Research UK. Limitations: Study does not include all non-invasive treatment options. The EQ-5D tariff used is not stated although as this is a UK study it is judged likely to be the UK tariff. Follow-up may not be sufficient to capture all benefits and costs - authors suggest that if participants continue to practice yoga it might continue to have an impact on their back function and they noted that 60% of participants in the yoga arm who answered the question continued practising yoga at home. Medication costs are not included. Within-trial analysis and so does not reflect full body available evidence for this comparison - Tilbrook is 1 of 7 studies that included this comparison.

Overall applicability^(a): partially applicable Overall quality^(b): potentially serious limitations

Abbreviations: 95% CI, 95% confidence interval; CUA, cost—utility analysis; EQ-5D, EuroQol 5 dimensions (scale: 0.0 [death] to 1.0 [full health], negative values mean worse than death); ICER, incremental cost-effectiveness ratio; NR, not reported; QALYs: quality-adjusted life years

- (a) Directly applicable/Partially applicable/Not applicable
- (b) Minor limitations/Potentially serious limitations/Very serious limitations

Table 8: Critchley 20079

Critchley DJ, Ratcliffe J, Noonan S, Jones RH, Hurley M, V. Effectiveness and cost-effectiveness of three types of physiotherapy used to reduce chronic low back pain disability: a pragmatic randomized trial with economic evaluation. Spine. 2007; 32(14):1474-1481. (Guideline Ref ID CRITCHLEY2007)

Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness
Economic analysis: CUA (health outcomes: QALYs) Study design: Withintrial analysis (RCT – clinical results in same paper) Approach to analysis: Analysis of individual level data for EQ-5D (adjusted for baseline differences in utility) and resource use. Unit costs applied. Perspective: UK NHS Follow-up: 18	Population: 18 years old or older, low back pain >12 weeks duration with or without leg symptoms or neurologic signs Patient characteristics N = 212 Mean age = 44 Male = 35.8% Intervention 1: Biomechanical exercise. Spinal stabilisation physiotherapy; individual transversus abdominis and multifidus muscle training, group spinal stability exercises, maximum of 8 supervised sessions of 90 minutes. (n=72) Intervention 2: Combination: Manual therapy plus self-management. Individual physiotherapy; a combination of joint	Total costs (mean per patient): Intervention 1: £379 Intervention 2: £474 Incremental 3: £165 Incremental (2–1): £95 (95% CI: NR; p=NR) Incremental (3–1): -£214 (95% CI: NR; p=0.16) Incremental (3–2): -£309 (95% CI: NR; p=0.16) Cost breakdown (initial treatment/other) Intervention 1: £80/£299	QALYs (mean per patient): Intervention 1: 0.90 Intervention 2: 0.99 Intervention 3: 1.00 Incremental (2–1): 0.09 (95% CI: NR; p=NR) Incremental (3–1): 0.10 (95% CI: NR; p=NR) Incremental (3–2): 0.01 (95% CI: NR; p=NR)	Fully incremental analysis MBR programme dominates both biomechanical exercise and combined manual therapy and self- management with higher QALYs and lower costs 95% CIs: NR Probability cost-effective (£20K/30K threshold): Intervention 1: ~33%/~35% Intervention 2: ~0%/~0% Intervention 3: 67%/65% Analysis of uncertainty: Sensitivity analysis testing multiple scenarios; a)

months

Discounting: Costs: 3.5%; Outcomes: 3.5%

mobilisations, joint manipulation and massage, trunk muscle retraining, stretching and spinal mobility exercises taught to perform at home, back care advice; up to 12 sessions of 30 minutes. (n=71)

Intervention 3:

MBR programme (3 elements: physical, cognitive, education). Structured back pain education, group general strengthening, stretching and aerobic exercises, cognitive-behavioural approach to reduce fear, encourage self-management; maximum of 8 supervised sessions of 90 minutes. (n=69)

Intervention 2: £90/£384 Intervention 3: £75/£90

Currency & cost year:

2003 UK pounds

Cost components
incorporated:

Physiotherapy, other healthcare visits (GP, consultant, other NHS, investigations, inpatient procedures), medication including patients with imputed missing data, b) excluding costly outliers In both cases the pain management program continues to be the most cost effective option.

Costs excluding spinal surgery patients:

Intervention 1: £188 Intervention 2: £401 Incremental 3: £165

Data sources

Health outcomes: QALYs were calculated using patient-level utility data collected at baseline, 6, 12 and 18 months and the area under the curve approach adjusted for baseline utility. Quality-of-life weights: EQ5D, tariff used not stated (although as this is a UK study it is judged likely to be UK tariff) Cost sources: resource use was captured through physiotherapy notes and cost questionnaires, unit costs were obtained from the personal social services research unit database, NHS reference costs, and British National Formulary

Comments

Source of funding: NR **Limitations:** Resource use data (2002-2005) and unit costs (2003/3) may not reflect the current NHS context. EQ-5D tariff used is not stated (although as UK study judged likely to be UK tariff). Study does not include all non-invasive treatment options. Time horizon may not be sufficient to capture all benefits and costs if benefits persist beyond 18 months. Within-trial analysis and so does not reflect full body of available evidence for this intervention; Critchley 2007 is 1 of 19 studies included in the clinical review for MBR.

Overall applicability^(a): partially applicable Overall quality^(b): minor limitations

Abbreviations: 95% CI: 95% confidence interval; CUA: cost—utility analysis; da: deterministic analysis; EQ-5D: Euroqol 5 dimensions (scale: 0.0 [death] to 1.0 [full health], negative values mean worse than death); ICER: incremental cost-effectiveness ratio; NR: not reported; pa: probabilistic analysis; QALYs: quality-adjusted life years; CSRI: client services receipt inventory (a) Directly applicable / Partially applicable / Not applicable

(b) Minor limitations / Potentially serious limitations / Very serious limitations

Table 9: Niemisto 2003^{41,41}/Niemisto 2005^{40,41}

Niemisto L, Lahtinen-Suopanki T, Rissanen P, Lindgren KA, Sarna S, Hurri H. A randomized trial of combined manipulation, stabilizing exercises, and physician consultation compared to physician consultation alone for chronic low back pain. Spine. 2003; 28(19):2185-2191. (Guideline Ref ID NIEMISTO2003)

Niemisto L, Rissanen P, Sarna S, Lahtinen-Suopanki T, Lindgren K-A, Hurri H. Cost-effectiveness of combined manipulation, stabilizing exercises, and physician

consultation compared to physician consultation alone for chronic low back pain: a prospective randomized trial with 2-year follow-up. Spine. 2005; 30(10):1109-1115. (Guideline Ref ID NIEMISTO2005)

1115. (Guideline Ref ID NIEMISTO2								
Study details	Population & interventions	Costs (d)	Health outcomes	Cost effectiveness				
Economic analysis: CCA (various health outcomes) Study design: Within-trial analysis (RCT – clinical results in same paper) Approach to analysis: Analysis of individual level data for health outcomes and resource use. Unit costs applied. Perspective: Dutch healthcare costs (societal costs analysed but not presented here) Follow-up 12/24 months Discounting: Costs: 0%; Outcomes: 0%	Population: 24-46 years with chronic low back pain (with or without sciatica) of at least 3 months duration with ODI was at least 16%. Severe sciatica in the straight leg raising test with less than 35 degrees was an exclusion criterion. Patient characteristics N = 204 Mean age = 37 years (SD: NR) Male = 46% Intervention 1: Self management programme. Physician consultation alone; clinical evaluation (60 minutes) plus educational booklet, instruction regarding posture and spinal exercise recommendation. (n=102) Intervention 2: Combination: Self management programme ,manual therapy (manipulation/mobilisation) and biomechanical exercise. As intervention 1 plus manipulation using muscle energy technique and muscle control and stabilising exercises, treatment and exercise weekly sessions for 5 weeks. (n=102)	12 months: total costs (mean per patient): Intervention 1: £278 Intervention 2: £303 Incremental (2–1): £25 (95% CI: NR; p=NS) 24 months: Annual total costs (mean per patient): Intervention 1: £234 Intervention 2: £289 Incremental (2–1): £56 (95% CI: NR; p=NS) Cost breakdown of intervention/other costs not reported. Currency & cost year: 2000 Finland Euros presented as 2000 US dollars (presented here as 2000 UK pounds ^(a)) Cost components incorporated: Visits to physicians, visits to physiotherapy, outpatient visits, inpatient care, x-ray examinations	24 months VAS (mean per patient): Intervention 1: NR Intervention 2: NR Incremental (2–1): 4.97 (95% CI: 4.83 to 5.12; p=NR) ODI (mean per patient): Intervention 1: NR Intervention 2: NR Incremental (2–1): 1.24 (95% CI: 1.18 to 1.30; p=NR) 15D (mean per patient): Authors report no difference in 15D.	Analysis of uncertainty: Uncertainty around the point estimates of incremental effects was assessed through bootstrapping but for societal costs not healthcare costs.				
Data sources								

Health outcomes: Within-trial analysis (measurements at baseline, 5, 12, 24 months). **Quality-of-life weights:** 15D utility instrument, Finnish population, VAS-based tariff. **Cost sources:** Within-trial analysis of resource use was captured through cost questionnaires administered at baseline, 12, 24 months. Finnish standard national prices used (average costs of Finnish healthcare providers).

Comments

Source of funding: The social insurance institute of Finland and Finska Lakarsallskapet. Limitations: Finnish resource use data (1999-2001) and unit costs (2000) may not reflect the current NHS context. Non-NICE reference case utility measure used (15D) and this uses a non-comparable valuation method (VAS) from the Finnish population. QALYs were not calculated using area under the curve. Discounting was not applied (24 month analysis). Study does not include all non-invasive treatment options. Within-trial analysis and so does not reflect full body of available evidence for this comparison Niemisto 2003 is 1 of several studies included in the clinical review for individual combinations. Limited sensitivity analysis.

Overall applicability^(a): partially applicable Overall quality^(c): potentially serious limitations

Abbreviations: CCA: cost—consequence analysis; 95% CI: 95% confidence interval; CUA: cost—utility analysis; da: deterministic analysis; NR: not reported; pa: probabilistic analysis; ODI: oswestry disability index; VAS: visual analogue scale

- (a) Converted using 2000 purchasing power parities⁴²
- (b) Directly applicable / Partially applicable / Not applicable
- (c) Minor limitations / Potentially serious limitations / Very serious limitations
- (d) Original analysis adopted a societal perspective, costs presented here were re-estimated to reflect NHS perspective only

Table 10: Smeets 2009⁴⁷

Smeets RJ, Severens JL, Beelen S, Vlaeyen JW, Knottnerus JA. More is not always better: Cost-effectiveness analysis of combined, single behavioral and single physical rehabilitation programs for chronic low back pain. European Journal of Pain. 2009; 13(1):71-81. (Guideline Ref ID SMEETS2009)

Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness
Economic analysis: CUA (health outcome: QALYs) Study design: Within-trial analysis (RCT – associated clinical paper Smeets 2006/2008a ^{48,49} Approach to analysis: Analysis of individual level data for EQ-5D (adjusted for baseline differences in	Population: 18-65 years, low back pain for more than 3 months resulting in disability (RDQ >3) and ability to walk at least 100m. With or without sciatica. Patient characteristics N = 160 Mean age: 42 years (SD: 10) Male: 55% Intervention 1:	Total costs (mean per patient): Intervention 1: £2089 Intervention 2: £1182 Intervention 3: £2618 Incremental (2–1): saves £908 (95% CI: NR; p=NR) Incremental (3–1): £530 (95% CI: £120 to £897; p=NR) Incremental (3–2): £1433 (95% CI: £1166 to £1688; p=NR)	QALYs (mean per patient): Intervention 1: 0.693 Intervention 2: 0.723 Intervention 3: 0.679 Incremental (2–1): 0.03 (95% CI: NR; p=NR) Incremental (3–1): -0.014 (95% CI: -0.094 to 0.066; p=NR) Incremental (3–2): -0.045 (95% CI: -0.119 to 0.029; p=NR)	Full incremental analysis: cognitive behavioural approaches dominates both exercise and combination treatment with higher QALYs and lower costs. 95% CI: NR Probability cost-effective (£20K/30K threshold): NR Analysis of uncertainty: Bootstrapping used to quantify uncertainty around ICER but for

utility) and resource use. Unit costs applied.

Perspective: Netherlands direct health care costs (societal also analysed but not presented here)

Follow-up: 62 weeks **Discounting:** Costs: n/a;

Outcomes: n/a

Mixed modality exercise. 30 minutes aerobic training on bicycle and 75 minutes strength and endurance training of their lower back and upper leg muscles, 3 times a week during 10 weeks.

Intervention 2:

Cognitive behavioural approach. Operant behavioural graded activity training (physiotherapist or occupational therapist, 3 group sessions and a maximum of 17 individual sessions of 30 minutes, no physical training element) and problem solving training (clinical psychologist or social worker, 10 sessions of 1.5 hours to a maximum of 4 patients at a time)

Intervention 3:

MBR programme (2 core elements: physical and cognitive). Combination of interventions 1 and 2. Therapists were told about the integrative nature of combination treatment.

Cost breakdown of intervention/other costs not reported.

Total lost productivity costs (mean per patient):

Incremental (3–1): -£1137 (95% CI: -£6706 to £4511; p=NR) Incremental (3–2): £3051 (95% CI: -£2933 to £8862; p=NR)

Currency & cost year:

2003 Netherlands euros (presented here as 2003 UK pounds(a))

Cost components incorporated:

Interventions, GP, medical specialist including radiology, occupational physician, physiotherapist, manual therapist, Cesar or Mensensieck therapist, psychologist, medication, hospitalisation, medical procedures.

societal costs not direct medical

Analysis where utility analysis was not adjusted for baseline utility: QALYs for 3-1 changed from -0.01 to 0.01. However, intervention 2 still had the highest QALYs and lowest costs.

Data sources

Health outcomes: QALYs were calculated using patient-level utility data collected at baseline, 3, 6 and 12 months and the area under the curve approach adjusted for baseline utility. Missing data was imputed. Quality-of-life weights: EQ-5D, UK tariff. Costs: Costs were calculated using patient-level resource use data collected during the 10 weeks treatment period, 1-12, 13-24, 25-36 and 37-52 weeks post treatment. Patients who did not return at least 3 cost diaries were excluded, otherwise missing data was imputed. Intervention cost was based on the number of attended sessions (mean intervention costs not reported). Unit costs were based on Dutch

national sources.

Comments

Source of funding: Netherlands Organization for Health Research and Development. **Limitations:** Dutch resource use data (2002-2004) and unit costs (2003) may not reflect current NHS context. Study does not include all non-invasive treatment options. Within-trial analysis and so does not reflect full body of available evidence for this intervention; Smeets 2006a is 1 of 7 studies included in the clinical review for mixed modality exercise, 1 of 5 where the mix was biomechanical + aerobic, although is the only one compared with cognitive behavioural approaches; 1 of 9 studies included in the clinical review for cognitive behavioural approach and one of 19 for MBR programmes. **Other:**

Overall applicability(b): partially applicable Overall quality(c): potentially serious limitations

Abbreviations: 95% CI: 95% confidence interval; CUA: cost—utility analysis; EQ-5D: Euroqol 5 dimensions (scale: 0.0 [death] to 1.0 [full health], negative values mean worse than death); ICER: incremental cost-effectiveness ratio; NR: not reported; QALYs: quality-adjusted life years

- (a) Converted using 2003 purchasing power parities⁴²
- (b) Directly applicable / Partially applicable / Not applicable
- (c) Minor limitations / Potentially serious limitations / Very serious limitations

I.6 Postural therapy

For Hollinghurst 2008²³ please see Table 5 (Self-management) above.

I.7 Orthotics

None.

I.8 Manual therapy

For Beam 2004⁶⁰ please see Table 6 (Exercise) above.

For Hollinghurst 2008²³ please see Table 5 (Self-management) above.

Table 11: Vavrek 2014⁶⁴

Vavrek D, Sharma R, Haas M. Cost-analysis related to dose-response for spinal manipulative therapy for chronic low back pain: outcomes from a randomized controlled trial. Journal of Alternative and Complementary Medicine. 2014; 20(5):A18. (Guideline Ref ID VAVREK2014)

Study details **Population & interventions** Costs **Health outcomes** Cost effectiveness **Economic analysis:** Population: Total costs (unadjusted mean per **QALYs** (unadjusted ICER: CCA (various health patient): mean per patient): Adults with low back pain 3 vs 1: £14,800 (calculated by NGC outcome) based on unadjusted data) without sciatica >3 months. Intervention 1: £206 Intervention 2: Intervention 1: 0.81 £540 Patient characteristics: Intervention 2: 0.80 Study design: Within-Intervention 3: £502 N = 400Intervention 3: 0.83 ICER based on adjusted data NR. Note trial analysis (RCT -Intervention 4: £586 that QALY gain in adjusted analysis Mean age (range between Intervention 4: 0.81 associated clinical potentially lower than in unadjusted arms): 40.9-41.8 (SD:13.8paper Haas 2014¹⁶) analysis. 14.8) Incremental (3-1): £296 Incremental (3-1): 0.02 Approach to analysis: Male (range between arms): (95% CI NR; p=NR) (95% CI NR; p=NR) Analysis of individual 48-51% Full incremental analysis was not level data for resource reported in study as differences in Cost breakdown QALYs (adjusted use. Unit costs applied. QALYs between interventions and **Intervention 1:** Sham Intervention cost/other costs: analysis) Costs imputed for across time was not statistically **Intervention 2:** Spinal Relative to weeks not covered by Intervention 1: £0/£206 significant. manipulation therapy (SMT) patient reports. Intervention 1 (sham) Intervention 2: £133/£407 6 sessions each dose of SMT Adjusted cost ratios Intervention 3: £266/£236 Probability CE was not reported. Intervention 2: SMT 12 and QALY based on vielded an additional Intervention 4: £399/£188 session 0.00 to 0.01 QALYs. No regression analyses. **Analysis of uncertainty:** A sensitivity Intervention 2: SMT 18 significant differences Perspective: USA **Adjusted cost ratios** analysis was conducted where the sessions between groups. direct medical costs weeks not covered by patient reports Follow-up: 1 year **1** Intervention 2 vs 1: 1.15 (95% CI: 0.63 were excluded from the cost analysis. **Discounting:** Costs: The results were similar to the base case to 2.11) n/a; Outcomes: n/a analysis. Intervention 3 vs 1: 1.18 (95% CI: 0.64 to 2.18) Intervention 4 vs 1: (95% CI: 0.78 (0.43 to 1.43) **Currency & cost year:** 2009 US dollars (presented here as 2009 UK pounds^(a)) **Cost components incorporated:** Interventions (reported separately in paper but added in to unadjusted costs

above; excluded from cost ratio analysis), primary care contacts (GP, practice nurse, physiotherapist, other), secondary care contacts (surgeon/neurologist and psychologist/psychiatrist consultations, emergency department visits and other), chiropractic manipulation, massage therapy and patient reported medication for low back pain.

Data sources

Health outcomes: QALYs were calculated using patient-level utility data collected at baseline, 12, 24, 39 and 52 weeks. Quality-of-life weights: Within-RCT analysis: EQ-5D, tariff not stated. Resource use: Within-RCT analysis. Intervention cost was based on the number of attended sessions. Cost sources: Within-trial resource use and 'resource-based relative value units'. Unit costs from Medicare 2009 national non-facility (i.e. non-hospital) payments.

Comments

Source of funding: NR. **Limitations:** Study does not include all non-invasive treatment options. USA resource use data (2007-2011) and unit costs (2009) may not reflect current NHS context. EQ-5D tariff used unclear. Within-trial analysis and so does not reflect full body of available evidence for this comparison; Haas 2014 is 1 of 8 included studies comparing manipulation/mobilisation to sham. Cost per QALY results were not reported (although QALYs were estimated); here the ICER has been calculated based on the reported unadjusted cost and QALY result however authors undertake a regression analysis to adjust costs and QALYs. Only minimal sensitivity analyses were carried out to quantify uncertainty.

Overall applicability^(b): Partially applicable Overall quality^(c): Potentially serious limitations

Abbreviations: CCA: cost—consequence analysis; 95% CI: 95% confidence interval; da: deterministic analysis; EQ-5D: Euroqol 5 dimensions (scale: 0.0 [death] to 1.0 [full health], negative values mean worse than death); ICER: incremental cost-effectiveness ratio; NR: not reported; pa: probabilistic analysis; QALYs: quality-adjusted life years

- (a) Converted using 2009 purchasing power parities⁴²
- (b) Directly applicable / Partially applicable / Not applicable
- (c) Minor limitations / Potentially serious limitations / Very serious limitations

I.9 Acupuncture

Table 12: Ratcliffe 2006^{45,45}, Thomas 2005^{54,56}

Thomas KJ, MacPherson H, Ratcliffe J, Thorpe L, Brazier J, Campbell Mea. Longer term clinical and economic benefits of offering acupuncture care to patients with chronic low back pain. Health Technology Assessment. 2005; 9:iii-x:iii-iix. (Guideline Ref ID THOMAS2005)

Ratcliffe J, Thomas KJ, MacPherson H, Brazier J. A randomised controlled trial of acupuncture care for persistent low back pain: cost effectiveness analysis. British

2016

Medical Journal. 2006; 333:626-628:626-628. (Guideline Ref ID RATCLIFFE2006) Study details **Population & interventions** Costs **Health outcomes Cost effectiveness Economic analysis:** Population: Total costs (mean per QALYs (mean per patient): ICER (Intervention 2 versus Intervention 1): CUA (health outcome: patient): Adults 18-65 years with low EQ-5D complete case EQ-5D complete case analysis QALYs) back pain (with or without EQ-5D complete case analysis analysis(n=85) £3598 per QALY gained sciatica) of 4-52 weeks (n=85)Intervention 1: NR 95% CI: £188 to £22,149 Study design: Withinduration. Intervention 1: NR Intervention 2: NR Probability Intervention 2 cost-effective trial analysis (RCT -Intervention 2: NR Incremental (2-1): 0.071 (£20K/30K threshold): NR associated clinical Patient characteristics: Incremental (2-1): £255 (95% CI -0.036 to 0.178; paper Thomas 2005⁵⁴ N = 241(95% CI £203 to £387: p=NR) SF-6D complete case analysis and Thomas 2006⁵⁵) Mean age: 43 years (SD: 11) p<0.05) £4241 per QALY gained Approach to analysis: Male: 40% SF-6D complete case 95% CI: £191 to £28,026 Analysis of individual SF-6D complete case analysis analysis (n=122) level data for EQ-Probability Intervention 2 cost-effective (n=122)Intervention 1: 1.426 Intervention 1: 5D/SF-6D and resource (£20K/30K threshold): ~97%/~100% Intervention 1: £345 use. Unit costs applied. Usual care (at discretion of Intervention 2: 1.453 GP). Intervention 2: £460 Incremental (2-1): 0.027 Analysis of uncertainty: Intervention 2: Incremental (2-1): £115 **Perspective: UK NHS** (95% CI -0.056 to 0.110; Bootstrapping was undertaken to estimate (95% CI -£40 to £269; p=NR) (societal also analysed Acupuncture (initial p=NR) uncertainty around the ICER. but not presented consultation and treatment plus up to nine further here) Cost breakdown (n=181) Alternative analyses: treatment) plus usual care. Follow-up: 2 years Intervention cost/other NHS • SF-6D analysis with missing data imputed **Discounting:** Costs: costs: for costs and QALYs: £4209 per QALY 3.5%; O outcomes: Intervention 1: £0/£332 gained (95% CI £182 to £27,899) 3.5% Intervention 2: £214/£257 • Excluding those permanently unable to work: £2104 per QALY gained (95% CI £128 to £19,340) **Currency & cost year:** 2002/3 UK pounds **Cost components** incorporated: Intervention, primary care contacts (GP, practice nurse, non-study intervention NHS

acupuncture, chiropractic, osteopathy, other) and secondary care contacts (emergency service, inpatient hospital stays, outpatient appointments (generic, pain clinic, physiotherapy), physiotherapy at GP surgery).

Data sources

Health outcomes: QALYs were calculated using patient-level utility data collected at baseline, 3, 12 and 24 months and the area under the curve approach adjusted for baseline differences across the groups. Those with complete case utility and cost data were used in the cost-effectiveness analysis base case. Quality-of-life weights: Within-RCT analysis: EQ-5D, UK tariff and SF-6D, UK tariff. Resource use: Within-trial analysis of prospectively collected data. Intervention cost was based on the number of attended sessions. Unit costs: Mostly UK national sources with some data from trial participants.

Comments

Source of funding: UK NHS Executive health technology programme. Limitations: Study does not include all non-invasive treatment options. Resource use data (1999-2002) and unit costs (2002/3) may not reflect the current NHS context. Within-trial analysis and so does not reflect full body of available evidence for this comparison; Thomas 2005/Thomas 2006 is 1 of 16 included studies comparing acupuncture to usual care. The probability cost effective is not reported for the EQ-5D based analysis. Other:

Overall applicability^(a): Partially applicable Overall quality^(b): potentially serious limitations

Abbreviations: 95% C,: 95% confidence interval; CUA, cost—utility analysis; EQ-5D, Eurogol 5 dimensions (scale: 0.0 [death] to 1.0 [full health], negative values mean worse than death); ICER, incremental cost-effectiveness ratio; NR, not reported; QALYs, quality-adjusted life years

- (a) Directly applicable / Partially applicable / Not applicable
- (b) Minor limitations / Potentially serious limitations / Very serious limitations

Electrotherapy 1.10

None.

1.11 **Psychological**

Table 13: Jellema2007^{24,26}

Jellema P, van der Roer N, Van Der Windt DAWM, van Tulder MW, Van Der Horst HE, Stalman WAB et al. Low back pain in general practice: Cost-effectiveness of a minimal psychosocial intervention versus usual care. European Spine Journal. 2007; 16(11):1812-1821. (Guideline Ref ID JELLEMA2007)

Study details

Population & interventions Population: Adults (18-65 years) with low back pain of >12 weeks duration or exacerbation of mild symptoms. With or without sciatica. Patient characteristics N = 250 (cost analysis complete cases)/213 (costs and QALYs complete cases) Mean age: 43 years (SD: NR) Male: 52% Intervention 1: Usual care (Provided by GP; Dutch national guidelines which recommend wait and

no explicit content but assumed would follow see <6weeks and referral for physical therapy 6-12weeks if persistent disability. Explicit guidance on psychosocial factors is lacking.) Intervention 2: Minimal intervention strategy (categorised as cognitive behavioural approaches) - 20 minute GP consultation aimed at

identification and discussion

Costs **Health outcomes Cost effectiveness** Total costs (mean per QALYs (mean per patient): ICER (Intervention 2 versus Intervention 1): patient): Intervention 1: 0.837 Intervention 1 dominant (lower costs and Intervention 1: £122 better health outcomes Intervention 2: 0.833 Intervention 2: £126 95% CI: NR Incremental (2-1): 0.004 Incremental (2-1): £4 **QALYs** lost Probability Intervention 2 cost-effective (£20K/30K threshold): NR (95% CI: -£45 to £51; p=NS) (95% CI: NR; p=NR) Analysis of uncertainty: Bootstrapping is Cost breakdown (primary reported as undertaken to estimate care/secondary uncertainty around the ICER but results are care/medication)(b) not reported for the cost per QALY analysis. Intervention 1: £106/£16/£6 Intervention 2: £111/£15/£6 As an alternative to the complete case analysis undertaken for the base case **Currency & cost year:** analysis, an analysis was undertaken where 2002 Dutch Euros all missing cost data was imputed. However, (presented here as 2002 UK results are reported for total costs only and pounds(a))

Cost components incorporated: Primary care (GP, intervention costs, physical therapist, manual therapist, exercise therapist, back school, chiropractor, physiofitness program, professional home carer, psychologist), secondary care (outpatient appointments, hospitalization, surgery, radiograph, MRI scan),

medication. (Other non-

direct healthcare costs alone are not available.

ь.		

of psychosocial factors	health care costs were
covering exploration,	complementary care,
information and self-care	informal care, equipment
aspects; a follow-up	aids and absenteeism from
appointment was	paid and unpaid work but
recommended.)	not reported here.)

Data sources

Health outcomes: QALYs were calculated using patient-level utility data collected at baseline, 3, 6 and 12 months and the area under the curve approach. Complete case analysis was used. **Quality-of-life weights:** EQ-5D, UK tariff. **Costs:** Costs were calculated using patient-level resource use data collected for periods of baseline-3 months, 3-6 months, 6-9 months and 9-12 months. Complete case analysis was used. Mean intervention costs were not reported separately. Unit costs were based on Dutch national sources.

Comments

Source of funding: Netherlands Organization for Health Research and Development. **Limitations:** Dutch resource use data (2001-2003) and unit costs (2002) may not reflect current NHS context. Study does not include all non-invasive treatment options. Within-trial analysis and so does not reflect full body of available evidence for this comparison; Jellema2005 is 1 of 9 studies included in the clinical review for cognitive behavioural approach - although 1 of 2 compared to usual care with EQ5D data. No exploration of uncertainty available relevant to guideline. **Other:**

Overall applicability(c): partially applicable Overall quality(d): potentially serious limitations

Abbreviations: 95% CI: 95% confidence interval; CUA: cost—utility analysis; EQ-5D: Euroqol 5 dimensions (scale: 0.0 [death] to 1.0 [full health], negative values mean worse than death); ICER: incremental cost-effectiveness ratio; NR: not reported; NS: not significant (at 0.05); QALYs: quality-adjusted life years

- (a) Converted using 2002 purchasing power parities⁴²
- (b) Intervention costs were not reported as a separate category
- (c) Directly applicable/Partially applicable/Not applicable
- (d) Minor limitations/Potentially serious limitations/Very serious limitations

Table 14: Lamb 2010^{29,30}

Lamb SE, Lall R, Hansen Z, Castelnuovo E, Withers EJ, Nichols V et al. A multicentred randomised controlled trial of a primary care-based cognitive behavioural programme for low back pain. the back skills training (BeST) trial. Health Technology Assessment. 2010; 14(41):1-281. (Guideline Ref ID LAMB2010A)

Lamb SE, Hansen Z, Lall R, Castelnuovo E, Withers EJ, Nichols V et al. Group cognitive behavioural treatment for low-back pain in primary care: a randomised controlled trial and cost-effectiveness analysis. Lancet. United Kingdom 2010; 375(9718):916-923. (Guideline Ref ID LAMB2010B)

Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness
Economic analysis: CUA (health outcome: QALYs)	Population: Adults (18+) with at least moderately troublesome low back pain	Total costs (mean per patient):	QALYs (mean per patient): Intervention 1: 0.604	ICER (Intervention 2 versus Intervention 1): £1786 per QALY gained

2016

Study design: Within-trial analysis (RCT – associated clinical paper Lamb 2012^{31,32}

Approach to analysis:

Analysis of individual level data for EQ-5D (adjusted for baseline differences) and resource use. Unit costs applied.

Perspective: UK NHS **Follow-up:** 1 year

Discounting: Costs: n/a;

Outcomes: n/a

of >6 weeks duration, and had consulted for low back pain in primary care within the preceding 6 months.

Patient characteristics

N = 528 (cases with complete follow-up at least for 3 months)

Mean age: 55 years (SD: NR)

Male: 41%

Intervention 1:

Self management. Active management in general practice (a 15-min session with a nurse or physiotherapist - advice to remain active, avoid bed rest and appropriate pain medication usage and symptom management; provision of the Back Book).

Intervention 2:

Self management (active management) + cognitive behavioural approach (1.5hr individual assessment and 6 group sessions; delivered by physiotherapist, nurse, psychologist or occupational therapist)

Intervention 1: £279 Intervention 2: £457 Incremental (2–1): £178

Cost (unadjusted) breakdown (initial treatment/other)

(95% CI: NR; p=NR)

Intervention 1: £17/£207 Intervention 2: £204/£217

Currency & cost year:

2008 UK pounds

Cost components incorporated:

Intervention costs (contact time, non-contact time [e.g. writing notes, admin, travel], supervisory support time, consumables, equipment, training); other NHS resource use (contacts with GPs, nurses, physiotherapists, psychologists, other healthcare consultations, diagnostic tests (x-rays, MRI scans, CT scans, blood tests), A&E attendances, hospital admissions; pharmacological treatments Intervention 2: 0.703 95% (Incremental (2–1): 0.099 Proba

(95% CI: NR; p=NR)

95% CI: NR

Probability Intervention 2 cost-effective (£20K/30K threshold): ~99%/99%

Analysis of uncertainty: Bootstrapping was undertaken to estimate uncertainty around the ICER.

Subgroup analyses were undertaken for:

- Males/females: £2422/£1461
- >60 / <60 years old: £1855/£1538
- Duration low back pain ≤3/>3 years: £1829/£1585
- RMQ scores <u>></u>4/<4: £1524/ AM+cognitive behavioural approaches dominated by AM (higher costs and lower QALYs)

Sensitivity analysis was undertaken: excluding cost outliers (above 90th percentile); excluding inverse weights in the estimation of costs and QALYs. This had very little impact on results.

Data sources

Health outcomes: QALYs were calculated using patient-level utility data collected at baseline, 3, 6 and 12 months and the area under the curve approach adjusted for

relevant baseline characteristics including utility. Missing data was imputed using multiple imputation techniques for those with at least one item response. **Quality-of-life weights:** EQ-5D, UK tariff. **Costs:** Costs were calculated using patient-level resource use data collected at baseline, 3, 6 and 12 months and were adjusted for relevant baseline characteristics including utility. Missing data was imputed using unconditional mean imputation methods if some resource use items were present. Intervention cost was based on the number of attended sessions (mean cost cognitive behavioural approaches £187). Unit costs were based on standard UK national sources.

Comments

Source of funding: NIHR HTA programme. **Limitations:** Study does not include all non-invasive treatment options. A longer time horizon may be preferable if differences seen at 1 year persist beyond this time. Within-trial analysis and so does not reflect full body of available evidence for this comparison; Lamb 2010 is 1 of 13 studies included in the clinical review for cognitive behavioural approach - although 1 of 2 compared to usual care with EQ5D data. **Other:**

Overall applicability^(a): partially applicable Overall quality^(b): potentially serious limitations

Abbreviations: 95% CI: 95% confidence interval; CUA: cost—utility analysis; EQ-5D: Euroqol 5 dimensions (scale: 0.0 [death] to 1.0 [full health], negative values mean worse than death); ICER: incremental cost-effectiveness ratio; NR: not reported; QALYs: quality-adjusted life years

- (a) Directly applicable / Partially applicable / Not applicable
- (b) Minor limitations / Potentially serious limitations / Very serious limitations

For Smeets 2009⁴⁷ please see Table 10 (Exercise) above.

I.12 Pharmacological

Table 15: Lloyd 2004³⁷

Lloyd A, Scott DA, Akehurst RL, Lurie-Luke E, Jessen G. Cost-effectiveness of low-level heat wrap therapy for low back pain. Value in Health. 2004; 7(4):413-422. (Guideline Ref ID LLOYD2004)

Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness
Economic analysis: CEA (health outcome: successful treatment - defined as a 2-point improvement in the 6 point pain NRS on at least 3 of the 4 days AND a 2-point improvement or	Population: Low back pain (without sciatica). Adults with acute uncomplicated, muscular, non-traumatic, low back pain. People with severe underlying morbidity or sciatica and other secondary	Total costs (mean per patient): Intervention 1: £34.22 Intervention 2: £36.04 Incremental (2–1): £1.84 (95% CI: NR; p=NR)	Proportion successfully treated: Intervention 1: 0.26 Intervention 2: 0.18 Incremental (2-1): -0.08 (95% CI: NR; p=NR)	ICER (Intervention 2 versus Intervention 1): Paracetamol dominates ibuprofen (lower costs and better health outcomes Analysis of uncertainty: PSA not conducted. An analysis was also undertaken with only initial drugs costs – the conclusion was essentially the same although the difference

better on the 24-point RMDQ from baseline at day 4)

Study design: Within-trial analysis (RCT – associated clinical paper Nadler 2002³⁹) with modelled post-trial extrapolation

Approach to analysis:

Patient level analysis of successful treatment and adverse events. Decision tree including three outcomes for patients: successful treatment, unsuccessful treatment or an AE. Each outcome was associated with different resource use in order to model the downstream cost implications of treatments.

Perspective: UK NHS
Time horizon: 4 days for
outcomes, cost
perspective not stated but
also short-term

Discounting: Costs: n/a;

Outcomes: n/a

causes of low back pain were excluded.

Patient characteristics:

N = 371 Mean age:

Intervention 1: 34.90 (SD:

11.29)

Intervention 2: 36.61 (SD:

10.4) Male:

Intervention 1: 43.4 Intervention 2: 40.6

Intervention 1:

Paracetamol 1000mg 4x daily for 2 days (n=113)

Intervention 2:

Ibuprofen (NSAID) 400mg 3x daily (n=106)

Note that study also included heat wrap but this comparator does not meet the guideline protocol.

Cost breakdown (initial treatment/other)
Intervention 1: £0.26

Currency & cost year: 2001/2002 UK pounds

Intervention 2: £0.28

Cost components incorporated:

Initial prescription costs (NHS price of treatment, plus dispensing charge, corrected for patient contribution; assuming nonexempt patients (76%) buy OTC and so zero cost to NHS), GP reconsultation for AE or unsuccessful treatment, referral to physiotherapy for unsuccessful treatment, paracetamol prescription costs for those not referred to physiotherapy initial treatment was unsuccessful. in cost was very small (2-1: £0.02). Sensitivity analyses were undertaken with: different definitions of success (range 2-1: 0.0 to -0.08); varying proportions of patients exempt from prescription charges (max 85%; increased difference in initial treatment costs 2-1 to £0.10).

Data sources

Health outcomes: Within trial analysis for health outcome of successfully treated patients (both analyses) and treatment-related AE rates (model only). Quality-of-life

weights: n/a. Cost sources: The proportion of patients exempt from prescription charges was stated as based on population data but not referenced; rate of reconsultation if not successful or AE was estimated (50%) but validated with UK survey data; rate of referral to physiotherapy was estimated (18%) and validated using NHS data: unit costs from standard UK national sources.

Comments

Source of funding: Proctor & Gamble Health Sciences Limited (manufacturers of the heat wrap in the study). Limitations: Study does not include all non-invasive treatment options; resource use data (pre-1999) and unit costs (2001/2) may not reflect current NHS context. QALYs were not used as the health outcome measure. Modelled extrapolation of within-trial analysis and so does not reflect full body of available evidence: 1 of 1 study identified in clinical review directly comparing ibuprofen and paracetamol (although no protocol outcomes available); however, a number of placebo controlled studies are available for ibuprofen and paracetamol and so indirect evidence is available that is not incorporated. Downstream resource use rates based on estimates, although validated with UK data. PSA was not undertaken. Other:

Overall applicability(a): Partially applicable Overall quality(b): Potentially serious limitations

Abbreviations: CEA: cost-effectiveness analysis; 95% CI: 95% confidence interval; da: deterministic analysis; ICER: incremental cost-effectiveness ratio; NR: not reported; NRS = numerical rating scale; QALYs: quality-adjusted life years

- (a) Directly applicable/Partially applicable/Not applicable
- (b) Minor limitations/Potentially serious limitations/Very serious limitations

Table 16: Morera-Dominguez 2010³⁸

Morera-Dominguez C, Ceberio-Balda F, Florez G, Masramon X, Lopez-Gomez V. A cost-consequence analysis of pregabalin versus usual care in the symptomatic treatment of refractory low back pain: sub-analysis of observational trial data from orthopaedic surgery and rehabilitation clinics. Clinical Drug Investigation. 2010; 30(8):517-531 (Guideline Ref ID MORERADOMINGUEZZ010)

Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness
Economic analysis: CCA (various health outcomes)	Population: Adults with low back pain due to radiculopathy	Total costs (mean change from baseline per patient): Intervention 1: £41	From clinical review (2 vs. 1): • Pain (BPI): MD -1.40 (CI:	ICER (Intervention 2 versus Intervention 1): n/a
Study design: within-trial analysis (cohort study – associated clinical paper Morera-Dominguez 2010 ³⁸ Approach to analysis: Analysis of individual level data for health outcomes	(sciatica) (>6 months) refractory to at least one course of previous analgesics Patient characteristics N = 683 Mean age: 55.0 years (SD:	Intervention 2: -£26 Incremental (2-1): -£68 (95% CI: -£280 to £145; p≤0.540) Cost breakdown — incremental (2-1):	-1.81, -0.99) • Quality of life (SF-12 physical summary score): MD 3.90 (CI: 2.21, 5.59) • Quality of life (SF-12 mental summary score):	Analysis of uncertainty:

and resource use. Unit 12.7) MD 5.30 (CI: 3.71, 6.89) Pharma treatment: £236 costs applied. Male: 50.5% Psychological distress Non-pharma treatment: (HADS - anxiety): MD --£94 1.80 (CI: -2.42, -1.18) Perspective: Spain direct Intervention 1: Care not Medical visits and hospital medical costs (societal Psychological distress including pregabalin admissions: -£243 also analysed but not (HADS - depression): MD Complementary tests: £34 presented here) -1.90 (CI: -2.58, -1.22) **Intervention 2:** Care Follow-up: 12 weeks including pregabalin (mean Currency & cost year: **Discounting:** Costs: n/a; dose 189.9 mg/day, SD 2007 Spanish Euros Outcomes: n/a 141.7) (gabapentinoid (presented here as 2007 UK anticonvulsant) pounds(a)) **Cost components** incorporated: Pharmacological treatment, non-pharmacological treatment, medical visits and hospital admissions and complementary tests (e.g. CT and MRI). Does not include any cost of adverse events of drugs.

Data sources

Health outcomes: Within-RCT analysis. **Quality-of-life weights:** n/a **Cost sources:** Costs were calculated using patient-level resource use data collected at baseline and 12 weeks. Unit costs were based on Spanish list prices for drugs and a healthcare cost database for other resource items.

Comments

Source of funding: Pfizer (manufacturer of pregabalin). **Limitations:** Spanish resource use data (2006-7) and unit costs (2007) may not reflect current NHS context. QALYs were not used as the health outcome measure. Study does not include all non-invasive treatment options. Analysis is based on a cohort study. Within-trial analysis and so does not reflect full body of available evidence for this comparison; Morera-Dominguez is 1 of 2 studies included in the clinical review for gabapentinoid anticonvulsants; 1 cohort and 1 RCT. No exploration of uncertainty. The analysis was funded by the manufacturer of pregabalin. **Other:** In the arm without pregabalin use of gabapentin was significantly higher.

Overall applicability(b): partially applicable Overall quality(c): potentially serious limitations

Abbreviations: BPI: brief pain index, 0-100; CCA: cost—consequence analysis; 95% CI: 95% confidence interval; HADS: hospital anxiety and depression scale, 0-21; ICER: incremental cost-effectiveness ratio; MD = mean difference; NR: not reported; QALYs: quality-adjusted life years; SF-12: short-form 12, 0-100

- (a) Converted using 2007 purchasing power parities⁴²
- (b) Directly applicable/Partially applicable/Not applicable
- (c) Minor limitations/Potentially serious limitations/Very serious limitations

Table 17: Wielage 201368

Wielage RC, Bansal M, Andrews JS, Wohlreich MM, Klein RW, Happich M. The cost-effectiveness of duloxetine in chronic low back pain: a US private payer perspective. Value in Health. 2013; 16(2):334-344. (Guideline Ref ID WIELAGE2013)

Study details	Population & interventions	Costs	Health outcomes	Cost	effectivene	ss			
Economic analysis: CUA (health outcome: QALYs) Study design: Probabilistic decision analytic model Approach to analysis: Markov model based on NICE Osteoarthitis (OA) 2008 clinical guideline. Health states include treatment, death and 12 states associated with persistent adverse events	interventions Population: Chronic low back pain (with or without sciatica), >3 months, post first line treatment with paracetamol Cohort settings: Start age: NR Male: NR Intervention 1: Duloxetine (SNRI), 60-	Total costs (mean per patient): Intervention 1: £35,920 Intervention 2: £35,213 Intervention 3: £34,989 Intervention 4: £35,842 Intervention 5: £36,188 Intervention 6: £36,876 Intervention 7: £38,090 Intervention 8: £35,758 For incremental analysis see cost effectiveness	Health outcomes QALYs (mean per patient): Intervention 1: 12.2123 Intervention 2: 12.1887 Intervention 3: 12.1899 Intervention 4: 12.1884 Intervention 5: 12.1973 Intervention 6: 12.1974 Intervention 7: 12.2029 Intervention 8: 12.2043 For incremental analysis see cost effectiveness	ICER	(Interventi	2 versus analysis(c) QALY 12.1884 12.1887 12.1899 12.1973 12.1974 12.2029 12.2043	Inc cost Domina Domina Baselin Domina Domina	Inc QALY ated by 2 ated by 3	ICER
(symptomatic ulcer, complicated GI bleed, myocardial infarction, stroke, heart failure and fracture). Proton-pump inhibitor usage and transient adverse events (dyspepsia, nausea, diarrhoea, constipation, insomnia, pruritus, vomiting, dizziness, somnolence and opioid abuse) were included in	120mg Intervention 2: Celecoxib (NSAID), 200mg once daily Intervention 3: Naproxen (NSAID), 500mg twice daily Intervention 4: Pregabalin (gabapentinoid anticonvulsant), 300mg twice daily	Currency & cost year: 2011 USA dollars (presented here as 2011 UK pounds(b)) Cost components incorporated: Drug costs and medical utilisation for management of adverse events, titration and	see cost effectiveness column	8 £3 1 £3 PSA not r For pairw (~£20K/3 Intervent Intervent Probabili Other con	£35,920 not reported airwise ana 0K/30K three vention 1 volume 1 v	12.2123 d for full inc lyses, prob	£931 crementa ability co /10%(e) %/95% r 5: 99.99 orted.	0.022 4 al analysis sst-effecti	£41,5 21 5.

Perspective: USA healthcare payer perspective

Time horizon: Lifetime

Treatment effect duration(a): Same as
treatment duration (see
intervention description).

Discounting: Costs: 3%;

Outcomes: 3%

Intervention 5:

Oxycodone/acetaminop hen (opioid/paracetamol), 7.5/325-15/650mg every 6 hours

Intervention 6:

Oxycodone extended release (opioid), 10-30mg twice daily

Intervention 7:

Tapentadol extended release (opioid), 300-600mg once daily

Intervention 8:

Tramadol immediate release (opioid), 200-300mg once daily.

Duration of treatment was the lesser of: 1 year, until discontinuation or until

occurrence of a

persistent AE.

discontinuation.

analyses conducted for duloxetine versus naproxen. When the probabilities of CV adverse events associated with NSAIDs were increased or when the start age in the model was increased to 65 years, duloxetine was cost effective compared to naproxen at £20,000 per QALY.

Probabilistic sensitivity analysis for duloxetine versus naproxen, duloxetine versus tramadol and duloxetine versus oxycodone/acetaminophen.

Data sources

Health outcomes: AE rates from OA 2008 NICE guideline and published literature (meta-analysis), with exception of duloxetine which was from chronic low back pain RCTs. Expert opinion used for small number of inputs (e.g. PPI usage). Discontinuation rates for initial 3 months taken from low back pain RCTs for duloxetine; OA RCTs for NSAIDs and opioids; neuropathic pain RCTs for pregabalin. Discontinuation for subsequent 3 months based on expert opinion. Age-dependent mortality taken from USA life tables and persistent AE-related mortality from published literature. Quality-of-life weights: Systematic review of pain scores from chronic low back pain RCTs conducted. Pain scores converted to EQ-5D (USA preference weight) using 'a transfer to utility' regression equation. Patient level data from three Eli Lilly sponsored trials of duloxetine versus placebo in low back pain used in this analysis to build regression and for validation. No trials reporting drug efficacy (pain scores) were identified for celecoxib, pregabalin, tramadol, oxycodone/acetaminophen. Celecoxib and naproxen assumed to have same efficacy as pooled efficacy of etoricoxib and naproxen, equivalent efficacies were assumed for tramadol and tramadol/acetaminophen, and for oxycodone/ acetaminophen and oxycodone. Pregabalin was

assumed to have same efficacy as placebo effect seen in placebo arms of the other RCTs. Population utility weights for age and sex from USA national source and for adverse events taken from literature (unclear if these utilities are EQ-5D). **Cost sources:** Drug costs from average 2011 wholesale USA prices, discounted at 16% to reflect actual acquisition prices. For titration and discontinuation-related medical costs Medicare reimbursement rates were used, adjusted by using a Medicare/private payer ratio. Published literature costs used for AE-related medical costs (inflated to 2011 USA dollars). Resource use from published data and expert opinion.

Comments

Source of funding: Eli Lilly and Company (manufacturer of duloxetine). Limitations: Study does not include all non-invasive treatment options. USA unit costs from 2011 and resource use from various time points may not reflect current NHS context. Utilities obtained by converting pain scores to EQ-5D with a US preference weight, other utilities were included in the model and methods were unclear. Costs and health effects were discounted at a non-reference case rate (3%), although similar. Important outcomes may not be captured by model. Adverse events included were symptomatic ulcer, complicated GI bleed, myocardial infarction, stroke, heart failure, fracture, dyspepsia, nausea, diarrhoea, constipation, insomnia, pruritus, vomiting, dizziness, somnolence and opioid abuse adverse events omitted were renal failure, opioid misuse related mortality, bleeding, hepatotoxicity and suicidality. Full effect of treatment may not be captured as a result of mapping pain scores only (e.g. impact of disability and mental distress). Relative treatment effects for QoL were based on a meta-analysis: Skljarevski 2009, 2010A and 2010B are 3 of 10 studies comparing antidepressants to placebo; Pallay 2004 and Birbara 2003 are 2 of 6 studies comparing NSAIDs to placebo; Peloso 2004 is 1 of 4 studies comparing opioid combinations to placebo; Buynak 2009, Ruoff 2003 and Webster 2006 are 3 of 9 studies comparing opioids to placebo. Four studies were used in the model, which were excluded from the clinical review (Skljarevski 2010C, Binsfield 2010, Wild 2010, Hale 2009). AE rates for all comparators with the exception of duloxetine were from a different patient population; efficacy data for five of the comparators were based on assumptions: celecoxib and naproxen assumed to have same efficacy as placebo effect seen in placebo arms of the other RCTs. Discontinuation rates in subsequent 3 months based on expert opinion. PSA results were not reported for the full incremental analysis. Study funded by Eli Lilly (manufacturer of duloxetine).

Overall applicability^(f): Partial applicability Overall quality^(g): Potentially serious limitations

Abbreviations: AE: adverse event; CUA: cost—utility analysis; CV: cardiovascular; EQ-5D: Euroqol 5 dimensions (scale: 0.0 [death] to 1.0 [full health], negative values mean worse than death); ICER: incremental cost-effectiveness ratio; NR: not reported; pa: probabilistic analysis; NSAID: non-steroidal anti-inflammatories; OA: osteoarthritis; SNRI: serotonin—norepinephrine reuptake inhibitors; QALYs: quality-adjusted life years.

- (a) For studies where the time horizon is longer than the treatment duration, an assumption needs to be made about the continuation of the study effect. For example, does a difference in utility between groups during treatment continue beyond the end of treatment and if so for how long.
- (b) Converted using 2011 purchasing power parities⁴²
- (c) Intervention number in order of least to most effective in terms of QALYs $\,$
- (d) Full incremental analysis of available strategies: first strategies are ruled out that are dominated (another strategy is more effective and has lower costs) or subject to extended dominance (the strategy is more effective and more costly but the incremental cost effectiveness ratio is higher than the next most effective option and so it would never be the most cost effective option); incremental costs, incremental effects and incremental cost effectiveness ratios are calculated for the remaining strategies by comparing each to the next most effective option
- (e) Estimated from graph
- (f) Directly applicable/Partially applicable/Not applicable
- (g) Minor limitations/Potentially serious limitations/Very serious limitations

I.13 MBR

For Critchley 2007⁹ please see Table 8 (Exercise) above.

For Smeets 2009⁴⁷ please see Table 10 (Exercise) above.

I.14 Return to work

For return to work interventions both an NHS and an employer perspective were considered relevant on the basis that potentially employers could provide such interventions – information relevant to both perspectives is therefore included in evidence tables for this intervention. Note that applicability and methodological quality assessment relate to the NHS perspective and NHS decision making only.

Table 18: Hlobil 2007²²

Hlobil H, Uegaki K, Staal JB, Bruyne M, Smid T, Mechelen W. Substantial sick-leave costs savings due to a graded activity intervention for workers with non-specific sub-acute low back pain. Eur Spine J.: Springer-Verlag. 2007; 16(7):919-924. (Guideline Ref ID HLOBIL2007)

Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness
Economic analysis: CCA (clinical outcomes reported in separate paper ⁵¹) Study design: Within-trial analysis (RCT – associated clinical paper Staal 2004 ⁵¹) Approach to analysis: Analysis of individual level data for resource use (and sick leave days) and clinical outcomes. Unit costs applied. Perspective: Direct healthcare costs (productivity costs also	Population: Sick listed employees who had low back pain for a minimum of 4 weeks without sciatica. Patient characteristics N = 134 Mean age: 38 years (SD: NR) Male: 94% Intervention 1: Usual care from GP and guidance from occupational physician. Not allowed to attend physiotherapy practice where intervention group were treated.	Total healthcare costs 12 months (mean per patient): Intervention 1: £515 Intervention 2: £576 Incremental (2–1): saves £60 (95% CI: -£336 to £181; p=NR) Cost breakdown (initial treatment/other) Intervention 1: £0/£515 Intervention 2: £342/£234 Total lost productivity costs 3 years (mean per patient): Gross lost productivity days (total days workers were completely or partially sick listed) Incremental (2–1): £5455	See clinical review Staal2004	ICER (Intervention 2 versus Intervention 1): n/a Analysis of uncertainty: Net productivity loss was reestimated assuming 25%/50% decreased work performance. Results for year 1 went from £719 to £1197 and £1674

reported). Follow-up: 1 year (healthcare costs) / 3 years (productivity costs) Discounting: Costs: none; Outcomes: none Discounting: Costs: none; Outcomes: none Intervention 2: Graded activity, a physical exprogramme based on operation conditioning behavioural principles. Physiotherapist. The hour sessions per week. Edu Exercises (aerobic, abdomination and leg) and individually tailed exercises to simulate and proproblematic tasks at work or gradually increased. Return to plan.	accounting for partial lost days) Incremental (2–1): £1195 (95% CI: -£2989 to £4974; p=NR) Currency & cost year: 1999 Netherlands Euros (presented here as 1999 UK pounds(a))	respectively. Other results not reported.
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Data sources

Health outcomes: Within-trial analysis (reported separately in Staal 2004). Quality-of-life weights: n/a Cost sources: Health care costs were calculated using patient-level resource use data collected in 3 cost diaries over the first 12 months with missing data imputed. Intervention cost was based on the number of attended sessions (mean intervention cost £342). Unit costs were based on Dutch national sources.

Comments

Source of funding: Dutch Health Insurance Executive Council. **Limitations:** Dutch resource use data (1999-2002) and unit costs (1999) may not reflect current NHS context. QALYs were not used as the health outcome measure. Within-trial analysis and so does not reflect full body of available evidence for this comparison. Staal 2004 is 1 of 8 studies included in the clinical review for return to work interventions. Limited sensitivity analyses were undertaken. **Other:**

Overall applicability(a): partially applicable Overall quality(b): potentially serious limitations

Abbreviations: CCA: cost—consequence analysis; 95% CI: 95% confidence interval; NR: not reported; pa: probabilistic analysis; QALYs: quality-adjusted life years

- (a) Converted using 1999 purchasing power parities⁴²
- (b) Directly applicable / Partially applicable / Not applicable
- (c) Minor limitations / Potentially serious limitations / Very serious limitations

Table 19: Lambeek 2010³³

Lambeek LC, Bosmans JE, van Royen BJ, van Tulder MW, van MW, Anema JR. Effect of integrated care for sick listed patients with chronic low back pain: economic evaluation alongside a randomised controlled trial. British Medical Journal. 2010; 341:c6414. (Guideline Ref ID LAMBEEK2010)

Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness
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2016

Economic analysis:

CUA (health outcome: QALYs)

Study design: Within-trial analysis (RCT – associated clinical paper Lambeek2010A³⁴)

Approach to analysis:

Analysis of individual level data for EQ-5D and resource use (and sick leave days). Unit costs applied.

Perspective: Dutch NHS (productivity costs also reported; informal care costs also reported but not reported here).

Follow-up: 12 months
Discounting: Costs: n/a;

Outcomes: n/a

Population:

Adults 18-65 years with low back pain lasting more than 12 weeks (with/without sciatica), had paid work and were on (partial) sick leave.

Patient characteristics

N = 134

Mean age: 46 years (SD: NR)

Male: 58%

Intervention 1:

Usual care. Delivered by occupational therapist and/or GP according to the Dutch guidelines for low back pain. (n=68)

Intervention 2:

Integrated care. Workplace intervention protocol based on ergonomics and a graded activity protocol with an aim to restore occupational functioning, delivered by a team of a medical specialist, occupational therapist, physiotherapist and clinical occupational physician. (n=66)

Total healthcare costs (mean per patient):

Intervention 1: £1104 Intervention 2: £1375 Incremental (2-1): £271 (95% CI: NR; p=NR)

Cost breakdown (initial treatment/other)

Intervention 1: £0/£1104 Intervention 2: £1077/£298

Total lost productivity costs 3 years (mean per patient):

Intervention 1: £17,213 Intervention 2: £11,686 Incremental (2–1): -£5527

(95% CI: -£10,042 to -£740; p=NR)

Currency & cost year:

2007 Dutch Euros (reported as 2007 UK pounds(a)).

Cost components incorporated:

GP, physiotherapist, occupational physician, manual therapy, psychologist, clinical occupational physician, diagnostic tests, hospital stay, medical specialist.

QALYs (mean per patient):

Intervention 1: 0.65 Intervention 2: 0.74 Incremental (2–1): 0.09 (95% CI: 0.01 to 0.16; p=NR)

Absenteeism from work (mean days per patient):

Intervention 1: 130.4 Intervention 2: 88.5 Incremental (2–1): -41.9 (95% CI: NR; p=NR)

ICER (Intervention 2 versus Intervention 1):

£3011 per QALY gained (da) 95% CI: NR Probability Intervention 2 cost-effective (£20K/30K threshold): NR for healthcare costs only

Analysis of uncertainty:

perspective.

Uncertainty was quantified for the full analysis but not for the healthcare costs only perspective.

A series of alternative analyses were also undertaken but again only from the aggregated cost perspective.

Data sources

Health outcomes: QALYs were calculated using patient-level utility data and the area under the curve approach. EQ-5D was administered to patients at four time points. **Quality-of-life weights:** EQ5D, Dutch tariff (TTO). **Cost sources:** Resource use captured from patient cost questionnaires at 3, 6, 9, 12 months. Unit costs were from Dutch national sources. Integrated care costs were constructed through a bottom-up approach (£1077).

Comments

Source of funding: funded by VU University medical centre, TNO work and employment, Dutch health insurance executive council, Stichting Instituut GAK, and the Netherlands organisation and development R&D **Limitations:** Dutch resource use data (2005-2009) and unit costs (2009) may not reflect current NHS context. Dutch EQ5D tariff used (time-trade off method). Within-trial analysis and so does not reflect full body of available evidence for this comparison. Lambeek2010A is 1 of 8 studies included in the clinical review for return to work interventions. Although uncertainty was explored in the analysis, no sensitivity analyses were available for the healthcare perspective relevant to the guideline. **Other:**

Overall applicability(b): partially applicable Overall quality(c): potentially serious limitations

Abbreviations: 95% CI: 95% confidence interval; CUA: cost—utility analysis; da: deterministic analysis; EQ-5D: Euroqol 5 dimensions (scale: 0.0 [death] to 1.0 [full health], negative values mean worse than death); ICER: incremental cost-effectiveness ratio; NR: not reported; QALYs: quality-adjusted life years; TTO: time-trade off

- (a) Converted by authors using 2007 purchasing power parities
- (b) Directly applicable / Partially applicable / Not applicable
- (c) Minor limitations / Potentially serious limitations / Very serious limitations

Table 20: Steenstra 2006^{52,53}

Steenstra IA, Anema JR FAU - van Tulder M, van Tulder MW FAU - Bongers P, Bongers PM FAU - de Vet H, de Vet HC FAU - van Mechelen W, van MW. Economic evaluation of a multi-stage return to work program for workers on sick-leave due to low back pain. Journal of Occupational Rehabilitation. 2006; 16(4):557-578. (Guideline Ref ID STREENSTRA2006A)

Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness
Economic analysis: CUA (health outcome: QALYs) Study design: Within-trial analysis (RCT – associated clinical paper Anema2007¹). Approach to analysis: Analysis of individual level data for EQ-5D and resource use (and sick leave days). Unit costs applied. Perspective: Dutch NHS (costs of lost paid work	Population: Workers with low back pain on sick leave from regular work for 2-6 weeks, 18-65 years. With/without sciatica. Patient characteristics N = 196 Mean age: 42 years (SD: NR) Male: 66% Intervention 1: Usual care. Recommendation to take sick-leave, resuming daily activities and work within two	Total costs (mean per patient): Intervention 1: £1,314 Intervention 2: £1,541 Incremental (2–1): £228 (95% CI: -£116 to £557; p=NR) Cost breakdown of intervention/other costs not reported. Total lost productivity costs (mean per patient):	QALYs (mean per patient): Intervention 1: 0.26 Intervention 2: 0.21 Incremental (2–1): -0.04 (95% CI: -0.12 to 0.04; p=NR)	ICER (Intervention 2 versus Intervention 1): Intervention 2 dominated by intervention 1 Analysis of uncertainty: Uncertainty was quantified using bootstrapping for some analyses but not for the healthcare costs only perspective. Three sensitivity analyses around the calculation of indirect costs were undertaken. Relevant numerical results were not reported.

days also reported; costs of lost unpaid work days and indirect healthcare costs also reported but not reported here).

Follow-up 12 months

Discounting: Costs: n/a; Outcomes: n/a

weeks, supervised by GP Intervention 2:

Usual care plus multidisciplinary programme with a return to work focus (individual workplace intervention). Workplace assessment with work modifications (involving ergonomist or occupational health nurse), co-ordination between occupational physician and worker's GP.

Note, this study has 2 randomisation stages; first randomisation occurred at 2 weeks for all recruited participants into the two intervention groups, second randomisation was at 8 weeks for only those people who were still off work due to their back pain. In this second randomisation they were rerandomised to either graded activity or usual care. Only the first randomisation is presented here.

Intervention 1: £3,879 Intervention 2: £3,413 Incremental (2–1): saves £467 (95% CI: -£1,381 to £495; p=NR)

Currency & cost year:

2002 (assumed cost year as not reported) Netherlands Euros (presented here as 2002 UK pounds(a)]

Cost components incorporated:

Direct healthcare costs: intervention costs, additional healthcare visits (GP, manual therapist, physiotherapist, medical specialist, other healthcare professionals), prescription medication, professional home care and hospitalisation. Productivity costs: days lost of paid work.

Data sources

Health outcomes: Health outcome questionnaires administered at baseline, 3, 6, 12 months, missing data was imputed. However it appears that the CUA is calculated using the mean difference in change in EQ-5D from baseline to 12 months rather than estimating QALYs taking into account the time spent at different utility levels. Quality-of-life weights: EQ5D, UK tariff. Cost sources: Analysis of individual-level resource use captured through questionnaires administered at 3, 6 and 12 months, missing data was imputed. Unit costs sources were the Dutch NHS prices based on Dutch guidelines, Dutch society of pharmacy and market prices (for graded activity). Other:

Comments

Source of funding: The Netherlands Organisation for Health Research and Development Limitations: Dutch resource use (2000-2003) and unit cost (year not stated) data may not reflect current NHS context. The CUA ICER is calculated as the difference in EQ5D utility between baseline and last follow-up rather than using the time spent at different EQ5D levels to calculate QALYs. There is a significant difference in baseline EQ5D between two of the arms. Within-trial analysis and so does not reflect full body of available evidence for this comparison; Anema2007 is 1 of 8 studies included in the clinical review for return to work interventions. Limited sensitivity analyses.

Overall applicability(b): partially applicable Overall quality(c): potentially serious limitations

Abbreviations: CUA: cost—utility analysis; da: deterministic analysis; EQ-5D: Euroqol 5 dimensions (scale: 0.0 [death] to 1.0 [full health], negative values mean worse than death); ICER: incremental cost-effectiveness ratio; NR: not reported; pa: probabilistic analysis; QALYs: quality-adjusted life years

- (a) Converted using 2002 purchasing power parities⁴²
- (b) Directly applicable / Partially applicable / Not applicable
- (c) Minor limitations / Potentially serious limitations / Very serious limitations

I.15 Spinal Injections

None.

I.16 Radiofrequency ablation

Table 21: van Wijk 2005⁶³

van Wijk RMAW, Geurts JWM, Wynne HJ, Hammink E, Buskens E, Lousberg R et al. Radiofrequency denervation of lumbar facet joints in the treatment of chronic low back pain: a randomized, double-blind, sham lesion-controlled trial. Clinical Journal of Pain. 2005; 21(4):335-344. (Guideline Ref ID VANWIJK2005)

Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness
Economic analysis: CCA (health outcomes: SF-36, VAS-back, global perceived effect on back pain, analgesic intake) Study design: RCT (within trial analysis)	Population: >17 year olds with low back pain, with/without sciatica, > 6 months with focal tenderness over the facet joints	Total costs (mean per patient): Intervention 1: £68 Intervention 2: £254 Incremental (2–1): £186 (95% CI: NR; p=NR)	See clinical review van Wijk 2005 (SF-36, VAS-back, global perceived effect on back pain, analgesic intake).	ICER (Intervention 2 versus Intervention 1): n/a Analysis of uncertainty: No sensitivity analysis conducted.
Approach to analysis: Health outcome and	Cohort settings: n: 81	Cost breakdown (mean per patient):		

resource collated through diaries and questionnaires administered prior to treatment and at 3 months. 1 year data for health outcomes was supposed to be reported by the study, however at this time-point most patients were un-blinded and there was loss-to follow-up. Dutch unit costs applied.

Perspective: Netherlands healthcare payer perspective

Follow-up: 3 months **Discounting:** Costs: n/a;

Outcomes: n/a

Start age: 48 Male: 28%

Intervention 1: (n=41)

Sham lesion

Intervention 2: (n=40) Radiofrequency lesion (80°C lesion for 60 seconds, lesion made on 1 or both sides).

Both groups given intraarticular joint injection prior to radiofrequency ablation. Responders were randomised. Intervention cost
Intervention 1: £0
Intervention 2: £197

Medical consumption over 3

months:

Intervention 1: £68 Intervention 2: £57

Currency & cost year:

Year NR assumed 2003 Euros (presented here as 2003 UK pounds^(a))

Cost components incorporated:

Intervention costs (including staff time, materials, overheads, administration, accommodation and day care facilities)
Additional medical consumption over 3 month follow-up (medical, paramedical, and pharmaceutical treatment).

Data sources

Health outcomes: Within-trial analysis (same paper). Health outcome collated through diaries and questionnaires administered prior to treatment and at 3, 6, 9 and 12 months. Data beyond 3 months not reported for all outcomes as at these later time points most patients were un-blinded and there was loss-to follow-up. Quality-of-life weights: n/a. Cost sources: Resource use for interventions recorded by trial investigators, other resource use captured from patient questionnaires. Source of unit costs not reported. Study reported the cost of sham lesion to be equal to radiofrequency ablation. Including the cost of a sham was deemed inappropriate and was excluded here.

Comments

Source of funding: Dutch Health Insurance Council and Pain Expertise Center, The Netherlands. Limitations: Dutch resource use data (1996-1999) and unit costs (year not reported, assumed to be 2003) may not reflect current NHS context. QALYs were not used as the health outcome measure (SF-36 reported, however QALYs were

not calculated). A longer time horizon may be preferable if effects may persist beyond 3 months. Within-trial analysis and so does not reflect full body of available evidence for this comparison; van Wijk 2005 is 1 of 7 studies included in the clinical review for radiofrequency ablation versus placebo sham. No sensitivity analyses undertaken. Source of unit costs unclear. **Other:** n/a

Overall applicability(b): Partially applicable Overall quality(c): Potentially serious limitations

Abbreviations: CCA: cost-consequence analysis; 95% CI: 95% confidence interval; ICER: incremental cost-effectiveness ratio; NR: not reported; QALYs: quality-adjusted life years

Price C, Arden N, Coglan L, Rogers P. Cost-effectiveness and safety of epidural steroids in the management of sciatica. Health Technology Assessment. United

- (a) Converted using 2003 purchasing power parities⁴²
- (b) Directly applicable / Partially applicable / Not applicable
- (c) Minor limitations / Potentially serious limitations / Very serious limitations

I.17 Epidurals

Table 22: Price 2005^{44,44}

Kingdom 2005; 9(33):iii, 1-ii	Kingdom 2005; 9(33):iii, 1-iii,58. (Guideline Ref ID PRICE2005)							
Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness				
Economic analysis: CUA (health outcome: QALYs) Study design: Within-trial analysis (associated clinical paper Arden 2005) Approach to analysis: Analysis of individual level data of SF-36 data (converted to SF-6D utility) at baseline, 3, 6, 12, 26 and 52 weeks. QALYs constructed through area under the curve method. Resource use captured from within trial and unit costs applied.	Population: Adults with low back pain and sciatica (unclear spinal pathology). Cohort settings: Start age: 43 Male: 47% Intervention 1: (n=108) Placebo (injection of 2ml of normal saline into the interspinous ligament) Intervention 2: (n=120) Steroid plus local anaesthetic epidural, non-	Total costs (mean per patient): Intervention 1: £0 Intervention 2: £265 Incremental (2–1): £265 (95% CI: NR; p=NR) Currency & cost year: 2002-2003 UK pounds Cost components incorporated: For those receiving intervention 2 only: assessment and review by clinician, medical and nursing time incurred during procedure, nursing time on	QALYs (mean per patient): Intervention 1: NR Intervention 2: NR Incremental (2–1): 0.0059350 (95% CI: NR; p=NR)	ICER (Intervention 2 versus Intervention 1): £44,701 per QALY gained (da) 95% CI: NR Analysis of uncertainty: No bootstrapping undertaken. A sensitivity analysis was conducted where the costs were adjusted assuming only one epidural injection was administered and the impact on QALYs is assumed to be unchanged. ICER = £25,746. Additional sensitivity analyses were undertaken, where the maximum healthcare professional resource use reported in the trial were used to estimate intervention costs and where the patient is assumed to require an overnight stay. In both cases this				

Perspective: UK NHS Follow-up: 1 year Discounting: Costs: n/a; Outcomes: n/a	image guided (lumbar epidural injection of 80mg triamcinolone acetonide and 10ml of 0.125% bupivacaine)	recovery post-procedure, drug and equipment use associated with procedure and pathology and radiology use.	increased the total cost of intervention 2 and therefore the ICER.
	All participants received a standard physiotherapy package prior (education and exercise) and analgesia as required. Injections were repeated at 3 and 6 weeks in relation to response. The indication for repeat injection was less than a 75% improvement in Oswestry Disability Questionnaire from the baseline visit.		

Data sources

Health outcomes: QALYs were calculated using patient-level SF-36 data, converted to SF-6D utility, collected at baseline, 3, 6, 12, 26 and 52 weeks. At 12 weeks the average scores converged for intervention 1 and 2. The area under the curve approach was used to calculate incremental QALYs. Quality-of-life weights: SF-6D, tariff used unclear. Cost sources: Resource use for interventions as reported by clinicians. Unit costs from NHS trusts finance departments and UK national published sources. No costs were collected for the placebo arm. Usual care cost not included as it was received by both groups and assumed to be the same.

Comments

Source of funding: NHS R&D HTA Programme. Limitations: UK resource use data (1999-2002) and unit costs (2002/3) may not reflect current NHS context. Non-NICE reference case utility measure used to estimate QALYs (SF-6D), unclear if UK population valuations were used. Within-trial analysis and so does not reflect full body of available evidence for this comparison; Arden 2005 is 1 of 2 studies included in the clinical review for steroid epidurals + local anaesthetic versus placebo (non-image guided). Limited sensitivity analyses undertaken. Other: None

Overall applicability^(a): Partially applicable Overall quality^(b): Potentially serious limitations

Abbreviations: 95% CI: 95% confidence interval; CUA: cost—utility analysis; da: deterministic analysis; ICER: incremental cost-effectiveness ratio; NR: not reported; QALYs: quality-adjusted life years; SF-6D: Short form 6 dimensions (scale: 0.0 [death] to 1.0 [full health]; SF-36: Short form 36 — quality of life questionnaire

- (a) Directly applicable / Partially applicable / Not applicable
- $(b) \ \ \textit{Minor limitations / Potentially serious limitations / Very serious limitations}$

Table 23: Spijker-Huiges 2014⁵⁰

Spijker-Huiges A, Vermeulen K, Winters JC, van WM, van der Meer K. Costs and cost-effectiveness of epidural steroids for acute lumbosacral radicular syndrome in general practice: an economic evaluation alongside a pragmatic randomized control trial. Spine. 2014; 39(24):2007-2012. (Guideline Ref ID SPIJKER2014)

Study details	Population & interventions	Costs (d)	Health outcomes	Cost effectiveness
Economic analysis: CEA (health outcome: 1 point improvement in NRS back pain score) Study design: Within-trial analysis (RCT, associated clinical paper Spijker- Huiges 2014A) Approach to analysis: Analysis of individual level data for health outcomes and resource use (based on patient questionnaire) collected at baseline, 2, 4, 6, 13, 26 and 52 weeks. Unit costs applied. Perspective: Dutch health care provider (societal costs analysed but not presented here) Follow-up: 1 year Discounting: Costs: n/a; Outcomes: n/a	Population: Adults with sciatica (unclear spinal pathology). Cohort settings: Start age: 44 Male: 45% Intervention 1: (n=33) Usual care provided by GP (pain treatment with analgesics, advice to maintain normal activities and referral if necessary) Intervention 2: (n=30) Steroid epidural, non-image guided (segmental epidural injection of 80mg of triamcinolone in normal saline)	Total costs (mean per patient): Intervention 1: £1,042 Intervention 2: £1,100 Incremental (2–1): £58 (95% CI: NR; p=NR) Currency & cost year: Year unclear, assumed to be 2007 Euros (presented here as 2007 UK pounds(a)) Cost components incorporated: Intervention cost (for intervention 2 only), GP care, hospital care, additional examinations, medication, physiotherapy, alternative therapies and home help visits.	NRS back pain score (mean change per patient): Intervention 1: NR Intervention 2: NR Incremental (2–1): 0.97	ICER (Intervention 2 versus Intervention 1): £60 per 1 point improvement in NRS back pain (da) 95% CI: NR Analysis of uncertainty: Bootstrapping undertaken but only from a societal perspective which is not presented here. No other sensitivity analyses were conducted.

Data sources

Health outcomes: Within-trial analysis (RCT, associated clinical paper Spijker-Huiges 2014A) measurements at baseline, 2, 4, 6, 13, 26 and 52 weeks. Mean change in NRS back pain score calculated from point estimate for the ICER reported in the study. Quality-of-life weights: n/a. Cost sources: Resource use from questionnaires

completed by participants. Unit costs sourced from Dutch guidelines for costs and Dutch national medication costs.

Comments

Source of funding: Department of General Practice, University Medical Center Groningen, Netherlands. Limitations: Dutch resource use data (2005-2007) and unit costs (date unclear) may not reflect current NHS context. QALYs were not used as the health outcome measure. Within-trial analysis and so does not reflect full body of available evidence for this comparison. No sensitivity analyses undertaken. Other: None

Overall applicability(b): Partially applicable Overall quality(c): Potentially serious limitations

Abbreviations: CEA: cost-effectiveness analysis; 95% CI: 95% confidence interval; da: deterministic analysis; EQ-5D: Eurogol 5 dimensions (scale: 0.0 [death] to 1.0 [full health], negative values mean worse than death); ICER: incremental cost-effectiveness ratio; NR: not reported; NRS: numerical rating scale; QALYs: quality-adjusted life years

- (a) Converted using 2007 purchasing power parities⁴²
- (b) Directly applicable / Partially applicable / Not applicable
- (c) Minor limitations / Potentially serious limitations / Very serious limitations
- (d) Original analysis adopted a societal perspective, costs presented here were re-estimated to reflect NHS perspective only

I.18 Surgery and prognostic factors

None.

1.19 **Spinal decompression**

Table 24: Tosteson 2008⁵⁹

Tosteson ANA, Skinner JS, Tosteson TD, Lurie JD, Andersson GB, Berven S et al. The cost effectiveness of surgical versus nonoperative treatment for lumbar disc herniation over two years: Evidence from the Spine Patient Outcomes Research Trial (SPORT), Spine, 2008: 33(19):2108-2115⁵⁹

Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness
Economic analysis: CUA (health outcome: QALY) Study design: both randomised and observational cohorts of the SPORT trial combined and analysed	Population: Adults with a diagnosis of intervertebral disc herniation. Cohort settings: N: Intervention 1: 775	Total costs (mean per patient): Intervention 1: £12,806 Intervention 2: £3,673 Incremental (2–1): £9,133 (95% CI: NR; p=NR)	QALYs (mean per patient): Intervention 1: 1.64 Intervention 2: 1.44 Incremental (2–1): 0.21 (95% CI: 0.16 – 0.25; p=NR)	ICER (Intervention 2 versus Intervention 1): £43,490 per QALY gained (da) 95% CI: NR – only reported for total costs which include indirect costs. Probability Intervention 2 costeffective (£20K/30K threshold): NR
according to treatment received using regression	Intervention 2: 416	Currency & cost year:		

models 2004 US dollars (presented Analysis of uncertainty: none here as 2004 UK pounds^(d)) Approach to analysis: Start age: Analysis of individual level **Cost components** Intervention 1: 40.7 data for EQ-5D and patientincorporated: Intervention 2: 43.8 reported resource use. Unit Surgery, health care visits, costs applied. Both costs and diagnostic test, medications, Male: EQ-5D are collected at 6 other health care services. Intervention 1: 56% weeks, 3, 6, 12 and 24 Indirect costs were included months. QALYs were Intervention 2: 59% but analysed separately and estimated through timenot reported here. weighted sums of EQ-5D Intervention 1: values adjusted to the overall Standard open mean baseline health state laminotomy/laminectomy with value. removal of the herniation and Perspective: USA health care examination of the involved Follow-up: 2 years nerve root. Surgeons only Treatment effect duration(c): performed other procedures 2 years when it was deemed necessary. **Discounting:** Costs: 3%; Outcomes: 3% Intervention 2:

Data sources

Health outcomes: within-trial analysis **Quality-of-life weights:** EQ-5D US tariff. **Cost sources:** resource use from patient-reported data; unit costs from Medicare payments and Redbook for drugs.

Comments

Source of funding: National institute of Arthritis and Musculoskeletal and Skin Diseases. Limitations: Study conducted in the USA; discount rate is 3%. Outcomes were based also on observational data, not on RCT; costs from US Medicare payments which may not reflect actual costs; resource use was based on patient-reported data which may not be accurate; unclear what parameters at baseline were used to adjust EQ5D data; no sensitivity analyses were conducted and the 95% CI of the ICER was reported only for the total costs (direct and indirect too). Other: it was reported that a total of 63 repeat surgeries occurred in 53 (6.8%) surgery patients. No difference in health care visits, physical therapy visits, chiropractor visits, acupuncture, device use; people in the surgery group reported more diagnostic test use and medication use.

Overall applicability^(a): Partially applicable Overall quality^(b): Potentially serious limitations

Usual care chosen individually by patients and physicians.

Abbreviations: 95% CI: 95% confidence interval; CUA: cost—utility analysis; da: deterministic analysis; EQ-5D: Euroqol 5 dimensions (scale: 0.0 [death] to 1.0 [full health], negative values mean worse than death); ICER: incremental cost-effectiveness ratio; N: sample size; NR: not reported; QALYs: quality-adjusted life years

- (c) For studies where the time horizon is longer than the treatment duration, an assumption needs to be made about the continuation of the study effect. For example, does a difference in utility between groups during treatment continue beyond the end of treatment and if so for how long.
- (d) Converted using 2013 purchasing power parities⁴²
- (e) Directly applicable / Partially applicable / Not applicable
- (f) Minor limitations / Potentially serious limitations / Very serious limitations

Table 25: Tosteson 2008 58

Tosteson AN, Lurie JD, Tosteson TD, Skinner JS, Herkowitz H, Albert T et al. Surgical treatment of spinal stenosis with and without degenerative spondylolisthesis: cost-effectiveness after 2 years. Annals of Internal Medicine. 2008; 149(12):845-853 58

Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness
Economic analysis: CUA (health outcome: QALY) Study design: both randomised and observational cohorts of the SPORT trial combined and analysed according to treatment received using regression models (analysed separately in a sensitivity analysis) Approach to analysis: Analysis of individual level data for EQ-5D and patient-reported resource use. Unit costs applied. Both costs and EQ-5D are collected at 6 weeks, 3, 6, 12 and 24 months. QALYs were estimated through time-weighted sums of EQ-5D values adjusted to baseline age, sex, comorbid stomach	Population: Adults with symptoms for at least 12 weeks and image-confirmed diagnosis of spinal stenosis without degenerative spondylolisthesis. Cohort settings: N: Intervention 1: 394 Intervention 2: 240 Start age: Intervention 1: 63.6 Intervention 2: 66.3 Male: Intervention 1: 61% Intervention 2: 60%	Total costs (mean per patient): Intervention 1: £11,193 Intervention 2: £4,531 Incremental (2–1): £6,661 (95% CI: NR; p=NR) Currency & cost year: 2004 US dollars (presented here as 2004 UK pounds ^(d)) Cost components incorporated: Surgery, health care visits, diagnostic test, medications, other health care services. Indirect costs were included but analysed separately and not reported here.	QALYs (mean per patient): Intervention 1: 1.54 Intervention 2: 1.37 Incremental (2–1): 0.17 (95% CI: NR; p=NR)	ICER (Intervention 2 versus Intervention 1): £44,865 per QALY gained (da) 95% CI: 31,617 – 66,191 Probability Intervention 2 costeffective (£20K/30K threshold): NR Analysis of uncertainty: indirect costs were included in all the sensitivity analyses conducted: observational and randomised cohorts were analysed separately and no major difference between the two ICERs was observed; adjusting for observed mortality decreased the ICER only slightly; the ICER increased when QALYs were estimated with SF-6D and when higher surgery cost was used.

conditions, straight leg raise or femoral tension sign, smoking, comorbid joint conditions, patient selfassessed health trend, annual income, compensation, BMI, EQ5D and centre.

Perspective: USA health care

Follow-up: 2 years

Treatment effect duration(c):

2 years

Discounting: Costs: 3%;

Outcomes: 3%

Standard posterior laminectomy.

Intervention 2:

Usual care chosen individually by patients and physicians.

Data sources

Health outcomes: within-trial analysis **Quality-of-life weights:** EQ-5D US tariff. **Cost sources:** resource use from patient-reported data; unit costs from Medicare payments and Redbook for drugs.

Comments

Source of funding: National institute of Arthritis and Musculoskeletal and Skin Diseases. **Limitations:** Study conducted in the USA; discount rate is 3%. Outcomes were based also on observational data, not on RCT; costs from US Medicare payments which may not reflect actual costs; resource use was based on patient-reported data which may not be accurate; sensitivity analyses were conducted using both direct and indirect costs. **Other:** No difference in health care visits, physical therapy visits, chiropractor visits, acupuncture, device use; people in the surgery group reported more diagnostic test use and medication use.

Overall applicability^(a): Partially applicable Overall quality^(b): Potentially serious limitations

Abbreviations: 95% CI: 95% confidence interval; CUA: cost—utility analysis; da: deterministic analysis; EQ-5D: Euroqol 5 dimensions (scale: 0.0 [death] to 1.0 [full health], negative values mean worse than death); ICER: incremental cost-effectiveness ratio; N: sample size; NR: not reported; QALYs: quality-adjusted life years

- (a) For studies where the time horizon is longer than the treatment duration, an assumption needs to be made about the continuation of the study effect. For example, does a difference in utility between groups during treatment continue beyond the end of treatment and if so for how long.
- (b) Converted using 2013 purchasing power parities⁴²
- (c) Directly applicable / Partially applicable / Not applicable
- (d) Minor limitations / Potentially serious limitations / Very serious limitations

Table 26: van den Hout 2008⁶²

van den Hout WB, Peul WC, Koes BW, Brand R, Kievit J, Thomeer RT. Prolonged conservative care versus early surgery in patients with sciatica from lumbar disc

Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness
Economic analysis: CUA (health outcome: QALY) Study design: Within-trial analysis (associated clinical paper Peul 2008 ⁴³) Approach to analysis: Analysis of individual level data for EQ-5D and patient- reported resource use. Unit costs applied. Both costs and EQ-5D are collected at 2, 4, 8, 12, 26, 38 and 52 weeks. Perspective: Dutch health care Follow-up: 1 years Treatment effect duration (c): 6 months Discounting: Costs: n/a; Outcomes: n/a	Population: patients aged 18 to 65 with a radiologically confirmed disc herniation and lumbosacral radicular syndrome that had lasted for 6 to 12 weeks. Cohort settings: N: Intervention 1: 141 Intervention 2: 142 Start age: Intervention 1: 42 Intervention 2: 43 Male: Intervention 1: 63% Intervention 2: 68% Intervention 2: 68% Intervention 5: Early surgery; disc herniation was removed through a unilateral transflaval approach using magnification. Intervention 2: Prolonged conservative care provided by the GP; if sciatica persisted at 6 months, microdiscectomy was offered.	Total costs (mean per patient): Intervention 1: £4,347 Intervention 2: £2,942 Incremental (2–1): £1,405 (95% CI: 651 – 2,156; p<0.001) Currency & cost year: 2008 Euros (presented here as 2008 UK pounds ^(d)) Cost components incorporated: Surgery with admissions to hospital, physical therapy, visits, homecare, drugs and aids. Indirect and societal costs were included but analysed separately and not reported here.	QALYs (mean per patient): Intervention 1: 0.78 Intervention 2: 0.73 Incremental (2–1): 0.044 (95% CI: 0.005-0.083; p=0.03)	ICER (Intervention 2 versus Intervention 1): £ 31,932 per QALY gained 95% CI: 10,817 – 332,249 Probability Intervention 2 costeffective (£20K/30K threshold): NR Analysis of uncertainty: when SF-6E was used as an alternative utility measure the QALY difference was 0.024, resulting in an ICER of £58,541.

Increasing leg pain not responsive to drugs and progressive neurological deficit were reasons for performing surgery earlier than 6 months.

Data sources

Health outcomes: within-trial analysis Quality-of-life weights: EQ-5D UK tariff. Cost sources: resource use from patient-reported data; unit costs from prices set up by the hospital for the intervention; other costs from Dutch standard prices.

Comments

Source of funding: Netherlands Organization for Health research and Development. Limitations: Study conducted in the Netherlands. Intervention not described in detail in this paper. Patients in the usual care group could have surgery after the initial 6 months and outcomes were collected up to 1 year. Short time horizon; resource use was based on patient-reported data which may not be accurate; hospital prices were used. Other: During the first year surgery was performed in 89% of patients in the early surgery group and 40% of the prolonged conservative care group.

Overall applicability^(a): Partially applicable Overall quality^(b): Potentially serious limitations

Abbreviations: 95% CI: 95% confidence interval; CUA: cost-utility analysis; da: deterministic analysis; EQ-5D: Eurogol 5 dimensions (scale: 0.0 [death] to 1.0 [full health], negative values mean worse than death); ICER: incremental cost-effectiveness ratio; N: sample size; n/a: not applicable; NR: not reported; QALYs: quality-adjusted life years

- (a) For studies where the time horizon is longer than the treatment duration, an assumption needs to be made about the continuation of the study effect. For example, does a difference in utility between groups during treatment continue beyond the end of treatment and if so for how long.
- (b) Converted using 2013 purchasing power parities⁴²
- (c) Directly applicable / Partially applicable / Not applicable
- (d) Minor limitations / Potentially serious limitations / Very serious limitations

Spinal fusion 1.20

Table 27: Fritzell 2011¹³ (also published by Berg 2011⁵)

Fritzell P, Berg S, Borgstrom F, Tullberg T, Tropp H. Cost effectiveness of disc prosthesis versus lumbar fusion in patients with chronic low back pain: randomized controlled trial with 2-year follow-up. European Spine Journal. 2011; 20(7):1001-1011. (Guideline Ref ID FRITZELL2011)

Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness
Economic analysis: CUA	Population:	Total costs (mean per	QALYs (mean per patient):	ICER (Intervention 2 versus Intervention 1):
(health outcome: QALYs)	Adults (21-55 years) with	patient):	Intervention 1: 0.41	Intervention 1 dominates intervention 2
	low back pain with/without	Intervention 1: £10,194	Intervention 2: 0.40	(lower costs and higher QALYs)
Study design: Within-trial	sciatica. Patients had	Intervention 2: £11,780	Incremental (2-1): -0.01	95% CI: NR
analysis (RCT, associated	suffered at least 12 months			Probability Intervention 2 cost-effective

clinical paper Berg 2009)

Approach to analysis: EQ-5D data collected preoperatively, 1 year and 2 years follow-up. QALYs constructed through area under the curve method. Resource use captured from patient cost diaries (at 1, 3, 6, 12, 18 and 24 months), unit costs applied. Surgical procedure resource use estimated from index

Perspective: Swedish healthcare payer perspective
Follow-up: 2 years
Discounting: No discounting applied in base case analysis

episode.

from what was understood to be discogenic low back pain in one or two motion segments between L3 and S1; they could also have additional nonspecific leg pain.

Cohort settings:

Start age: 39 Male: 59%

Intervention 1: (n=80) Total disc replacement surgery

Intervention 2: (n=72)
Fusion (either ALIF or PLIF according to surgeon preference)

Incremental (2-1): £1,587 (95% CI: £83 to £2,971; p=NR)

(95% CI: NR; p=NR)

Cost breakdown (mean per patient):
Hospital cost index procedure:

Intervention 1: £7,287 Intervention 2: £7.390

Hospital costs after index procedure:

Intervention 1: £1,070 Intervention 2: £2,301

Primary/Private care:

Intervention 1: £1,666 Intervention 2: £1,844

Back-related drugs:

Intervention 1: £172 Intervention 2: £246

Currency & cost year:

2006 Swedish Krona (presented here as 2006 UK pounds^(a))

Cost components incorporated:

Intervention cost (index procedure for surgery),

(£20K/30K threshold): NR

Analysis of uncertainty: Bootstrapping of ICER conducted but only from a societal perspective not a health care provider perspective. Therefore this is not reported here.

Two additional sensitivity analyses were conducted.

- The costs were discounted at 3%, this did not impact the total cost difference between the two comparators.
- Reoperation costs were excluded from total healthcare costs. The total costs (mean per patient) were:

Intervention 1: £9,710 Intervention 2: £10,235 Incremental (2–1): £525

(95% CI: -£827 to £1,710; p=NR)

(including private care) and back-related drug costs.	
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Data sources

Health outcomes: Within-trial analysis (RCT, Berg 2009)⁶. Health outcomes included patient reported EQ-5D collected pre-operatively, 1 year and 2 years follow-up, other outcomes included Oswestry Disability Index, back pain (VAS) and patient-reported outcome (see clinical review, Berg 2009). QALYs were calculated using the area under the curve approach adjusted for baseline utility. Quality-of-life weights: EQ-5D, Swedish tariff. Cost sources: Resource use and cost for interventions and post-surgery hospital stay based on index procedures/episodes (within-trial and Stockholm Spine Center). Other resource use captured from patient cost diaries. Unit costs from Swedish national board of health and welfare and Swedish published drug costs.

Comments

Source of funding: DePuySpine, Medtronic and Synthesis, manufacturers of surgical devices. **Limitations:** Swedish resource use data (2002-2005) and unit costs (2006) may not reflect current NHS context. No discounting applied in base case analysis, discounting of costs at 3% applied in sensitivity analysis, however this is not in line with NICE reference case. Within-trial analysis and so does not reflect full body of available evidence for this comparison; Berg 2009 is one of the studies included in the clinical review for disc replacement surgery. Bootstrapping of ICER not undertaken from a healthcare payer perspective. Potential conflict of interest, study funded by manufacturers of surgical devices. **Other:** n/a

Overall applicability(b)(a): Partially applicable Overall quality(c): Potentially serious limitations

Abbreviations: 95% CI: 95% confidence interval; CUA: cost—utility analysis; EQ-5D: Euroqol 5 dimensions (scale: 0.0 [death] to 1.0 [full health], negative values mean worse than death); ICER: incremental cost-effectiveness ratio; NR: not reported; QALYs: quality-adjusted life years

- (a) Converted using 2006 purchasing power parities⁴²
- (b) Directly applicable / Partially applicable / Not applicable
- (c) Minor limitations / Potentially serious limitations / Very serious limitations

Table 28: Rivero-Arias 2005⁴⁶

Rivero-Arias O, Campbell H, Gray A, Fairbank J, Frost H, Wilson-MacDonald J. Surgical stabilisation of the spine compared with a programme of intensive rehabilitation for the management of patients with chronic low back pain: cost utility analysis based on a randomised controlled trial. British Medical Journal. 2005; 330: 1239-1243:1239-1243. (Guideline Ref ID RIVEROARIAS2005)

Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness
Economic analysis: CUA	Population:	Total costs (mean per	QALYs (mean per patient):	ICER (Intervention 2 versus Intervention 1):
(health outcome: QALYs)	Adults with chronic low back	patient):	Intervention 1: 0.936	£48,515 per QALY gained (pa)
	pain	Intervention 1: £4,419	Intervention 2: 1.004	95% CI: NR
Study design: Within-trial	Cohort settings:	Intervention 2: £7,718	Incremental (2-1): 0.068	Probability Intervention 2 cost-effective

(RCT, associated clinical paper Fairbank 2005)

Approach to analysis: EQ-5D data collected at baseline, 6, 12 and 24 months follow-up. QALYs constructed through area under the curve method. Within-trial reported resource use, including patient-reported resource use for medication use, over 24 months, unit costs applied.

Perspective: UK NHS Follow-up: 2 years Discounting: Costs: 3.5%;

Outcomes: 3.5%

Age range: 18-55 years

Male: 49%

Intervention 1: (n=139)

Intensive rehabilitation programme-3 element MBR program (paced exercise and education programme based on cognitive behavioural approaches). Total duration approximately 75 hours.

Intervention 2: (n=151)
Fusion(technique based on surgeon preference)

Incremental (2–1): £3,299 (95% CI: £2,322 to £4,267; p<0.001)

Cost breakdown (mean per patient):

Intervention cost:

Intervention 1: £1,410 Intervention 2: £6,011

Other back-related related NHS contacts (up to 24 months):

Intervention 1: £3,009 Intervention 2: £1,707

Currency & cost year: 2002-2003 UK pounds

Cost components incorporated:

Intervention costs (including staff time and other resource use such as surgical implants and equipment) and other back pain related NHS contacts up to 24 months (including surgical follow-up appointments, physiotherapy outpatient appointments, unplanned or other back-related hospital admission, HCP contacts,

(95% CI: -0.02 to 0.156; p=0.13)

(£20K): ~5% (reading from graph) – see caveat regarding perspective below.

Analysis of uncertainty: Bootstrapping of ICER conducted but only using a total costs including patient-related costs (broader perspective) not a NHS perspective.

Sensitivity analyses were conducted assuming different surgical technique costs:

- posterolateral technique (least expensive procedure): ICER 2 vs 1 = £35,338 per QALY

- 360 degree fusion (most expensive procedure): ICER 2 vs 1 = £60,765 per QALY

Further sensitivity analysis by varying the time horizon to 4 years (assuming treatment differences for utilities were maintained):

Finally, they examined impact of patients receiving other interventions subsequent to allocated intervention (at 2 years 45 patients had received both interventions) by assuming that people in each arm continued to receive both treatments in years 3,4 and 5 at rates observed in year 1 and 2: ICER =£16,824 per QALY. The same sensitivity analysis was done but assuming half the rate observed at year 1 and 2 applied: ICER = £31,838 per QALY.

ICER = £25,398 per QALY.

Note, these were all conducted using the broader perspective (including patient-related costs).

prescriptions).

Data sources

Health outcomes: Within-trial analysis (RCT, Fairbank 2005)¹⁰. Health outcomes included patient reported EQ-5D collected baseline, 6, 12 and 24 months follow-up. QALYs were calculated using the area under the curve approach adjusted for baseline utility. **Quality-of-life weights:** EQ-5D UK tariff. **Cost sources:** Within-trial reported resource use and patient-reported resource use for medication use, over 24 months. UK national average unit costs.

Comments

Source of funding: UK Medical Research Council. **Limitations:** UK NHS resource use data (1996-2002) and unit cost (2002-2003) may not reflect current NHS context. Within-trial analysis and so does not reflect full body of available evidence for this comparison; Fairbank 2005 is 1 of 4 studies included in the clinical review for spinal fusion versus other treatments. Sensitivity analyses were conducted using a broader perspective which included patient-related costs. **Other:**

Overall applicability(a): Partially applicable Overall quality(b): Potentially serious limitations

Abbreviations: 95% CI: 95% confidence interval; CUA: cost—utility analysis; da: deterministic analysis; EQ-5D: Euroqol 5 dimensions (scale: 0.0 [death] to 1.0 [full health], negative values mean worse than death); ICER: incremental cost-effectiveness ratio; NR: not reported; QALYs: quality-adjusted life years

- (a) Directly applicable / Partially applicable / Not applicable
- (b) Minor limitations / Potentially serious limitations / Very serious limitations

I.21 Disc replacement

For Fritzell 2011¹³ (also published by Berg 2011⁵) please see Table 27 (Spinal fusion) above.

Table 29: Johnsen 2014²⁸

Johnsen LG, Hellum C, Storheim K, Nygaard OP, Brox JI, Rossvoll I et al. Cost-effectiveness of total disc replacement versus multidisciplinary rehabilitation in patients with chronic low back pain: A norwegian multicenter RCT. Spine. 2014; 39(1):23-32²⁸

•				
Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness
Economic analysis: CUA (health outcome: QALYs) Study design: Within-trial analysis (RCT, same paper and other associated	Population: Patients with chronic low back pain for more than one year and degenerative changes in lumbosacral intervertebral discs.	Total costs (mean per patient): Intervention 1: £8299 Intervention 2: £5054 Incremental (2–1): £3245 (95% CI: NR; p=NR)	QALYs (mean per patient): Intervention 1: 1.29 Intervention 2: 0.95 Incremental (2–1): 0.34 (95% CI: 0.18-0.5; p<0.001)	ICER (Intervention 2 versus Intervention 1): £9544 per QALY gained (da) Analysis of uncertainty: Bootstrapping analysis was conducted using a societal perspective and therefore the 95% CI around
clinical paper Hellum 2011 ^{17-19,27} Approach to analysis:	Cohort settings: Start age: 41	Currency & cost year: 2012 euros (presented here		the ICER is not reported. Using the intention to treat analysis total disc replacement was more costly but also more

EQ-5D data collected at baseline, 6 weeks, and 3, 6, 12, 24 months follow-up. QALYs constructed through area under the curve method. Resource use captured from patient cost diaries (at 6 weeks, and at 3, 6, 12, 18 and 24 months), unit costs applied. Multiple imputation was used when data were missing.

Perspective: Norwegian healthcare payer
Follow-up: 2 years
Discounting: none

Male: 47%

Intervention 1:

Total disc replacement

Intervention 2:

3-element MBR (outpatient programme with cognitive, physical and education components; the treatment was interdisciplinary and directed by a team of physiotherapists and specialists in physical medicine and rehabilitation and lasted for approximately 60 hours during 3 to 5 weeks)

as 2012 UK pounds(d))

Cost components incorporated:

Cost of intervention, hospital follow up (reoperations, admissions, visits), GP consultations, physical therapist consultations, visits to complementary practitioners, medications. effective, however the costs included the societal perspective therefore results are reported.

Where missing data were not inputed but dropped, the effectiveness of total disc replacement was lower, however the costs included the societal perspective therefore results are reported.

When SF-6D instead of EQ5D was used, the incremental QALY gain was 0.11, and the ICER was £29,500.

Data sources

Health outcomes: within-trial analysis (same study and Hellum 2011^{17-19,27} **Quality-of-life weights:** EQ-5D UK tariff and SF-6D **Cost sources:** For rehab a top-down approach was used, that is the total cost of a spine clinic was estimated and then how much of the clinic's costs were associated with MDR was determined; spare capacity was included; Norwegian national sources were used.

Comments

Source of funding: national funds through the Norwegian Back Pain association funds. **Limitations:** Norwegian resource use data (2004-2007) and unit costs may not reflect current NHS context. No discounting conducted. Within-trial analysis and so does not reflect full body of available evidence for this comparison. Bootstrapping of ICER not undertaken. **Other:**

Overall applicability(a): Partially applicable Overall quality(b): Potentially serious limitations

Abbreviations: 95% CI: 95% confidence interval; CUA: cost—utility analysis; da: deterministic analysis; EQ-5D: Euroqol 5 dimensions (scale: 0.0 [death] to 1.0 [full health], negative values mean worse than death); ICER: incremental cost-effectiveness ratio; NR: not reported; QALYs: quality-adjusted life years

- (g) Converted using 2012 purchasing power parities⁴²
- (h) Directly applicable / Partially applicable / Not applicable
- (i) Minor limitations / Potentially serious limitations / Very serious limitations

Appendix J: GRADE tables

J.1 Clinical examination

None.

J.2 Risk assessment tools and stratification

Table 30: Clinical evidence profile: Hicks/Delitto classification versus no risk tool stratification

			Qualit	y assessment			No of patients			Effect	Ovality	Importono
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Stratified treatment versus non-stratified treatment-Delitto Classification	Control	Relative (95% CI)	Absoluto	Quanty	Importance
QoL (SF	-36, PCS,0-	100) ≤4	months (follow-up	4 weeks; Bette	er indicated by lo	ower values)						
	randomised trials		no serious inconsistency	no serious indirectness	very serious ²	none	37	41	-	MD 6.2 higher (8.74 lower to 21.14 higher)	⊕OOO VERY LOW	CRITICAL
QoL(SF	-36,PCS,0-1	00) >4 m	nonths - 1 year (fol	low-up >4 mon	ths - 1 year; Bett	er indicated by lower v	values)					
	randomised trials		no serious inconsistency		no serious imprecision	none	111	123	-	MD 0.59 lower (3.7 lower to 2.52 higher)		CRITICAL
QoL (SF	-36, MCS,0	-100) ≤4	months (follow-up	mean 4 weeks	; Better indicate	d by lower values)						
	randomised trials		no serious inconsistency	no serious indirectness	very serious ²	none	37	41	-	MD 1.6 higher (13.34 lower to 16.54 higher)	⊕000 VERY LOW	CRITICAL

QoL(SF	-36,MCS,0-1	100) >4 n	nonths - 1 year (fo	ollow-up >4 mor	nths - 1 year; Bet	ter indicated by lower	values)						
2	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none		111	123	-	MD 0.94 higher (2.24 lower to 4.12 higher)	⊕⊕OO LOW	CRITICAL
Pain(NI	RS,0-10) ≤ 4	months	(follow-up 8 week	s; Better indica	ated by lower val	ues)							
1	randomised trials		no serious inconsistency	no serious indirectness	serious ²	none	156		-	-	MD 0.49 lower (1.34 lower to 0.36 higher)		CRITICAL
Pain(NI	RS,0-10) >4 ı	months -	- 1 year (follow-up	1 year; Better	indicated by low	er values)							
1	randomised trials		no serious inconsistency	no serious indirectness	serious ²	none		156	-	-	MD 0.13 higher (0.83 lower to 1.09 higher)	⊕OOO VERY LOW	CRITICAL
Functio	on(ODI,0-100) ≤ 4 mo	nths (follow-up ≤₄	4 months; Bette	er indicated by lo	wer values)							
2	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none		111	123	-	MD 1.16 lower (5.13 lower to 2.82 higher)		CRITICAL
Functio	on(ODI,0-100) > 4 mo	onth (follow-up >4	months - 1 yea	r; Better indicate	ed by lower values)							
2	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	,	111	123	-	MD 0.23 higher (4.09 lower to 4.54 higher)	⊕⊕OO LOW	CRITICAL
Respor	nder criteria(NRS>30	% improvement)	≤ 4 months (foll	low-up 8 weeks)								
1	randomised trials		no serious inconsistency	no serious indirectness	serious ²	none	44/74 (59.5%)	73.2%	(0.	0.81 65 to .02)	139 fewer per 1000 (from 256 fewer to 15 more)	⊕OOO VERY LOW	IMPORTAN T
Respor	nder criteria(NRS>30	% improvement)>	4 months - 1 ye	ear (follow-up 1 y	/ears)							
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	57/74 (77%)	74.4%	R 1.0 (0.)4 87	nore per 1000 (from 97 fewer to 179 more)	⊕⊕OO LOW	IMPORTAN T

spo	nder criteria(ODI>30	% improvement)	≤ 4 months (foll	ow-up 8 weeks)	<u> </u>					T	
	randomised trials	,	no serious inconsistency	no serious indirectness	serious²	none	27/74 (36.5%)	45.1%	RR 0.81 (0.55 to 1.19)	86 fewer per 1000 (from 203 fewer to 86 more)	⊕OOO VERY LOW	IMPORTA T
spo	nder criteria(ODI>30	% improvement)	>4 months - 1 ye	ar (follow-up 1 y	ears)					T	
	randomised trials		no serious inconsistency	no serious indirectness	serious ²	none	60/74 (81.1%)	68.3%	RR 1.19 (0.99 to 1.43)	130 more per 1000 (from 7 fewer to 294 more)	⊕OOO VERY LOW	IMPORTA T
umbe	r of therapy	appoint	ments ≤ 4 montl	ns (follow-up 4 w	eeks; Better ind	icated by lower values			·			
					serious ²	none		37	41	- MD 0.3 lower (1.68	Φ000	
	randomised trials	,	no serious inconsistency	no serious indirectness	Serious	none			7.	lower to 1.08 higher)		IMPORTA T
umbe	trials	serious ¹	inconsistency	indirectness		ter indicated by lower v					VERY	IMPORTA T

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias 2 Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

|--|

			Quality ass	sessment			No of patients			Effect	Quality	Importanc
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Stratified treatment versus non- stratified treatment-O'Sullivan Classification	Contro I	Relative (95% CI)	Absolute		е

	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	51	43	-	MD 2.1 lower (2.83 to 1.37 lower)	⊕⊕OO LOW	CRITICA
in(VAS,0-10)>4 mor	nths - 1 ye	ear (follow-up 1	years; Better in	dicated by low	er values)						
	randomised trials	very serious	no serious inconsistency	no serious indirectness	serious ²	none	51	43	-	MD 1.5 lower (2.33 to 0.67 lower)	⊕000 VERY LOW	CRITICA
ıc	tion(ODI,0-100) ≤	4 months	s (follow-up 3 m	onths; Better in	dicated by low	er values)						
	randomised trials	very serious	no serious inconsistency	no serious indirectness	no serious imprecision	none	51	43	-	MD 10.9 lower (13.94 to 7.86 lower)	⊕⊕OO LOW	CRITICA
	(0.01.0.400)	4 months	- 1 year (follow	-up 1 years; Bet	ter indicated by	lower values)		—				ļ
10	tion(ODI,0-100)>											CRITICA

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

Table 32: Clinical evidence profile: STarT Back classification versus no risk tool classification

			Quality ass	sessment			No of patients			Effect	Quality	Importance
No of studies	Design Inconsistancy Indirectness Imprecision Versus non-stratitied Absolute											
Quality of	f life (SF-12,	PCS,0-10	0) <4 months (fol	low-up 4 month	s; Better indic	ated by lower valu	ues)					1

1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	568	283	-	MD 2.3 higher (0.42 to 4.18 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	f life (SF-12,	PCS,0-10	0) >4 months (fo	llow-up 12 mon	ths; Better indi	cated by lower va	lues)					
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	568	283	-	MD 2.3 higher (0.73 to 3.87 higher)	⊕000 VERY LOW	CRITICAL
Quality o	f life (SF-12,	MCS,0-10	0) <4 months (fo	llow-up 4 mont	hs; Better indic	ated by lower val	ues)					
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	568	283	-	MD 0 higher (1.58 lower to 1.58 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	f life (SF-12,	MCS,0-10	0) >4 months (fo	ollow-up 12 mon	ths; Better ind	icated by lower va	ilues)					
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	568	283	-	MD 0.5 higher (1.39 lower to 2.39 higher)	⊕⊕OO LOW	CRITICAL
Pain(VAS	6/NRS,0-10)<	4 months	(follow-up <4 m	onths; Better in	dicated by low	er values)						
2	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	635	316	-	not pooled	⊕⊕OO LOW	CRITICAL
Pain(VAS	5,0-10)>4 moi	nths (folio	ow-up 12 months	; Better indicate	ed by lower val	ues)						
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	568	283	-	MD 0.2 lower (0.58 lower to 0.18 higher)	⊕⊕⊕O MODERATE	CRITICAL
Function	(RMDQ/ODI,	0-24)< 4 m	l nonths (follow-up	o <4 months; Be	etter indicated l	by lower values)						
	randomised trials	very serious ¹	serious ³	no serious indirectness	serious ²	none	635	316	-	SMD 0.34 lower (0.47 to 0.2 lower)	⊕000 VERY LOW	CRITICAL
Function	(RMDQ,0-24)	>4 month	s (follow-up 12 r	months; Better i	ndicated by lov	ver values)						

	randomised	very	no serious	no serious	no serious	none	568	283	-	MD 1 lower (1.89 to ⊕⊕OO	
	trials	serious ¹	inconsistency	indirectness	imprecision					0.11 lower) LOW	
ychol	ogical Distres	ss (HADS	, anxiety subsca	ale, 0-21)< 4 mo	nths (follow-up	4 months; Bette	r indicated by lower val	lues)			
		serious ¹	no serious	no serious	no serious	none	568	283	-	MD 0.5 lower (1.05 ⊕⊕⊕O	CRITICA
	trials		inconsistency	indirectness	imprecision					lower to 0.05 MODERATE higher)	
sychol	ogical Distres	ss (HADS	, anxiety subsca	 ale, 0-21)> 4 mo	nths (follow-up	12 months; Bett	er indicated by lower va	alues)			
	randomised	very	no serious	no serious	no serious	none	568	283	-	MD 0.3 lower (0.9 ⊕⊕OO	CRITICA
	trials	serious ¹	inconsistency	indirectness	imprecision					lower to 0.3 higher) LOW	
sychol	ogical Distres	ss (HADS	depression su	bscale, 0-21)< 4	months (follow	w-up 4 months; E	setter indicated by lowe	r values)			
	randomised	very	no serious	no serious	no serious	none	568	283	-	MD 0.3 lower (0.87 ⊕⊕OO	CRITICA
	trials	serious ¹	inconsistency	indirectness	imprecision					lower to 0.27 LOW higher)	
sychol	ogical Distres	ss (HADS	, depression su	bscale, 0-21) >4	months (follow	w-up 12 months;	Better indicated by low	er values)			
	randomised	very	no serious	no serious	serious ²	none	568	283	-	MD 2.3 lower (2.88 ⊕OOO	CRITICA
	trials	serious ¹	inconsistency	indirectness						to 1.72 lower) VERY LOW	
uality	of life (SF-12,	PCS,0-10	0) <4 months(sf	tratified) - Low-	Risk (Better inc	dicated by lower	/alues)				
	randomised	very	no serious	no serious	serious ²	none	148	73	-	MD 1.4 higher (1.31 ⊕OOO	CRITICA
	trials	serious ¹	inconsistency	indirectness						lower to 4.11 VERY LOW higher)	
	of life (SF-12	PCS,0-10	0) <4 months(st	ratified) - Medi	um-risk (follow	up 4 months; Be	tter indicated by lower	values)			
uality	or line (31 -12,				1 . 2	none	263	131	-	MD 2.7 higher (0.39 ⊕OOO	CRITICA
uality		very	no serious	no serious	serious ²	none					
uality		very serious ¹	no serious inconsistency	no serious indirectness	serious ²	lione				to 5.01 higher) VERY LOW	

1	randomised	very	no serious	no serious	serious ²	none	157	79	_	MD 2.5 higher (1.71	⊕OOO	CRITICAL
	trials	serious ¹	inconsistency	indirectness	Scrious	none	107	13	-	lower to 6.71 higher)	VERY LOW	ONTIOAL
Quality o	f life (SF-12,	PCS,0-10	0) >4 months(st	ratified) - Low-R	isk (follow-up 1	2 months; Better	indicated by lower value	es)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	148	73	-	MD 1.6 higher (1.19 lower to 4.39 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	f life (SF-12,	PCS,0-10	0) >4 months(str	ratified) - Mediu	m-risk (follow-เ	ip 12 months; Bet	ter indicated by lower va	alues)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	261	131	-	MD 3.1 higher (0.66 to 5.54 higher)	⊕000 VERY LOW	CRITICAL
Quality o	f life (SF-12,	PCS,0-10	0) >4 months(st	ratified) - High-ri	isk (follow-up 1	2 months; Better	indicated by lower value	es)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	157	79	-	MD 1.8 higher (1.66 lower to 5.26 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	f life (SF-12,	MCS,0-10	0) <4 months(st	ratified) - Low-R	lisk (follow-up	4 months; Better i	ndicated by lower value	s)				
1		very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	148	73	-	MD 1.5 lower (4.58 lower to 1.58 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	f life (SF-12,	MCS,0-10	0) <4 months(st	ratified) - Mediu	m-risk (follow-	up 4 months; Bett	er indicated by lower va	lues)		<u> </u>		
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	263	131	-	MD 0.4 higher (2.01 lower to 2.81 higher)	⊕⊕OO LOW	CRITICAL
Quality o	f life (SF-12,	MCS,0-10	0) <4 months(st	ratified) - High-r	isk (follow-up	l months; Better i	ndicated by lower values	5)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	very serious ²	none	157	79	-	MD 0.7 higher (3.01 lower to 4.41 higher)	⊕OOO VERY LOW	CRITICAL

Quality o	of life (SF-12,	MCS,0-10	0) <4 months(str	atified) - Low-Ri	isk (follow-up 1	2 months; Better	indicated by lower value	s)				
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	148	73	-	MD 1.7 lower (4.55 lower to 1.15 higher)	⊕000 VERY LOW	CRITICA
uality o	of life (SF-12,	MCS,0-10	0) <4 months(str	atified) - Mediur	n-risk (follow-u	p 12 months; Bet	ter indicated by lower va	lues)				
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	263	131	-	MD 1.1 higher (1.53 lower to 3.73 higher)	⊕OOO VERY LOW	CRITICA
uality o	of life (SF-12,	MCS,0-10	0) <4 months(str	atified) - High-ri	sk (follow-up 1	2 months; Better	indicated by lower value	s)				
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	157	79	-	MD 1.9 higher (1.83 lower to 5.63 higher)	⊕OOO VERY LOW	CRITICA
in(VA	5,0-10)< 4 mc	onths(stra	tified) - Low-Risl	र (follow-up <4 ।	months; Better	indicated by lowe	er values)					
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	163	87	-	MD 0.14 lower (0.68 lower to 0.4 higher)	⊕OOO VERY LOW	CRITICA
ain(VA	6,0-10)< 4 mc	onths(stra	l tified) - Medium-	risk (follow-up	<4 months; Bet	ter indicated by lo	ower values)	1				
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	294	143	-	MD 0.81 lower (1.25 to 0.37 lower)	⊕OOO VERY LOW	CRITICA
ain(VA	5,0-10)< 4 mc	onths(stra	tified) - High-risl	(follow-up <4 r	months; Better	indicated by lowe	r values)	1		<u>'</u>		
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	178	86	-	MD 0.76 lower (1.43 to 0.1 lower)	⊕OOO VERY LOW	CRITICA
ain(VA	6,0-10)>4 mo	nths(strat	ified) - Low-Risk	(follow-up 12 n	nonths; Better	indicated by lowe	r values)	1				
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	148	73	-	MD 0 higher (0.66 lower to 0.66	⊕⊕OO LOW	CRITICA
	1		1	1	1	1		1		1		

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in(VA	S,0-10)>4 mo	nths(stra	tified) - High-ris	k (follow-up 12 i	months; Better	indicated by lo	wer values)					
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	157	79	-	MD 0.1 lower (0.92 lower to 0.72 higher)	⊕⊕OO LOW	CRITICA
ınctio	n(RMDQ/ODI)	< 4 mont	hs (stratified) - L	ow-Risk (follow	v-up <4 months	; Better indicate	ed by lower values)					
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	163	87	-	SMD 0.22 lower (0.48 lower to 0.05 higher)	⊕⊕OO LOW	CRITICA
ınctio	n(RMDQ/ODI)	< 4 mont	hs (stratified) - N	ledium-risk (fol	low-up <4 mon	ths; Better indic	cated by lower values)					
	randomised trials	very serious ¹	serious ³	no serious indirectness	serious ²	none	294	143	-	SMD 0.39 lower (0.59 to 0.18 lower)	⊕OOO VERY LOW	CRITICA
ınctio	n(RMDQ/ODI)	< 4 mont	hs (stratified) - H	ligh-risk (follow	-up <4 months	; Better indicate	d by lower values)					
unctio	randomised trials	< 4 montl very serious ¹	hs (stratified) - F	ligh-risk (follow no serious indirectness	-up <4 months serious ²	none	d by lower values)	86	-	SMD 0.38 lower (0.64 to 0.12 lower)	⊕OOO VERY LOW	CRITICA
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none		86	-			CRITICA
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	178	73	-			CRITICA
unctio	randomised trials n(RMDQ,0-24) randomised trials	very serious ¹ > 4 mont very serious ¹	no serious inconsistency ths (stratified) - I no serious inconsistency	no serious indirectness Low-Risk (follow no serious indirectness	serious ² v-up 12 months no serious imprecision	none s; Better indicate	178 ed by lower values)		-	(0.64 to 0.12 lower) MD 0.4 lower (1.72 lower to 0.92	VERY LOW	
unctio	randomised trials n(RMDQ,0-24) randomised trials	very serious ¹ > 4 mont very serious ¹	no serious inconsistency ths (stratified) - I no serious inconsistency	no serious indirectness Low-Risk (follow no serious indirectness	serious ² v-up 12 months no serious imprecision	none s; Better indicate	ed by lower values)		-	(0.64 to 0.12 lower) MD 0.4 lower (1.72 lower to 0.92	VERY LOW	
unctio	randomised trials n(RMDQ,0-24) randomised trials n(RMDQ,0-24) randomised trials	very serious¹ very serious¹ > 4 mont very serious¹	no serious inconsistency Ins (stratified) - Ins (s	no serious indirectness Low-Risk (follow no serious indirectness Medium-risk (follow no serious indirectness	serious² v-up 12 months no serious imprecision llow-up 12 mor no serious imprecision	none s; Better indicate none nths; Better indi	178 ed by lower values) 148 cated by lower values)	73	-	(0.64 to 0.12 lower) MD 0.4 lower (1.72 lower to 0.92 higher)	⊕⊕OO LOW	CRITICA

	trials	serious ¹	inconsistency	indirectness	imprecision					higher)	LOW	
Psycholo	ogical Distres	s (HADS,	anxiety subsca	ale, 0-21)< 4 mor	nths(stratified)	· Low-Risk (follow	-up 4 months; Better in	dicated by	lower valu	es)		
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	148	73	-	MD 0.3 higher (0.66 lower to 1.26 higher)	⊕⊕OO LOW	CRITICAL
Psycholo	ogical Distres	s (HADS,	anxiety subsca	nle, 0-21)< 4 mor	nths(stratified) -	· Medium-risk (foll	ow-up 4 months; Better	indicated	l by lower v	alues)		
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	263	131	-	MD 0.9 lower (1.68 to 0.12 lower)	⊕⊕OO LOW	CRITICAL
Psycholo	ogical Distres	s (HADS,	anxiety subsca	nle, 0-21)< 4 mor	nths(stratified)	· High-risk (follow	-up 4 months; Better ind	licated by	lower value	es)		L
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	157	79	-	MD 0.6 lower (1.8 lower to 0.6 higher)	⊕⊕OO LOW	CRITICAL
Psycholo	ogical Distres	s (HADS,	anxiety subsca	ale, 0-21)> 4 mor	nths(stratified)	Low-Risk (follow	-up 12 months; Better in	ndicated b	y lower val	ues)		
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	148	73	-	MD 0.3 higher (0.75 lower to 1.35 higher)	⊕⊕OO LOW	CRITICAL
Psycholo	ogical Distres	s (HADS,	anxiety subsca	ale, 0-21)> 4 mor	nths(stratified)	· Medium-risk (foll	ow-up 12 months; Bette	er indicate	ed by lower	values)		
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	263	131	-	MD 0.7 lower (1.58 lower to 0.18 higher)	⊕⊕OO LOW	CRITICAL
Psycholo	ogical Distres	s (HADS,	anxiety subsca	nle, 0-21)> 4 mor	nths(stratified)	High-risk (follow	-up 12 months; Better in	ndicated b	y lower valu	ues)		
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	157	79	-	MD 0.4 lower (1.71 lower to 0.91 higher)	⊕⊕OO LOW	CRITICAL
Psycholo	ogical Distres	ss (HADS,	depression sul	bscale, 0-21)> 4	months(stratifi	ed) - Low-Risk (fo	llow-up 4 months; Bette	r indicate	d by lower	values)		

1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	148	73	-	MD 0.1 lower (1.02 lower to 0.82 higher)	⊕⊕OO LOW	CRITICAL
Psychological	ogical Distres	s (HADS,	depression sub	oscale, 0-21)> 4	months(stratif	ied) - Medium-ri	isk (follow-up 4 months; B	Better indica	ated by low	er values)		
1		very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	263	131	-	MD 0.5 lower (1.24 lower to 0.24 higher)	⊕⊕OO LOW	CRITICAL
Psychol	ogical Distres	s (HADS,	depression sub	oscale, 0-21)> 4	months(stratif	ied) - High-risk	(follow-up 4 months; Bette	er indicated	d by lower v	ralues)		
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	157	79	-	MD 1.1 lower (2.17 to 0.03 lower)	⊕OOO VERY LOW	CRITICAL
Psychological	ogical Distres	s (HADS,	depression sub	oscale, 0-21)> 4	months(stratif	ied) - Low-Risk	(follow-up 12 months; Bet	tter indicat	ed by lower	values)		
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	148	73	-	MD 0 higher (0.96 lower to 0.96 higher)	⊕⊕OO LOW	CRITICAL
Psychol	ogical Distres	s (HADS,	depression sub	oscale, 0-21)> 4	months(stratif	ied) - Medium-ri	isk (follow-up 12 months;	Better indi	cated by lov	ver values)		
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	263	131	-	MD 0.3 lower (1.09 lower to 0.49 higher)	⊕⊕OO LOW	CRITICAL
Psycholo	ogical Distres	s (HADS,	depression sub	oscale, 0-21)> 4	months(stratif	ied) - High-risk	(follow-up 12 months; Bet	ter indicate	ed by lower	values)		
1		- ,	no serious inconsistency	no serious indirectness	serious ²	none	157	79	-	MD 1.2 lower (2.43 lower to 0.03 higher)	⊕OOO VERY LOW	CRITICAL
Respond	ler criteria(pa	tients wit	h > 30% improv	 ement in pain)<	4 months (foli	ow-up <4 mont	hs)					
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	32/67 (47.8%)	7/33 (21.2%)	RR 2.25 (1.11 to 4.55)	265 more per 1000 (from 23 more to 753 more)	⊕OOO VERY LOW	IMPORTAN'

Respond	er criteria(pa	itients wit	:h > 30% improv	ement in pain-S	TRATIFIED)< 4	months - low risk	(follow-up <4 months)					
4		1			. 2	T	4/45	4/4.4	DD 0 00	T 00 5 4000		IN ADODE A NE
	randomised	very	no serious	no serious	serious ²	none	4/15	4/14	RR 0.93	20 fewer per 1000		IMPORTAN1
	trials	serious ¹	inconsistency	indirectness			(26.7%)	(28.6%)	(0.29 to 3.03)	(from 203 fewer to 580 more)	VERY LOW	
									3.03)	580 more)		
Respond	er criteria(pa	itients wit	h > 30% improv	ement in pain-S	TRATIFIED)< 4	months - medium	risk (follow-up <4 mor	iths)				
	randomised	verv	no serious	no serious	very serious ³	none	20/31	2/12	RR 3.87	478 more per 1000	⊕OOO	IMPORTAN ⁻
	trials	serious ¹	inconsistency	indirectness	, , , , , , , , , , , , , , , , , , , ,		(64.5%)	(16.7%)	(1.06 to	(from 10 more to	VERY LOW	
			,				(* ***)	(14.09)	1000 more)		
enond	er criteria(na	tionts wit	h > 30% improv	ement in nain-S	TRATIFIED)< 4	months - high ris	k (follow-up <4 months	\ \				
езропи	er criteria(pa	itients wit	.ii > 30 /6 iiiipi 0 v	ement in pain-o		months - mgm ns	k (lollow-up <4 illollillis	,				
	randomised	very	no serious	no serious	serious ²	none	8/21	1/7	RR 2.67	239 more per 1000	⊕OOO	IMPORTAN [*]
	trials	serious1	inconsistency	indirectness			(38.1%)	(14.3%)	(0.4 to	(from 86 fewer to	VERY LOW	
			,				, ,		17.74)	1000 more)		
Respond	er criteria(pa	tients wit	h > 30% improv	ement in function	on)< 4 months	follow-up <4 mon	ths)					
	randomised	verv	no serious	no serious	serious ²	none	41/67	11/33	RR 1.84	280 more per 1000	⊕000	IMPORTANT
	trials	serious ¹	inconsistency	indirectness	3011003	Horic	(61.2%)	(33.3%)	(1.09 to	(from 30 more to	VERY LOW	IIVII OITTAIN
			-				, ,	, ,	3.08)	693 more)	VEIXI LOW	
espond	er criteria(%	age of pa	itients with > 30	% improvement	in ODI-STRATI	FIEDI)< 4 months	- low risk (follow-up <4	months)				
	randomised	very	no serious	no serious	serious ²	none	8/15	6/14	RR 1.24	103 more per 1000	⊕OOO	IMPORTAN ⁻
	trials	serious1	inconsistency	indirectness			(53.3%)	(42.9%)	(0.58 to	(from 180 fewer to	VERY LOW	
									2.68)	720 more)		
espond	er criteria(%	age of pa	tients with > 30	 % improvement	in ODI-STRATI	FIEDI)< 4 months	- medium risk (follow-ւ	p <4 mont	ths)			
					1							
	randomised	very	no serious	no serious	very serious ²	none	22/31	2/12	RR 4.26	543 more per 1000	\oplus OOO	IMPORTAN [*]
	trials	serious1	inconsistency	indirectness			(71%)	(16.7%)	(1.18 to	(from 30 more to	VERY LOW	
									15.39)	1000 more)		
espond	er criteria(%	age of pa	tients with > 30	│ % improvement	in ODI-STRATI	FIEDI)< 4 months	- high risk (follow-up <	4 months)				
	randomias d	l.on.	no oprious	no porious	aariaua ²	hana	11/21	2/7	DD 1 22	04 mars par 1000	0000	IMPODTANI
	randomised	very	no serious	no serious	serious ²	none	11/21	3/7	RR 1.22	94 more per 1000	\oplus OOO	IMPORTANT
									(0.47 to	(from 227 fewer to		

trials	serious1	inconsistency	indirectness		(52.4%)	(42.9%)	3.15)	921 more)	VERY LOW	
		_								

Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 33: Clinical evidence profile: STarT Back classification versus no risk tool classification (IMPaCT cohort)

			Quality ass	essment			No of	patients		Effect	Quality	Importan
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	STarT Back Group	Usual Care (IMPaCT)	Relative (95% CI)	Absolute	,	е
QoL (SF-1	 12, PCS,0-100)	4 months	- 1 year (follow-up	6 months; Bette	r indicated by lo	wer values)						
I	observational studies	very serious	no serious inconsistency	no serious indirectness	no serious imprecision	none	554	368	-	MD 0.2 lower (2 lower to 1.6 higher)	⊕000 VERY LOW	CRITICAL
QoL (SF-1	12, MCS,0-100) >	•4 months	- 1 year (follow-up	6 months; Bette	r indicated by lo	wer values)						
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision ²	none	554	368	-	MD 0.2 lower (2.05 lower to 1.65 higher)	⊕OOO VERY LOW	CRITICAL
Pain(VAS	,0-10)>4 months	s - 1 year (follow-up 6 months	s; Better indicate	d by lower value	es)						
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	554	368	-	MD 0.2 lower (0.59 lower to 0.19 higher)	⊕OOO VERY LOW	CRITICAL
Function(RMDQ,0-24)>4	months - 1	year (follow-up 6	months; Better in	ndicated by lowe	r values)						
	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	554	368	-	MD 0.5 lower (1.27 lower to 0.27 higher)	⊕000 VERY LOW	CRITICAL

² Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

³ Downgraded by 1 or 2 increments because of Heterogeneity, I2=50%, p=0.04, unexplained by subgroup analysis.

	observational	very	no serious	no serious	no serious	none	554	368	-	MD 0.2 lower (0.8	\oplus OOO	CRITICA
	studies	serious ¹	inconsistency	indirectness	imprecision					lower to 0.4 higher)	VERY LOW	
hol	ogical Distress (HADS, dep	pression subscal	e, 0-21) >4 month	ıs - 1 year (follow	-up 6 months; I	Better indicated b	y lower valu	es)			
	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	554	368	-	MD 0.4 lower (0.91 lower to 0.11 higher)	⊕000 VERY LOW	CRITICA
(EC	0-5D,0-1) ≤4 mont	ths(stratifi	ed) - Low Risk (fo	ollow-up 2 month	s; Better indicate	ed by lower valu	ies)					•
	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision ²	none	554	368	-	MD 0.01 higher (0.03 lower to 0.04 higher)	⊕000 VERY LOW	CRITICA
(EC	0-5D,0-1) ≤4 mont	ths(stratifi	ed) - Medium risk	(follow-up 2 mo	nths; Better indic	cated by lower	values)					
	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision ²	none	554	368	-	MD 0.02 lower (0.06 lower to 0.02 higher)	⊕000 VERY LOW	CRITICA
(EC	n-5D,0-1) ≤4 mont	ths(stratifi	ed) - High risk (fo	ollow-up 2 month	s; Better indicate	ed by lower valu	es)					
	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision ²	none	554	368	-	MD 0.06 higher (0.01 to 0.12 higher)	⊕OOO VERY LOW	CRITICA
(EC	1-5D,0-1) >4 mon	ths - 1 yea	r(stratified) - Low	Risk (follow-up	6 months; Better	indicated by lo	wer values)					
	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision ²	none	554	368	-	MD 0 higher (0.03 lower to 0.04 higher)	⊕000 VERY LOW	CRITICA

1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision ²	none	554	368	-	MD 0.01 higher (0.03 lower to 0.04 higher)	⊕OOO VERY LOW	CRITICAL
QoL (EC	Q-5D,0-1) >4 mont	hs - 1 year	r(stratified) - High	n risk (follow-up 6	6 months; Better	indicated by lo	ower values)					
1	observational studies	very serious ²	no serious inconsistency	no serious indirectness	no serious imprecision ²	none	554	368	-	MD 0.07 higher (0.02 to 0.12 higher)	⊕OOO VERY LOW	
QoL (SF	F-12, PCS,0-100) >	•4 months	- 1 year(stratified	l) - Low Risk (foll	ow-up 6 months;	Better indicat	ed by lower values	s)				
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision ²	none	214	136	-	MD 0.4 higher (2.98 lower to 3.78 higher)	⊕000 VERY LOW	CRITICAL
QoL (SF	-12, PCS,0-100) >	4 months	- 1 year(stratified	l) - Medium risk (follow-up 6 mont	hs; Better indi	cated by lower val	ues)				
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision ²	none	232	151	-	MD 1.7 lower (4.39 lower to 0.99 higher)	⊕000 VERY LOW	CRITICAL
QoL (SF	F-12, PCS,0-100) >	•4 months	- 1 year(stratified	l) - High risk (foll	ow-up 6 months;	Better indicat	ed by lower values	5)				
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	108	81	-	MD 3.8 higher (0.19 lower to 7.79 higher)	⊕OOO VERY LOW	CRITICAL
QoL (SF	-12,MCS,0-100) >	4 months	- 1 year(stratified	l) - Low Risk (Bet	ter indicated by I	ower values)						
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision ²	none	214	136	-	MD 0.9 lower (3.87 lower to 2.07 higher)	⊕OOO VERY LOW	CRITICAL
QoL (SF	-12,MCS,0-100) >	4 months	- 1 year(stratified	l) - Medium risk (Better indicated b	y lower value	s)		1			
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision ²	none	232	151	-	MD 0.8 higher (1.95 lower to 3.55 higher)	⊕OOO VERY	CRITICAL

Low back pain and sciatica in over 16s GRADE tables

											LOW	
QoL (SI		4 months	- 1 year(stratified	d) - High risk (foll	ow-up 6 months	; Better indicate	ed by lower values	s)				
•	, , ,		· ·	, , ,	•	•		•				
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	108	81	-	MD 1.6 higher (2.78 lower to 5.98 higher)	⊕OOO VERY	CRITICAL
											LOW	
Pain(V <i>A</i>	S,0-10)>4 months	s - 1 year(s	stratified) - Low F	Risk (follow-up 6	months; Better i	ndicated by low	er values)					
1	observational	very	no serious	no serious	no serious	none	214	136	T -	MD 0.2 higher (0.43	⊕000	CRITICAL
	studies	serious ¹	inconsistency	indirectness	imprecision					lower to 0.83 higher)	VERY	
											LOW	
Pain(VA	S,0-10)>4 months	s - 1 year(s	stratified) - Mediu	ım risk (follow-up	o 6 months; Bett	er indicated by	lower values)					
1	observational	luon.	no serious	no serious	no serious	none	232	151	1	MD 0.1 lower (0.72	⊕000	CRITICAL
l	studies	very serious ¹	inconsistency	indirectness	imprecision	none	232	151	-	lower to 0.52 higher)	VERY	CRITICAL
	studies	Serious	inconsistency	indirectiless	Imprecision					lower to 0.32 higher)	LOW	
											LOW	
Pain(V <i>A</i>	S,0-10)>4 months	s - 1 year(s	stratified) - High	risk (follow-up 6;	Better indicated	by lower values	s)					
1	observational	very	no serious	no serious	serious ²	none	108	81	T -	MD 1 lower (1.84 to	⊕OOO	CRITICAL
	studies	serious ¹	inconsistency	indirectness						0.16 lower)	VERY	
											LOW	
Functio	n(RMDQ,0-24)>4	months - 1	year (stratified)	- Low Risk (follow	w-up 6 months;	Better indicated	by lower values)			I.		
1	observational	very	no serious	no serious	no serious	none	214	136	Τ.	MD 0 higher (1.15	⊕000	CRITICAL
•	studies	serious ²	inconsistency	indirectness	imprecision	none	217	100		lower to 1.15 higher)	VERY	OTTITIO/ LE
			,								LOW	
Functio	n(RMDQ,0-24)>4	months - 1	vear (stratified)	- Medium risk (fo	ollow-up 6 month	ns; Better indica	ted by lower valu	es)				
	,		, , ,			,		,				
1	observational	very	no serious	no serious	no serious	none	232	151	-	MD 0.1 lower (1.37	\oplus OOO	CRITICAL
	studies	serious ²	inconsistency	indirectness	imprecision					lower to 1.17 higher)	VERY	
		1									LOW	ĺ
							1				_	

Low back pain and sciatica in over 16s GRADE tables

	observational studies	very serious ²	no serious inconsistency	no serious indirectness	serious ²	none	108	81	-	MD 2.5 lower (4.3 to 0.7 lower)	⊕000 VERY	CF
											LOW	
olog	gical Distress (HADS, anx	ciety subscale, 0-	21)>4 months - 1	year(stratified)	- Low Risk (foll	ow-up 6 months;	Better indica	ted by low	ver values)		
	observational	very	no serious	no serious	no serious	none	214	136	-	MD 0.1 higher (0.79	⊕000	CI
	studies	serious ²	inconsistency	indirectness	imprecision					lower to 0.99 higher)	VERY LOW	
olog	gical Distress (HADS, anx	iety subscale, 0-	21)>4 months - 1	year(stratified)	- Medium risk (follow-up 06 mon	ths; Better in	dicated by	/ lower values)		
	observational	very	no serious	no serious	no serious	none	232	151	-	MD 0.2 lower (0.98	⊕000	CI
	studies	serious ¹	inconsistency	indirectness	imprecision					lower to 0.58 higher)	VERY LOW	
olog	gical Distress (HADS, anx	riety subscale, 0-	21)>4 months - 1	year(stratified)	- High risk (foll	ow-up 6 months;	Better indica	ted by low	ver values)		
	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	108	81	-	MD 0.6 lower (2.05 lower to 0.85 higher)	⊕000 VERY	С
	studies	Serious	inconsistency	munectness	Imprecision					lower to 0.65 fligher)	LOW	
olog	gical Distress (HADS, dep	ression subscale	e, 0-21)>4 month	s - 1 year(stratifi	ed) - Low Risk	(follow-up 6 mont	ths; Better in	dicated by	lower values)		
	observational	very	no serious	no serious	no serious	none	214	136	-	MD 0.2 lower (1.06	⊕OOO	CI
	studies	serious ¹	inconsistency	indirectness	imprecision					lower to 0.66 higher)	VERY LOW	
olog	gical Distress (HADS, dep	pression subscale	e, 0-21)>4 month	s - 1 year(stratifi	ed) - Medium ri	sk (follow-up mea	an 6 months;	Better inc	 licated by lower values	s)	
	observational	very	no serious	no serious	no serious	none	232	151	-	MD 0 higher (0.68	⊕000	С
	studies	serious ¹	inconsistency	indirectness	imprecision					lower to 0.68 higher)	VERY LOW	

Low		observational studies	serious ¹		no serious indirectness ²	serious ²	none	108	81	-	MD 1.5 lower (2.66 to 0.34 lower)	VERY	CRITICAL
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Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

J.3 Imaging

Table 34: Clinical evidence profile: Imaging versus No imaging for Low back pain and/or sciatica (RCTs)

			Quality as	sessment			No of pa	atients		Effect	Quality	Importance			
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	lmaging	Control	Relative (95% CI)	Absolute	Š				
Health-rela	ated quality of	f life (SF-3	6 bodily pain, 0-10	0) ≤ 4 months (fol	low-up 6 weeks;	range of scores:	0-100; Be	tter indi	cated by highe	r values)					
	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	57	67	-	MD 0 higher (8.31 lower to 8.31 higher)	⊕⊕OO LOW	CRITICAL			
Health-rela	ated quality of	f life (SF-3	6 general health pe	erception, 0-100)	≤ 4 months (folio	w-up 6 weeks; ran	ige of sco	ores: 0-1	00; Better indi	cated by higher values)					
	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness		none	55	65	-	MD 2 higher (6.31 lower to 10.31 higher)	⊕⊕OO LOW	CRITICAL			
Health-rela															
-		very seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	57	66	-	MD 8 higher (0.93 to 15.07 higher)	⊕OOO VERY LOW	CRITICAL			
Health-rela	ated quality of	f life (SF-3	6 role-physical fun	ctioning, 0-100) ≤	4 months (follo	w-up 6 weeks; ran	ge of sco	res: 0-1	00; Better indic	ated by higher values)					
		- ,	no serious inconsistency	no serious indirectness	no serious imprecision	none	55	64	-	MD 4 lower (19.31 lower to 11.31 higher)	⊕⊕OO LOW	CRITICAL			
Health-rela	ated quality of	f life (SF-3	6 social functionin	g, 0-100) ≤ 4 mon	ths (follow-up 6	weeks; range of se	cores: 0-1	100; Bet	ter indicated by	higher values)					

² Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

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i	1	1	ı	T	1	T	1		l .	T		1
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	57	67	-	MD 5 higher (4.78 lower to 14.78 higher)	⊕000 VERY LOW	CRITICAL
Health-rel	ated quality o	f life (SF-3	6 mental health, 0-	·100) ≤ 4 months (follow-up 6 weel	ks; range of scores	s: 0-100;	Better in	ndicated by hig	her values)		
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	57	66	-	MD 9 higher (3.46 to 14.54 higher)	⊕OOO VERY LOW	CRITICAL
Health-rel	ated quality o	f life (SF-3	6 physical function	ning, 0-100) ≤ 4 m	onths (follow-up	6 weeks; range of	scores:	0-100; E	Better indicated	by higher values)		
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	56	65	-	MD 2 higher (6.31 lower to 10.31 higher)	⊕⊕OO LOW	CRITICAL
Health-rel	ated quality o	f life (SF-3	6 role-emotional fu	unctioning, 0-100)	≤ 4 months (foll	ow-up 6 weeks; ra	nge of s	cores: 0	-100; Better ind	icated by higher values)		
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	54	64	-	MD 10 higher (3.85 lower to 23.85 higher)	⊕OOO VERY LOW	CRITICAL
Health-rel	ated quality o	f life (EQ-5	5D VAS, 0-100) ≤ 4	months (follow-u	p 6 weeks; meas	ured with: EQ-5D \	/AS; ran	ge of sc	ores: 0-100; Be	tter indicated by higher va	lues)	
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	57	64	-	MD 7 higher (1.31 lower to 15.31 higher)	⊕OOO VERY LOW	CRITICAL
Pain seve	rity (ALBP sc	ore, 0-100)	>4 months - 1 yea	r (follow-up 24 m	onths; range of s	scores: 0-100; Bett	er indica	ited by l	ower values)			
1	randomised trials	very serious ^a	no serious inconsistency	Serious ^c	no serious imprecision	none	357	335	-	MD 4.2 lower (7.17 to 1.23 lower)	⊕000 VERY LOW	CRITICAL
Function	(RMDQ, 0-24)	≤ 4 month	s (follow-up 6 wee	ks; range of score	es: 0-24; Better i	ndicated by lower	values)	•				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	59	67	-	MD 1 lower (3.08 lower to 1.08 higher)	⊕OOO VERY LOW	CRITICAL
Function	(RMDQ, 0-24)	>4 months	s - 1 year (follow-u	p 1 years; range o	of scores: 0-24; E	Setter indicated by	lower va	lues)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	46	57	-	MD 0.2 higher (1.88 lower to 2.28 higher)	⊕⊕OO LOW	CRITICAL

gical distress	(HADS An	xiety Score, 0-21) :	≤ 4 months (follow	v-up 6 weeks; ra	nge of scores: 0-2	1; Better	indicate	ed by lower valu	ies)		
randomised trials	very seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	57	65	-	MD 0.9 lower (2.43 lower to 0.63 higher)	⊕OOO VERY LOW	CRITICAL
gical distress	(HADS An	xiety Score, 0-21)	>4 months - 1 yea	ır (follow-up 1 ye	ears; range of score	es: 0-21;	Better i	ndicated by low	er values)		
randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	46	53	-	MD 0.4 lower (2.08 lower to 1.28 higher)	⊕⊕OO LOW	CRITICAL
gical distress	(HADS De	pression Score, 0-	21) ≤ 4 months (fe	ollow-up 6 weeks	s; range of scores:	0-21; Be	tter indi	cated by lower	values)		
randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	57	65	-	MD 0.4 lower (1.65 lower to 0.85 higher)	⊕⊕OO LOW	CRITICAL
gical distress	(HADS De	pression Score, 0-	21) >4 months - 1	year (follow-up	1 years; range of s	cores: 0	-21; Bett	ter indicated by	lower values)		
randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	46	56	-	MD 0.3 lower (1.68 lower to 1.08 higher)	⊕⊕OO LOW	CRITICAL
ated quality of	f life (SF-3	6 bodily pain, 0-10	0) >4 months - 1 <u>y</u>	year (range of so	ores: 0-100; Better	indicate	ed by hig	jher values)			
randomised trials	very serious ^a	no serious inconsistency	Serious ^c	Serious ^b	none	403	389	-	MD 3.97 higher (0.36 to 7.59 higher)	⊕OOO VERY LOW	CRITICAL
ated quality of	f life (SF-3	6 mental health, 0-	100) >4 months -	1 year (range of	scores: 0-100; Bet	ter indic	ated by	higher values)			
randomised trials	very serious ^a	Serious ^d	Serious ^c	Serious ^b	none	403	387	-	MD 2.77 higher (0.03 to 5.51 higher)	⊕OOO VERY LOW	CRITICAL
ated quality of	f life (SF-3	6 physical function	ning, 0-100) >4 m	onths - 1 year (ra	inge of scores: 0-1	00; Bette	er indica	ted by higher v	alues)		
randomised trials	very serious ^a	no serious inconsistency	Serious ^e	Serious ^b	none	403	387	-	MD 3.25 higher (0.6 lower to 7.11 higher)	⊕OOO VERY LOW	CRITICAL
ated quality of	f life (SF-3	6 social functionin	g, 0-100) >4 mon	ths - 1 year (rang	je of scores: 0-100	; Better i	ndicated	d by higher valu	ies)		
randomised trials	very serious ^a	no serious inconsistency	Serious ^c	Serious ^b	none	403	391	-	MD 4.25 higher (0.16 to 8.33 higher)	⊕000 VERY	CRITICAL
	randomised trials gical distress randomised trials gical distress randomised trials gical distress randomised trials ated quality of trials	randomised trials very seriousa very very seriousa very very seriousa very very very very very very very very	randomised trials very serious inconsistency gical distress (HADS Anxiety Score, 0-21): randomised trials very serious no serious inconsistency gical distress (HADS Depression Score, 0- randomised very no serious inconsistency gical distress (HADS Depression Score, 0- randomised very no serious inconsistency gical distress (HADS Depression Score, 0- randomised very no serious inconsistency ated quality of life (SF-36 bodily pain, 0-10 randomised very serious inconsistency ated quality of life (SF-36 mental health, 0- randomised very serious Serious inconsistency ated quality of life (SF-36 physical function randomised very serious inconsistency ated quality of life (SF-36 social function in randomised very no serious inconsistency	randomised trials very serious inconsistency indirectness gical distress (HADS Anxiety Score, 0-21) >4 months - 1 year andomised very inconsistency indirectness gical distress (HADS Depression Score, 0-21) ≤ 4 months (for andomised very inconsistency indirectness gical distress (HADS Depression Score, 0-21) ≤ 4 months (for andomised very inconsistency indirectness gical distress (HADS Depression Score, 0-21) >4 months - 1 randomised very inconsistency indirectness gical distress (HADS Depression Score, 0-21) >4 months - 1 randomised very inconsistency indirectness ated quality of life (SF-36 bodily pain, 0-100) >4 months - 1 randomised very inconsistency inconsistency ated quality of life (SF-36 mental health, 0-100) >4 months - 1 randomised very serious inconsistency ated quality of life (SF-36 physical functioning, 0-100) >4 months - 1 randomised very inconsistency inconsistency ated quality of life (SF-36 social functioning, 0-100) >4 months - 1 randomised very inconsistency inconsistency serious Seriou	randomised trials very serious inconsistency indirectness indirectness imprecision no serious indirectness imprecision no serious indirectness imprecision indirectness imprecision serious indirectness imprecision no serious indirectness imprecision no serious indirectness imprecision no serious indirectness imprecision indirectness imprecision no serious i	randomised trials very serious inconsistency indirectness Serious Serious Inone gical distress (HADS Anxiety Score, 0-21) > 4 months - 1 year (follow-up 1 years; range of score indirectness imprecision in one imprecision	randomised trials very serious inconsistency indirectness	randomised trials very serious inconsistency indirectness serious indirectness serious indirectness serious indirectness serious indirectness serious indirectness serious indirectness ind	randomised very serious inconsistency indirectness indire	trials seríous* inconsistency indirectness to 0.63 higher) glical distress (HADS Anxiety Score, 0-21) >4 months -1 year (follow-up 1 years; range of scores: 0-21; Better indicated by lower values) randomised very inconsistency indirectness in one serious indirectness inconsistency indirectness indirectness inconsistency indirectness indirectness inconsistency indirectness indirectness indirectness indirectness inconsistency indirectness indirec	randomised very trials very tr

		1	1			1	1	1	T	<u> </u>	1.014	
											LOW	
Health-rel	ated quality o	f life (SF-3	6 role reported hea	alth transition, 0-1	100) >4 months -	1 year (follow-up 2	24 month	s; range	of scores: 0-1	00; Better indicated by hig	her value	es)
1	randomised trials	very serious ^a	no serious inconsistency	Serious ^c	no serious imprecision	none	357	335	-	MD 1.9 higher (1.77 lower to 5.57 higher)	⊕OOO VERY LOW	CRITICAL
Health-rel	ated quality o	f life (SF-3	6 vitality, 0-100) >4	months - 1 year	(range of scores	s: 0-100; Better indi	icated by	higher	values)			
2	randomised trials	very serious ^a	no serious inconsistency	Serious ^c	Serious ^b	none	403	387	-	MD 3.72 higher (0.54 to 6.9 higher)	⊕OOO VERY LOW	CRITICAL
Health-rel	ated quality o	f life (SF-3	6 general health po	erception, 0-100)	>4 months - 1 ye	ear (range of score	s: 0-100;	Better i	ndicated by hig	her values)		
2	randomised trials	very serious ^a	no serious inconsistency	Serious ^c	Serious ^b	none	402	388	-	MD 1.59 higher (1.76 lower to 4.93 higher)	⊕OOO VERY LOW	CRITICAL
Health-rel	ated quality o	f life (SF-3	6 role-physical fun	ctioning, 0-100) >	4 months - 1 ye	ar (range of scores	s: 0-100; l	Better in	dicated by high	ner values)		
2	randomised trials	very serious ^a	no serious inconsistency	Serious ^c	Serious ^b	none	401	388	-	MD 4.76 higher (1.24 lower to 10.75 higher)	⊕OOO VERY LOW	CRITICAL
Health-rel	ated quality o	f life (SF-3	6 role-emotional fu	ınctioning, 0-100)) >4 months - 1 y	ear (range of score	es: 0-100	; Better	indicated by hi	gher values)		
2	randomised trials	very serious ^a	no serious inconsistency	Serious ^c	Serious ^b	none	401	388	-	MD 5.54 higher (0.51 lower to 11.58 higher)	⊕OOO VERY LOW	CRITICAL
Health-rel	ated quality o	f life (EQ-5	D, 0-1) >4 months	- 1 year (follow-u	p 24 months; rai	nge of scores: 0-1;	Better in	ndicated	by higher value	es)		
1	randomised trials	very serious ^a	no serious inconsistency	Serious ^c	no serious imprecision	none	357	335	-	MD 0.06 higher (0.01 to 0.11 higher)	⊕OOO VERY LOW	CRITICAL
Health-rel	ated quality o	f life (EQ-5	D VAS, 0-100) >4 r	months - 1 year (f	ollow-up 1 years	; measured with: E	Q-5D VA	AS; rang	e of scores: 0-1	00; Better indicated by hi	gher value	es)
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	46	54	-	MD 2 lower (9.06 lower to 5.06 higher)	⊕OOO VERY LOW	CRITICAL

Haalthaa	ro utilization (r	hvoiothar	apy) ≤ 4 months (fallow up 2 mont	·ha)							
1	randomised trials	Serious ^e	no serious inconsistency	no serious indirectness	Serious ^f	none	67/199 (33.7%)	29.1%	RR 1.16 (0.87 to 1.55)	47 more per 1000 (from 38 fewer to 160 more)	⊕⊕OO LOW	
Healthca	re utilisation (a	cupunctu	re) ≤ 4 months (fo	llow-up 3 months	s)							
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ⁹	none	3/199 (1.5%)	3.5%	RR 0.44 (0.11 to 1.67)	20 fewer per 1000 (from 31 fewer to 23 more)	⊕OOO VERY LOW	IMPORTAN
Healthca	re utilisation (d	hiropracti	ic) ≤ 4 months (fo	low-up 3 months	s)							
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ⁹	none	4/199 (2%)	3%	RR 0.68 (0.19 to 2.37)	10 fewer per 1000 (from 24 fewer to 41 more)	⊕000 VERY LOW	IMPORTAN'
Healthca	re utilisation (h	nospital ad	lmission) ≤ 4 mon	ths (follow-up 3 i	months)							
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness		none	0/199 (0%)	0%	-	-		IMPORTAN ⁻
Healthca	re utilisation (d	steopathy	/) ≤ 4 months (foll	ow-up 3 months)								
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^g	none	7/199 (3.5%)	4.4%	RR 0.79 (0.3 to 2.09)	9 fewer per 1000 (from 31 fewer to 48 more)	⊕000 VERY LOW	IMPORTAN
Healthca	re utilisation (d	outpatient	attendance) ≤ 4 m	onths (follow-up	3 months)							
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^g	none	6/199 (3%)	3.5%	RR 0.87 (0.3 to 2.56)	5 fewer per 1000 (from 24 fewer to 55 more)	⊕000 VERY LOW	IMPORTAN'
Healthca	re utilisation (d	over the co	ounter drug) ≤ 4 m	onths (follow-up	3 months)							
1	randomised trials	Serious ^e	no serious inconsistency	no serious indirectness	Serious ^b	none	68/199 (34.2%)	33%	RR 1.04 (0.79 to 1.36)	13 more per 1000 (from 69 fewer to 119 more)	⊕⊕OO LOW	IMPORTAN'
Healthca	re utilisation (p	rescribed	drug) ≤ 4 months	(follow-up 3 mo	nths)							
1	randomised trials	Serious ^e	no serious inconsistency	no serious indirectness	Serious ^b	none	63/199 (31.7%)	29.1%	RR 1.09 (0.81 to 1.47)	26 more per 1000 (from 55 fewer to 137 more)	⊕⊕OO LOW	IMPORTAN'

Healthca	re utilisation (r	referral to	nhysiotheranist or	other health prof	essional) < 4 mo	onths (follow-up 6	weeks)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^g	none	22/69 (31.9%)	28.2%	RR 1.13 (0.68 to 1.88)	37 more per 1000 (from 90 fewer to 248 more)	⊕000 VERY LOW	IMPORTAN
Healthca	re utilisation (s	subsequer	nt doctor consultat	ion for back pain)	≤ 4 months							
2	randomised trials	Serious ^e	very serious ⁶	no serious indirectness	no serious imprecision	none	129/268 (48.1%)	33.1%	RR 1.53 (1.24 to 1.9)	175 more per 1000 (from 79 more to 298 more)	⊕000 VERY LOW	IMPORTAN
Healthca	re utilisation (d	outpatient	consultation) >4 m	nonths - 1 year			•					
2	randomised trials	very serious ^a	no serious inconsistency	Serious	serious ²	none	346/588 (58.8%)	37%	RR 1.24 (1.14 to 1.35)	89 more per 1000 (from 52 more to 130 more)	⊕000 VERY LOW	IMPORTAN
Healthca	re utilisation (p	ohysiother	apy) >4 months - 1	l year								
2	randomised trials	very serious ^a	no serious inconsistency	Serious ^c	no serious imprecision	none	279/588 (47.4%)	36.7%	RR 1.07 (0.95 to 1.19)	26 more per 1000 (from 18 fewer to 70 more)	⊕000 VERY LOW	IMPORTAN
Healthca	re utilisation (a	cupunctu	re) >4 months - 1 y	/ear (follow-up 9 r	nonths)	1						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ⁹	none	1/195 (0.51%)	1%	RR 0.51 (0.05 to 5.58)	5 fewer per 1000 (from 9 fewer to 46 more)	⊕000 VERY LOW	IMPORTAN
Healthca	re utilisation (p	orimary ca	re consultation) >4	1 months - 1 year	(follow-up 24 m	onths)						
1	randomised trials	Serious ^a	no serious inconsistency	Serious ^c	no serious imprecision	none	261/369 (70.7%)	70.1%	RR 1.01 (0.92 to 1.11)	7 more per 1000 (from 56 fewer to 77 more)	⊕⊕OO LOW	IMPORTAN
Healthca	re utilisation (s	subsequer	nt doctor consultat	ion for back pain	>4 months - 1 y	/ear						
2	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	64/264 (24.2%)	31.5%	RR 0.87 (0.66 to 1.16)	41 fewer per 1000 (from 107 fewer to 50 more)	⊕000 VERY LOW	IMPORTAN
Healthca	re utilisation (r	eferral to	physiotherapist or	other health prof	essional) >4 mo	nths - 1 year (follo	w-up 1 ye	ears)				

1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ⁹	none	31/69 (44.9%)		RR 0.97 (0.67 to 1.39)	14 fewer per 1000 (from 153 fewer to 181 more)	⊕OOO VERY LOW	IMPORTANT
Healthca	re utilisation (d	chiropracti	c) >4 months - 1 y	/ear (follow-up 9 r	nonths)							
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ⁹	none	6/195 (3.1%)	2.5%	RR 1.22 (0.38 to 3.95)	6 more per 1000 (from 16 fewer to 74 more)	⊕000 VERY LOW	IMPORTANT
Healthca	re utilisation (I	nospital ad	lmission) >4 mont	ths - 1 year		•	,		•			•
2	randomised trials	very serious ^a	no serious inconsistency	Serious ^c	Serious ^b	none	33/588 (5.6%)	3.3%	RR 1.25 (0.77 to 2.05)	8 more per 1000 (from 8 fewer to 35 more)	⊕000 VERY LOW	IMPORTANT
Healthca	re utilisation (d	osteopathy	v) >4 months - 1 y	ear (follow-up 9 m	onths)	•	,		•			
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^g	none	6/195 (3.1%)	3.5%	RR 0.87 (0.3 to 2.56)	5 fewer per 1000 (from 24 fewer to 55 more)	⊕000 VERY LOW	IMPORTANT
Healthca	re utilisation (d	over the co	ounter drug) >4 m	onths - 1 year (fol	low-up 9 months							
1	randomised trials	Seriouse	no serious inconsistency	no serious indirectness	Serious ^b	none	69/195 (35.4%)	28.6%	RR 1.24 (0.92 to 1.65)	69 more per 1000 (from 23 fewer to 186 more)	⊕⊕OO LOW	IMPORTANT
Healthca	re utilisation (orescribed	drug) >4 months	- 1 year (follow-u	o 9 months)							
1	randomised trials	Serious ^e	no serious inconsistency	no serious indirectness	Serious ^b	none	56/195 (28.7%)	24.6%	RR 1.17 (0.84 to 1.62)	42 more per 1000 (from 39 fewer to 153 more)	⊕⊕OO LOW	IMPORTANT
Healthca	re utilisation (CT imaging	g) >4 months - 1 y	ear (follow-up 24	months)	·						
1	randomised trials	very serious ^a	no serious inconsistency	Serious ^c	Serious ^b	none	29/393 (7.4%)	5.1%	RR 1.44 (0.83 to 2.49)	22 more per 1000 (from 9 fewer to 76 more)	⊕000 VERY LOW	IMPORTANT
Healthca	re utilisation (i	maging at	least once) >4 mo	onths - 1 year (foll	ow-up 24 month	s)			,			•
1	randomised trials	very serious ^a	no serious inconsistency	Serious ^c	no serious imprecision	none	353/393 (89.8%)		RR 3.04 (2.6 to 3.55)	604 more per 1000 (from 474 more to 755 more)	⊕000 VERY LOW	IMPORTANT

Healthcar	e utilisation (i	njection) >	4 months - 1 year	(follow-up 24 mo	nths)			ı	T		T	
1	randomised trials	very serious ^a	no serious inconsistency	Serious ^c	Serious ^b	none	70/393 (17.8%)	19.5%	RR 0.91 (0.68 to 1.22)	18 fewer per 1000 (from 62 fewer to 43 more)	⊕000 VERY LOW	IMPORTAN
Healthcar	e utilisation (N	/IRI imagir	ng) >4 months - 1 y	ear (follow-up 24	months)							
1	randomised trials	very serious ^a	no serious inconsistency	Serious ^c	no serious imprecision	none	324/393 (82.4%)	24.4%	RR 3.38 (2.82 to 4.04)	581 more per 1000 (from 444 more to 742 more)	⊕000 VERY LOW	IMPORTAN
Healthcar	e utilisation (s	surgery) >4	4 months - 1 year (follow-up 24 mon	ths)							
1	randomised trials	very serious ^a	no serious inconsistency	Serious ^c	Serious ^b	none	27/393 (6.9%)	5.1%	RR 1.34 (0.76 to 2.34)	17 more per 1000 (from 12 fewer to 68 more)	⊕000 VERY LOW	IMPORTAN
Healthcar	e utilisation (e	quipment	: back support) ≤ ₄	l months (follow-	up 3 months)							
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ⁹	none	4/199 (2%)	3.9%	RR 0.51 (0.16 to 1.67)	19 fewer per 1000 (from 33 fewer to 26 more)	⊕000 VERY LOW	IMPORTAN
Healthcar	e utilisation (c	lay-case ti	reatment) ≤ 4 mon	ths (follow-up 3 m	nonths)							
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	0/199 (0%)	0%	-	-	⊕000 VERY LOW	IMPORTAN
Healthcar	e utilisation (a	romather	apy) ≤ 4 months (fo	ollow-up 3 months	s)		,					•
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ⁹	none	4/199 (2%)	1.5%	RR 1.36 (0.31 to 6)	5 more per 1000 (from 10 fewer to 75 more)	⊕000 VERY LOW	IMPORTAN
Healthcar	e utilisation (s	ocial serv	rices, reflexology,	massage) ≤ 4 moı	nths (follow-up 3	3 months)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ⁹	none	7/199 (3.5%)	3%	RR 1.19 (0.41 to 3.48)	6 more per 1000 (from 18 fewer to 74 more)	⊕000 VERY LOW	IMPORTAN
Healthcar	e utilisation (c	lay-case tı	reatment) >4 mont	hs - 1 year (follow	v-up 3 months)							

		,	no serious inconsistency	no serious indirectness	very serious ^g	none	1/195 (0.51%)		RR 3.06 (0.1 to 74.69)	-	⊕OOO VERY LOW	IMPORTANT		
Healthcare	Healthcare utilisation (aromatherapy) >4 months - 1 year (follow-up 3 months)													
		,	no serious inconsistency	no serious indirectness	very serious ⁹	none	5/195 (2.6%)	0.5%	RR 5.10 (0.6 to 43.28)	20 more per 1000 (from 2 fewer to 211 more)	⊕OOO VERY LOW	IMPORTANT		
Healthcare	Healthcare utilisation (equipment: back support) >4 months - 1 year (follow-up 3 months)													
		,	no serious inconsistency	no serious indirectness	very serious ^g	none	11/195 (5.6%)	6%	RR 0.94 (0.42 to 2.07)	4 fewer per 1000 (from 35 fewer to 64 more)	⊕OOO VERY LOW	IMPORTANT		
Healthcare utilisation (social services) >4 months - 1 year (follow-up 3 months)														
		serious ^a	no serious inconsistency	indirectness	very serious ^g	none	3/195 (1.5%)	0%	RR 7.14 (0.37 to 137.38)	-	⊕OOO VERY LOW	IMPORTANT		

a Downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 35: Clinical evidence profile: Imaging versus No imaging for Low back pain and/or sciatica (Cohort studies)

			Quality ass	essment		No of p	oatients		Effect	Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	lmaging	No imaging	Relative (95% CI)	Absolute	,	
Healthcar	Healthcare utilisation (advanced imaging) ≤ 4 months (follow-up 3 months)											
1	observational	very	no serious	no serious	no serious	none	63/782	0.6%	RR 14.64 (7.55	82 more per 1000 (from	⊕000	IMPORTANT

b Downgraded by 1 increment if the confidence interval crossed one MID

c Downgraded by 1 increment because the majority of the evidence included an indirect population

d Heterogeneity, 12=66%, p=0.09. Different imaging techniques used in the 2 studies.

e Downgraded by 1 increment if the majority of the evidence was at high risk of bias

f Heterogeneity, I2=82%, p=0.01

g Downgraded by 2 increments if the confidence interval crossed both MIDs

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	studies	seriousª	inconsistency	indirectness	imprecision		(8.1%)		to 28.38)	39 more to 164 more)	VERY	
											LOW	
Healthca	re utilisation (ne	rve testing	g) ≤ 4 months (fol	ow-up 3 months)				1	,		
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	82/782 (10.5%)	0.3%	RR 31.75 (13.92 to 72.44)	92 more per 1000 (from 39 more to 214 more)	⊕OOO VERY LOW	IMPORTANT
Healthca	re utilisation (inj	ections) ≤	4 months (follow	-up 3 months)								
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	270/782 (34.5%)	1.2%	RR 28.52 (18.62 to 43.68)	330 more per 1000 (from 211 more to 512 more)	⊕000 VERY LOW	IMPORTANT
Healthca	re utilisation (su	rgery) ≤ 4	months (follow-u	p 3 months)								
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	70/782 (9%)	0.3%	RR 32.53 (13.18 to 80.28)	95 more per 1000 (from 37 more to 238 more)	⊕000 VERY LOW	IMPORTANT
Healthca	re utilisation (inj	ections) >	4 months - 1 year	(follow-up 6 mor	nths)							
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	329/782 (42.1%)	1.8%	RR 23.89 (16.78 to 34.01)	412 more per 1000 (from 284 more to 594 more)	⊕000 VERY LOW	IMPORTANT
Healthca	re utilisation (su	rgery) >4 ı	months - 1 year (f	ollow-up 6 month	ns)							
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	113/782 (14.5%)	0.55%	RR 26.26 (13.83 to 49.85)	139 more per 1000 (from 71 more to 269 more)	⊕000 VERY LOW	IMPORTANT
Healthca	re utilisation (ad	vanced im	aging) >4 months	s - 1 year (follow-	up 6 months)							
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	121/782 (15.5%)	0.7%	RR 21.63 (12.28 to 38.08)	144 more per 1000 (from 79 more to 260 more)	⊕OOO VERY LOW	IMPORTANT
Healthca	re utilisation (ref	erral to he	ealthcare professi	onal) ≤ 4 months	(follow-up 6 we	eks)	1		,			
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	40/91 (44%)	23.3%	RR 1.88 (1.39 to 2.56)	205 more per 1000 (from 91 more to 363 more)	⊕OOO VERY LOW	IMPORTANT

Healthca	re utilisation (re	ferral to he	ealthcare profession	onal) >4 months	- 1 year							
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	53/91 (58.2%)	37.4%	RR 1.56 (1.24 to 1.95)	209 more per 1000 (from 90 more to 355 more)	⊕OOO VERY LOW	IMPORTANT
Healthca	re utilisation (ne	erve testing	g) >4 months - 1 ye	ear (follow-up 6 n	nonths)							
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	113/782 (14.5%)	0.5%	RR 29.17 (14.87 to 57.22)	141 more per 1000 (from 69 more to 281 more)	⊕OOO VERY LOW	IMPORTAN ⁻
Healthca	re utilisation (su	ıbsequent	consultation for b	ack pain) ≤ 4 mo	nths (follow-up	6 weeks)						
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	38/91 (41.8%)	29.4%	RR 1.42 (1.06 to 1.91)	123 more per 1000 (from 18 more to 268 more)	⊕OOO VERY LOW	IMPORTAN ⁻
Healthca	re utilisation (su	bsequent	consultation for b	ack pain) >4 mor	nths - 1 year							
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	40/91 (44%)	28.4%	RR 1.55 (1.16 to 2.07)	156 more per 1000 (from 45 more to 304 more)	⊕OOO VERY LOW	IMPORTAN [*]
Health-re	elated quality of	life (SF-36	Bodily pain, 0-100) ≤ 4 months (fol	low-up 6 weeks	measured with: S	SF-36 Boo	dily pain;	range of scores	: 0-100; Better indicated	by highe	er values)
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	73	274	-	MD 7 lower (14.06 lower to 0.06 higher)	⊕000 VERY LOW	CRITICAL
Health-re	elated quality of	life (SF-36	Emotional role, 0-	100) ≤ 4 months	(follow-up 6 wee	eks; measured wit	h: SF-36 ∣	Emotiona	l role; range of	scores: 0-100; Better ind	licated b	y higher
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	70	262	-	MD 3 higher (8.42 lower to 14.42 higher)	⊕OOO VERY LOW	CRITICAL
Health-re	elated quality of	life (SF-36	General health, 0-	100) ≤ 4 months	(follow-up 6 wee	eks; measured wit	h: SF-36	General h	ealth; range of	scores: 0-100; Better ind	licated b	y higher
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	69	263	-	MD 1 higher (3.38 lower to 5.38 higher)	⊕OOO VERY LOW	CRITICAL

Health- values)		life (SF-36	Mental health, 0-	100) ≤ 4 months ((follow-up 6 wee	ks; measured with	n: SF-36 N	lental hea	Ith; range of so	cores: 0-100; Better indic	ated by h	igher
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	73	270	-	MD 3 higher (1.38 lower to 7.38 higher)	⊕000 VERY LOW	CRITICAL
	related quality of values)	life (SF-36	Physical functio	ning, 0-100) ≤ 4 m	nonths (follow-u	p 6 weeks; measur	red with:	SF-36 Phy	sical functionii	ng; range of scores: 0-10	0; Better	indicated b
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	69	265	-	MD 8 lower (15.07 to 0.93 lower)	⊕OOO VERY LOW	CRITICA
Health-	related quality of	life (SF-36	Physical role, 0-	100) ≤ 4 months (follow-up 6 wee	ks; measured with	: SF-36 P	hysical ro	le; range of sco	ores: 0-100; Better indica	ited by hi	gher value
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	70	259	-	MD 8 lower (19.42 lower to 3.42 higher)	⊕000 VERY LOW	CRITICAL
	related quality of values)	life (SF-36	Social functioning	ıg, 0-100) ≤ 4 mor	nths (follow-up 6	weeks; measured	l with: SF	-36 Social	functioning; ra	ange of scores: 0-100; Be	etter indic	cated by
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	74	274	-	MD 5 lower (12.07 lower to 2.07 higher)	⊕OOO VERY LOW	CRITICA
Health-	related quality of	life (SF-36	Vitality, 0-100) ≤	4 months (follow	-up 6 weeks; me	easured with: SF-3	6 Vitality;	range of s	scores: 0-100;	Better indicated by highe	er values)	
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	73	273	-	MD 2 higher (2.38 lower to 6.38 higher)	⊕000 VERY LOW	CRITICA
Health-	related quality of	life (EQ-5E	O VAS, 0-100) ≤ 4	months (follow-u	p 6 weeks; meas	sured with: EQ-5D	VAS; ran	ge of scor	res: 0-100; Bett	er indicated by higher va	alues)	
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	73	270	-	MD 2 lower (6.38 lower to 2.38 higher)	⊕OOO VERY LOW	CRITICAI
Health- values)	•	life (SF-36	Bodily pain, 0-10	0) >4 months - 1	year (follow-up	1 years; measured	with: SF	-36 Bodily	pain; range of	scores: 0-100; Better inc	dicated by	/ higher
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	63	252	-	MD 7 lower (14.06 lower to 0.06 higher)	⊕OOO	CRITICAL

	1			1		1	I				VERY	
											LOW	
Health-re		life (SF-36	Emotional role, 0	-100) >4 months -	· 1 year (follow-เ	ıp 1 years; measur	ed with:	SF-36 Em	otional role; ra	nge of scores: 0-100; Be	tter indica	ated by
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	58	233	-	MD 1.00 higher (9.56 lower to 11.56 higher)	⊕OOO VERY LOW	CRITICAI
Health-re	•	life (SF-36	General health, 0	-100) >4 months -	· 1 year (follow-เ	ıp 1 years; measur	ed with:	SF-36 Ger	neral health; ra	nge of scores: 0-100; Be	tter indica	ated by
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	58	244	-	MD 1 lower (7.19 lower to 5.19 higher)	⊕OOO VERY LOW	CRITICA
Health-re values)	elated quality of	life (SF-36	Mental health, 0-1	00) >4 months -	1 year (follow-uբ	o 1 years; measure	d with: S	F-36 Ment	tal health; rang	e of scores: 0-100; Bette	r indicate	d by highe
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	62	249	-	MD 0 higher (4.37 lower to 4.37 higher)	⊕OOO VERY LOW	CRITICA
	elated quality of d by higher value		Physical function	ing, 0-100) >4 mc	onths - 1 year (fo	llow-up 1 years; m	easured	with: SF-	36 Physical fun	ctioning; range of score	s: 0-100;	Better
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	60	240	-	MD 4.00 lower (11.06 lower to 3.06 higher)	⊕OOO VERY LOW	CRITICA
Health-re	elated quality of	life (SF-36	Physical role, 0-1	00) >4 months - 1	year (follow-up	1 years; measured	d with: S	F-36 Phys	ical role; range	of scores: 0-100; Better	indicated	d by highe
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	59	238	-	MD 8.00 lower (19.43 lower to 3.43 higher)	⊕OOO VERY LOW	CRITICA
	elated quality of er values)	life (SF-36	Social functioning	g, 0-100) >4 mont	hs - 1 year (folio	ow-up 1 years; mea	sured w	ith: SF-36	Social function	ning; range of scores: 0-	100; Bette	er indicate
1	observational	verv	no serious	no serious	no serious	none	63	252	_	MD 4.00 lower (10.2	⊕000	CRITICA

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Health-re	lated quality of I	ife (SF-36	Vitality, 0-100) >4	months - 1 year ((follow-up 1 year	rs; measured with:	SF-36 V	itality; ran	ge of scores: 0	-100; Better indicated by	y higher v	values)
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	62	250	-	MD 3.00 lower (9.19 lower to 3.19 higher)	⊕000 VERY LOW	CRITICAL
Health-re	lated quality of I	ife (EQ-5D	VAS, 0-100) >4 m	onths - 1 year (fo	ollow-up 1 years	; measured with: E	Q-5D VA	S; range	of scores: 0-100); Better indicated by hi	gher valu	es)
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	62	250	-	MD 3.00 lower (7.37 lower to 1.37 higher)	⊕OOO VERY LOW	CRITICAL
Function	disability (RMD)	Q, 0-24) ≤	4 months (follow-u	ıp 6 weeks; meas	sured with: Rola	nd Morris Disabilit	y Questi	onnaire; r	ange of scores	: 0-24; Better indicated b	y lower v	ralues)
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	76	276	-	MD 1.30 higher (0.01 lower to 2.61 higher)	⊕OOO VERY LOW	CRITICAL
Function values)	disability (RMD)	Q, 0-24) >4	1 months - 1 year (follow-up 1 years	s; measured witl	n: Roland Morris D	isability	Questionr	naire; range of	scores: 0-24; Better indi	cated by	lower
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	63	254	-	MD 1.40 higher (0.08 to 2.72 higher)	⊕OOO VERY LOW	CRITICAL
Psycholo	gical distress (F	IADS Anx	iety, 0-21) ≤ 4 mon	ths (follow-up 6 v	weeks; measure	d with: HADS Anxi	iety; ranç	ge of score	es: 0-21; Better	indicated by lower value	es)	
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	71	269	-	MD 0.10 lower (1.08 lower to 0.88 higher)	⊕OOO VERY LOW	CRITICAL
Psycholo	gical distress (H	IADS Anx	iety, 0-21) >4 mont	ths - 1 year (follo	w-up 1 years; m	easured with: HAD	S Anxie	ty; range o	of scores: 0-21;	Better indicated by low	er values)	
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	61	248	-	MD 0.20 lower (1.34 lower to 0.94 higher)	⊕OOO VERY LOW	CRITICAL
Psycholo	gical distress (H	IADS Dep	ression, 0-21) ≤ 4 ı	months (follow-u	p 6 weeks; meas	sured with: HADS I	Depressi	on; range	of scores: 0-21	; Better indicated by lov	ver values	s)
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	72	269	-	MD 0.30 lower (1.28 lower to 0.68 higher)	⊕OOO VERY LOW	CRITICAL

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Psychological distress (H	ADS Depr	ression, 0-21) >4 n	onths - 1 year (fo	ollow-up 1 years	s; measured with: I	HADS De	pression;	range of score	s: 0-21; Better indicated	l by lower	values)
I	- ,	no serious inconsistency		no serious imprecision	none	62	248	-	MD 0.40 lower (1.29 lower to 0.49 higher)	⊕OOO VERY LOW	CRITICAL

a Downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 36: Clinical evidence profile: Imaging versus No imaging or Deferred imaging for Low back pain and/or sciatica (Cohort studies)

			Quality ass	essment				No of patients		Effect	Ouality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Imaging	No imaging or Deferred imaging for Low back pain with/without sciatica	Relative (95% CI)	Absolute	Quanty	importance
Quality o	f life (EuroQuo	l 5D Inde	x, 0-1) ≤ 4 months	s (follow-up 3 n	nonths; measu	red with: EuroQu	ol 5D Inc	lex, 0-1; range of scores: 0-	1; Better in	dicated by higher v	/alues)	
	observational studies		no serious inconsistency		no serious imprecision	none	1523	1523	-	MD 0 higher (0.01 lower to 0.01 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	f life (EuroQuo	I 5D VAS,	, 0-100) ≤ 4 montl	ns (follow-up 3	months; meas	ured with: EuroQ	uol 5D V	AS, 0-100; range of scores:	0-100; Bett	er indicated by hig	her valu	es)
	observational studies		no serious inconsistency	no serious indirectness	no serious imprecision	none	1523	1523	-	MD 0.63 higher (0.72 lower to 1.97 higher)	⊕000 VERY LOW	CRITICAL
Quality o	f life (EuroQuo	l 5D Inde	x, 0-1) >4 months	s - 1 year (follow	/-up 1 years; m	neasured with: Eu	ıroQuol 5	5D Index, 0-1; range of score	es: 0-1; Bet	ter indicated by lo	wer value	es)
	observational studies		no serious inconsistency	no serious indirectness	no serious imprecision	none	1523	1523	1	MD 0.01 higher (0 to 0.02 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	f life (EuroQuo	I 5D VAS,	0-100) >4 month	ıs - 1 year (follo	w-up 1 years;	measured with: E	uroQuol	5D VAS, 0-100; range of sc	ores: 0-100	; Better indicated b	y lower	values)
1	observational studies	Seriousª	Serious ^b		no serious imprecision	none	1523	1523	-	MD 1.33 higher (0.01 lower to 2.66 higher)	⊕000 VERY LOW	CRITICAL

b Downgraded by 1 increment if the confidence interval crossed one MID

Pain sev	verity (Back Pai	n NRS, 0-	10) ≤ 4 months (1	follow-up 3 mor	nths; measured	I with: Back Pain	NRS, 0-1	0; range of scores: 0-10; Be	tter indicat	ed by lower values)	
1	observational studies	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	1523	1523	-	MD 0.09 lower (0.28 lower to 0.1 higher)	⊕OOO VERY LOW	CRITICAL
Pain sev	verity (Leg pain	NRS, 0-10	0) ≤ 4 months (fo	llow-up 3 mont	hs; measured	with: Leg pain NR	S, 0-10; ı	range of scores: 0-10; Bette	r indicated	by lower values)		
1	observational studies	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	1523	1523	-	MD 0.29 lower (0.5 to 0.08 lower)	⊕OOO VERY LOW	CRITICAL
	verity (Brief Pair r values)	n Invento	ry Interference, 0)-10) ≤ 4 months	s (follow-up 3 r	nonths; measured	d with: B	rief Pain Inventory Interfere	nce, 0-10; r	ange of scores: 0-1	0; Bette	r indicated
1	observational studies	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	1523	1523	-	MD 0 higher (0.18 lower to 0.17 higher)	⊕OOO VERY LOW	CRITICAL
Pain sev	verity (Back Pai	n NRS, 0-	10) >4 months -	1 year (follow-u	p 1 years; mea	sured with: Back	Pain NR	S, 0-10; range of scores: 0-1	0; Better ir	dicated by lower v	alues)	
1	observational studies	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	1523	1523	-	MD 0.17 lower (0.36 lower to 0.02 higher)	⊕OOO VERY LOW	CRITICAL
Pain sev	verity (Leg pain	NRS, 0-10	0) >4 months - 1	year (follow-up	1 years; meas	ured with: Leg pa	in NRS, (0-10; range of scores: 0-10;	Better indi	cated by lower valu	ies)	
1	observational studies	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	1523	1523	-	MD 0.23 lower (0.44 to 0.02 lower)	⊕OOO VERY LOW	CRITICAL
	verity (Brief Pair d by lower valu		ry Interference, 0	9-10) >4 months	- 1 year (follow	v-up 1 years; mea	sured w	ith: Brief Pain Inventory Inte	rference, 0	-10; range of score	s: 0-10;	Better
1	observational studies	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	1523	1523	-	MD 0.11 lower (0.29 lower to 0.07 higher)	⊕OOO VERY LOW	CRITICAL
Function	n (RMDQ, 0-24)	≤ 4 month	ns (follow-up 3 m	onths; measur	ed with: RMDC	, 0-24; range of so	cores: 0-	24; Better indicated by lowe	er values)			
1	observational studies	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	1523	1523	-	MD 0.02 higher (0.44 lower to 0.49 higher)	⊕OOO VERY LOW	CRITICAL

Function (RMDQ, 0-24) >4 months - 1 year (follow-up 1 years; measured with: RMDQ, 0-24; range of scores: 0-24; Better indicated by lower values)

1		- 7	no serious inconsistency		no serious imprecision	none	336/336 (100%)	17.8%		821 more per 1000 (from 716 more to 938 more)	IMPORTANT
Healthca	re utilisation (s	urgery) >	4 months - 1 yea	r (follow-up 12	months)						
1		- 7	no serious inconsistency		no serious imprecision	none	67/336 (19.9%)	2.5%	RR 7.94 (5.39 to 11.7)	174 more per 1000 (from 110 more to 268 more)	IMPORTANT

a Downgraded by 1 increment if the majority of the evidence was at high risk of bias

Table 37: Clinical evidence profile: Imaging versus No imaging or Deferred imaging for Low back pain without sciatica (Cohort studies)

			Quality asso	essment			N	o of patients		Effect	Over186	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Imaging	No imaging or Deferred imaging	Relative (95% CI)	Absolute	Quality	Importance
Quality of values)	life (SF-36v2 Ro	le-physica	al, 0-100) >4 month	s - 1 year (follow	-up 1 years; mea	sured with: SF-36	/2 Role-p	hysical, 0-100; rang	ge of sco	ores: 0-100; Better in	dicated b	y higher
	observational studies	very serious ^a		no serious indirectness	Serious ^b	none	121	834	-	MD 7.7 lower (10.16 to 5.24 lower)	⊕OOO VERY LOW	CRITICAL
•	life (SF-36v2 Ph	•	ctioning, 0-100) >4	months - 1 year	(follow-up 1 year	rs; measured with:	SF-36v2	Physical functioni	ng, 0-100	; range of scores: 0	-100; Bett	ter
-	observational studies	very serious ^a		no serious indirectness	Serious ^b	none	121	834	-	MD 7.7 lower (10.09 to 5.31 lower)	⊕OOO VERY LOW	CRITICAL
Pain seve values)	rity (Graded chr	onic pain s	scale, 0-10) >4 mor	nths - 1 year (follo	ow-up 1 years; m	easured with: Gra	ded chro	nic pain scale, 0-10); range o	of scores: 0-10; Bette	er indicat	ed by lower
1	observational	very	no serious	no serious	no serious	none	121	834	-	MD 0.9 higher (0.3	⊕000	CRITICAL

b Heterogeneity, I²=81%, p=0.02

c Downgraded by 2 increments if the majority of evidence was at very high risk of bias

d Downgraded by 1 increment if the confidence interval crossed one MID

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	studies	seriousª	inconsistency	indirectness	imprecision					to 1.5 higher)	VERY LOW	
Function	(RMDQ, 0-24) >4	months -	1 year (follow-up 1	years; measured	with: RMDQ, 0-	24; range of scores	s: 0-24; E	Better indicated by I	ower val	ues)		
	observational studies	- ,		no serious indirectness	Serious ^c	none	121	834	-	MD 4.6 higher (3.25 to 5.95 higher)	⊕000 VERY LOW	CRITICAL

a Downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 38: Clinical evidence profile: Imaging versus Deferred imaging for Low back pain and/or sciatica (Cohort studies)

	Quality assessment							No of patients		Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Imaging	Deferred imaging for Low back pain with/without sciatica	Relative (95% CI)	Absolute	Quality	Importance
Healthca	re utilisation (ir	njections)	≤ 4 months (folio	ow-up 3 months	3)							
	observational studies	- ,	no serious inconsistency	no serious indirectness	serious ^b	none	270/782 (34.5%)	26.5%	RR 1.3 (1.08 to 1.57)	79 more per 1000 (from 21 more to 151 more)	⊕OOO VERY LOW	IMPORTANT
Healthca	re utilisation (a	dvanced	imaging) ≤ 4 mor	nths (follow-up :	3 months)							
	observational studies	,	no serious inconsistency	no serious indirectness	serious ^b	none	63/782 (8.1%)	6.2%	RR 1.31 (0.84 to 2.04)	19 more per 1000 (from 10 fewer to 64 more)	⊕OOO VERY LOW	IMPORTANT
Healthca	re utilisation (n	erve testi	ng) ≤ 4 months (follow-up 3 mor	nths)							
		- ,	no serious inconsistency	no serious indirectness	serious ^b	none	82/782 (10.5%)	7.8%	RR 1.34 (0.91 to 1.98)	27 more per 1000 (from 7 fewer to 76 more)		IMPORTANT
Healthca	re utilisation (s	urgery) ≤	4 months (follow	/-up 3 months)								

b Downgraded by 1 increment if the confidence interval crossed one MID

c Downgraded by 1 increment if the confidence interval crossed one MID

1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	70/782 (9%)	3.1%	RR 2.91 (1.63 to 5.2)	59 more per 1000 (from 20 more to 130 more)	⊕000 VERY LOW	IMPORTANT
Healthca	are utilisation (i	njections)	>4 months - 1 ye	ear (follow-up 6	months)							
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	329/782 (42.1%)	36.2%	RR 1.16 (1 to 1.35)	58 more per 1000 (from 0 more to 127 more)	⊕OOO VERY LOW	IMPORTANT
Healthca	are utilisation (a	dvanced	imaging) >4 mon	ths - 1 year (fol	low-up 6 month	ıs)						
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	121/782 (15.5%)	11.6%	RR 1.34 (0.98 to 1.82)	39 more per 1000 (from 2 fewer to 95 more)		IMPORTANT
Healthca	are utilisation (r	erve test	ing) >4 months -	1 year (follow-u	p 6 months)							
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	113/782 (14.5%)	12.5%	RR 1.15 (0.85 to 1.56)	19 more per 1000 (from 19 fewer to 70 more)	⊕OOO VERY LOW	IMPORTANT
Healthca	are utilisation (s	surgery) >	4 months - 1 yea	r (follow-up 6 m	onths)							
1	observational studies		no serious inconsistency	no serious indirectness	no serious imprecision	none	113/782 (14.5%)	5.7%	RR 2.55 (1.67 to 3.89)	88 more per 1000 (from 38 more to 165 more)	⊕OOO VERY LOW	IMPORTANT

a Downgraded by 2 increments if the majority of the evidence was at very high risk of bias b Downgraded by 1 increment if the confidence interval crossed one MID

Table 39: Clinical evidence profile: Imaging versus No imaging for sciatica (Cohort studies)

			Quality assess	sment			N	o of patients		Effect	Overlife v	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	lmaging	No imaging or	Relative (95% CI)	Absolute	Quality	Importance
Quality of	life (SF-36v2 Phy	sical func	tioning, 0-100) >4 n	nonths - 1 year (f	ollow-up 1 ye	ears; measured wi	th: SF-36	v2 Physical function	ning, 0-10	0; range of scores: 0	-100; Bett	er

indicated	by higher values)										
1	observational studies	very seriousª	no serious inconsistency	no serious indirectness	serious ^b	none	107	164	-	MD 5 lower (7.94 to 2.06 lower)	⊕OOO VERY LOW	CRITICAL
Quality of	life (SF-36v2 Ro	le-physical	l, 0-100) >4 months	- 1 year (measur	ed with: SF-3	36v2 Role-physical	, 0-100; r	ange of scores: 0-10	0; Better	indicated by higher	/alues)	
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	107	164	-	MD 5.4 lower (8.35 to 2.45 lower)	⊕OOO VERY LOW	CRITICAL
Pain seve	rity (Graded chro	nic pain s	cale, 0-10) (follow-	up 1 years; meas	ured with: Gı	aded chronic pain	scale, 0-	10; range of scores:	0-10; Be	tter indicated by lowe	er values))
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	107	164	-	MD 0.8 higher (0.15 to 1.45 higher)	⊕OOO VERY LOW	CRITICAL
Function ((RMDQ, 0-24) >4	months - 1	year (follow-up 1	years; measured	with: Roland	Morris Questionna	aire, 0-24	; range of scores: 0-	24; Bette	er indicated by lower	values)	
1	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	107	164	-	MD 2.3 higher (0.58 to 4.02 higher)	⊕OOO VERY LOW	CRITICAL

a Downgraded by 2 increments if the majority of the evidence was at very high risk of bias

J.4 Self-management

Self-management programmes J.4.1

Table 40: Self-management versus usual care for low back pain with or without sciatica

			Quality ass	sessment			No of patient	s		Effect	Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Self-management versus usual care	Control	Relative (95% CI)	Absolute			
Quality of	uality of life (SF-36 physical health, 0-100) ≤ 4 months (range of scores: 0-100; Better indicated by higher values)												

b Downgraded by 1 increment if the confidence interval crossed one MID

						1	ı			1		
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	25	24	-	MD 27.24 higher (16.41 to 38.07 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life (SF-36 n	nental hea	alth, 0-100) ≤ 4 m	onths (range of	scores: 0-100; I	Better indicated by	y higher values)					
1	randomised trials	very seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	25	24	-	MD 7.49 higher (0.16 to 14.82 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	of life (SF-36 e	nergy do	main, 0-100) > 4 r	nonths (range o	f scores: 0-100	; Better indicated	by higher values)					
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	42	38	-	MD 5.9 higher (4.33 lower to 16.13 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life (SF-36 v	vell-being	domain, 0-100) >	4 months (rang	ge of scores: 0-	100; Better indicat	ed by higher values)					
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	42	38	-	MD 8.5 higher (0.35 to 16.65 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality o	of life (SF-36 g	eneral he	alth domain, 0-10	00) > 4 months (range of scores	s: 0-100; Better inc	licated by higher val	ues)				
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	42	38	-	MD 4.4 lower (11.33 lower to 2.53 higher)	⊕⊕OO LOW	CRITICAL
Pain sev	erity (low bac	k pain, V	AS 0-10) ≤ 4 mont	hs (range of sc	ores: 0-10; Bett	er indicated by lov	ver values)					
2	randomised trials	very serious ^a	Serious ^c	no serious indirectness	no serious imprecision	none	54	52	-	MD 0.16 lower (0.81 lower to 0.49 higher)	⊕OOO VERY LOW	CRITICAL
Pain sev	erity (low bac	k pain, V	AS 0-10) > 4 mont	ths (range of sc	ores: 0-10; Bett	er indicated by lov	ver values)					
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	54	47	-	MD 0.1 lower (1.07 lower to 0.87 higher)	⊕⊕⊕O MODERATE	CRITICAL
Function	(modified vo	n Korff 0-	100) >4 months (range of scores	: 0-100; Better i	ndicated by lower	values)					
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	54	47	-	MD 8.0 lower (19.28 lower to 3.28 higher)	⊕⊕OO LOW	CRITICAL
Function	(number not	working)	>4 months									
1	randomised	very	no serious	no serious	very serious ^b	none	14/217	5.9%	RR 1.09	5 more per 1000	⊕000	CRITICAL
					-							

	trials	serious ^a	inconsistency	indirectness			(6.5%)		(0.51 to 2.29)	(from 29 fewer to 76 more)	VERY LOW	
Function	(RMDQ/ODQ) ≤ 4 mon	ths (Better indica	ted by lower va	lues)							
2	randomised trials	very serious ^a	very serious ^d	no serious indirectness	very serious ^b	none	53	53	-	MD 0.02 lower (0.78 lower to 0.73 higher)	⊕000 VERY LOW	CRITICAL
Function	(RMDQ, 0-24) - 4-12 m	onths (range of s	cores: 0-24; Be	tter indicated b	y lower values)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	190	231	-	MD 1.26 lower (2.18 to 0.34 lower)	⊕⊕OO LOW	CRITICAL
Respond	er criteria (no	pain) ≤ 4	months									
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	46/62 (74.2%)	71.7%	RR 1.04 (0.83 to 1.29)	29 more per 1000 (from 122 fewer to 208 more)	⊕⊕OO LOW	CRITICAL
Respond	er criteria (no	pain) > 4	months									
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	34/59 (57.6%)	64.8%	RR 0.89 (0.66 to 1.19)	71 fewer per 1000 (from 220 fewer to 123 more)	⊕⊕OO LOW	CRITICAL
Healthca	re utilisation	(consulta	tion for back pair) > 4 months				•				
4	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	215/716 (30%)	22.7%	RR 0.86 (0.74 to 1.01)	32 fewer per 1000 (from 59 fewer to 2 more)	⊕000 VERY LOW	IMPORTANT
Healthca	re utilisation	(hospitali:	sation) > 4 month	s	_							
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	11/483 (2.3%)	4.2%	RR 0.54 (0.26 to 1.13)	19 fewer per 1000 (from 31 fewer to 5 more)	⊕OOO VERY LOW	IMPORTANT
Healthca	re utilisation	(physicia	visits for back)	> 4 months (Be	tter indicated b	y lower values)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	190	231	-	MD 0.89 lower (1.63 to 0.15 lower)	⊕⊕OO LOW	IMPORTANT
Healthca	re utilisation	(chiropra	ctor visits for bac	k) > 4 months (Better indicated	i by lower values)						

1	randomised trials	very serious ^a			no serious imprecision	none	190	231		MD 0.52 lower (2.52 lower to 1.47 higher)	⊕⊕OO LOW	IMPORTANT
Healthca	re utilisation	(physical	therapist visits fo	or back) > 4 mon	iths (Better indi	icated by lower va	lues)					
1	randomised trials	very serious ^a	no serious inconsistency		no serious imprecision	none	190	231	-	MD 0.68 lower (2.16 lower to 0.8 higher)	⊕⊕OO LOW	IMPORTANT
Healthca	re utilisation	(hospital	days) > 4 months	(Better indicate	d by lower valu	ies)						
1	randomised trials	very serious ^a		no serious indirectness	no serious imprecision	none	190	231	1	MD 0.24 lower (0.48 lower to 0 higher)	⊕⊕OO LOW	IMPORTANT

a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 41: Self-management versus sham for low back pain with or without sciatica

			VC13u3 31lu111 10	7. 1011 Buck pu								
			Quality as	sessment			No of patients	S		Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Self-management versus sham	Control	Relative (95% CI)	Absolute	Quality	Importance
Pain seve	rity (VAS 0-10) ≤ 4 mont	hs (Better indicat	ed by lower value	es)							
1	randomised trials		no serious inconsistency	no serious indirectness	serious	none	63	68	-	MD 0.6 lower (1.2 lower to 0 higher)	⊕⊕OO LOW	CRITICAL
Pain seve	rity (VAS 0-10) >4 mont	hs (Better indicate	d by lower value	s)							
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	63	68	-	MD 0.4 lower (1 lower to 0.2 higher)	⊕⊕OO LOW	CRITICAL
Disability	(RMDQ 0-24)	≤ 4 month	s (Better indicated	l by lower values	s)							
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	63	68	-	MD 0.9 lower (2.1 lower to 0.3 higher)	⊕⊕⊕O MODERATE	CRITICAL

b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

^c Downgraded by 1 or 2 increments because of heterogeneity, 1²=54%, p=0.14, unexplained by subgroup analysis ^d Downgraded by 2 increments because of heterogeneity, 1²=74%, p=0.05, unexplained by subgroup analysis

Disability	(RMDQ 0-24)	>4 months	s (Better indicated	by lower values								
1	randomised trials				no serious imprecision	none	63	68	-	MD 0.6 lower (1.9 lower to 0.7 higher)	⊕⊕⊕O MODERATE	CRITICAL

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Downgraded by 1 increment if the 95% CI crossed one MID, and downgraded by 2 increments if the 95% CI crossed both MIDs

Table 42: Self-management versus bed rest for low back pain with or without sciatica

			Quality as:	sessment			No of patient	s		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Self-management versus bed rest	Control	Relative (95% CI)	Absolute		
Responde	er outcome (r	no pain) ≤	4 months									
	randomised trials	Serious ^a		no serious indirectness	no serious imprecision	none	46/62 (74.2%)	77.2%	RR 0.96 (0.78 to 1.18)	31 fewer per 1000 (from 170 fewer to 139 more)	⊕⊕⊕O MODERATE	IMPORTANT
Responde	er outcome (r	no pain) >	4 months									
1	randomised trials	Serious ^a		no serious indirectness	very serious ^b	none	34/59 (57.6%)	60.4%	RR 0.95 (0.7 to 1.3)	30 fewer per 1000 (from 181 fewer to 181 more)	⊕OOO VERY LOW	IMPORTANT

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 43: Self-management versus exercise for low back pain with sciatica

			Quality asse	ssment			No of patients			Effect	0	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Self-management versus exercise	Control	Relative (95% CI)	Absolute	Quality	Importance

Pain seve	erity (VAS, 0-10) ≤ 4 mont	ths (range of score	es: 0-10; Better in	dicated by lo	wer values)						
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	40	43	-	MD 0.4 higher (0.65 lower to 1.45 higher)	⊕⊕OO LOW	CRITICAL
Pain seve	erity (VAS, 0-10) >4 mont	hs (range of score	s: 0-10; Better inc	licated by lo	wer values)						
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	40	43	-	MD 1 higher (0.02 lower to 2.02 higher)	⊕⊕OO LOW	CRITICAL
Function	(ODI 0-100) ≤ 4	l months (range of scores: 0	-100; Better indic	ated by lowe	r values)						
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	40	43	-	MD 2 higher (2.52 lower to 6.52 higher)	⊕⊕OO LOW	CRITICAL
Function	(ODI 0-100) >4	months (E	Setter indicated by	lower values)								
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	40	43	-	MD 2 higher (3.02 lower to 7.02 higher)	⊕⊕OO LOW	CRITICAL
Quality o	f life (15-D, 0-1)) ≤ 4 montl	hs (range of score	s: 0-1; Better indi	cated by hig	her values)						
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	40	43	-	MD 0.01 lower (0.04 lower to 0.02 higher)	⊕⊕OO LOW	CRITICAL
Quality o	f life (15-D, 0-1)) >4 month	ns (range of scores	: 0-1; Better indic	ated by high	ner values)						
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	40	43	-	MD 0.02 lower (0.05 lower to 0.01 higher)	⊕⊕OO LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the 95% CI crossed one MID, and downgraded by 2 increments if the 95% CI crossed both MIDs

Table 44: Self-management versus exercise for back pain without sciatica

			Quality ass	sessment			No of patient	s		Effect	Quality	Importance
No of studies							Self-management versus exercise	Control	Relative (95% CI)	Absolute	-	·

Function	(RMDQ. 0-24)	≤ 4 mont	hs (range of score	es: 0-24: Better i	ndicated by low	ver values)						
	randomised	very	no serious	no serious	no serious imprecision	none	63	117	-	MD 0.2 higher (1.3 lower to 1.7 higher)	⊕⊕OO LOW	CRITICAL
Responde	er criteria (>5	0% impro	vement in RMDQ)	≤ 4 months								
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	9/30 (30%)	15/30 (50%)	RR 0.6 (0.31 to 1.15)	200 fewer per 1000 (from 345 fewer to 75 more)	⊕⊕OO LOW	IMPORTANT
Healthcar	e utilisation (medicatio	on use) > 4 month	s								
1	randomised trials		no serious inconsistency	no serious indirectness	very serious ^b	none	17/29 (58.6%)	16/32 (50%)	RR 1.17 (0.74 to 1.86)	85 more per 1000 (from 130 fewer to 430 more)	⊕OOO VERY LOW	IMPORTANT

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 45: Self-management versus massage for low back pain without sciatica

			Quality as	sessment			No of patients	.		Effect	Ovality	Immontonco
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Self-management versus massage	Control	Relative (95% CI)	Absolute	Quanty	Importance
Function	(RMDQ, 0-24)	≤ 4 month	s (range of scores	s: 0-24; Better inc	dicated by lower	values)						
		,	no serious inconsistency	no serious indirectness	Serious ^b	none	83	77	-	MD 2.5 higher (0.65 to 4.35 higher)	⊕OOO VERY LOW	CRITICAL
Function	(RMDQ, 0-24)	> 4 month	s (range of scores	s: 0-24; Better inc	dicated by lower	values)						
		- ,	no serious inconsistency		no serious imprecision	none	83	76	-	MD 0.4 lower (2.23 lower to 1.43 higher)	⊕⊕OO LOW	CRITICAL

Healthcar	re utilisation (provider v	risits) > 4 months (Better indicated	by lower values	3)						
1		- ,	no serious inconsistency	no serious indirectness	no serious imprecision	none	83	76	-	MD 0.5 higher (0.48 lower to 1.48 higher)	⊕⊕OO LOW	IMPORTANT
Healthcar	re utilisation (low back _l	pain medication fil	ls) > 4 months (E	Better indicated	by lower values)						
1		very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	83	76	-	MD 1.5 higher (0.52 lower to 3.52 higher)	⊕⊕OO LOW	IMPORTANT

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 46: Self-management versus yoga for back pain without sciatica

			Quality as:	sessment			No of patien	ts		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Self-management versus yoga	Control	Relative (95% CI) Absolute			
Responde	er criteria (>5	0% impro	vement in RMDQ)	≤ 4 months								
	randomised trials	Serious ^a		no serious indirectness	no serious imprecision	none	9/30 (30%)	69.4%	RR 0.43 (0.24 to 0.78)	396 fewer per 1000 (from 153 fewer to 527 fewer)	⊕⊕⊕O MODERATE	IMPORTANT
Healthcar	e utilisation (Medicatio	on use) > 4 month	s				<u> </u>				
	randomised trials	Seriousª		no serious indirectness	no serious imprecision	none	17/29 (58.6%)	20.6%	RR 2.85 (1.38 to 5.89)	381 more per 1000 (from 78 more to 1000 more)	⊕⊕⊕O MODERATE	IMPORTANT

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 47: Self-management versus acupuncture for low back pain without sciatica

Quality acceptament	No of nationto	Effect	Quality	Importance
Quality assessment	No of patients	Effect	Quanty	Importance

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Self-management versus acupuncture	Control	Relative (95% CI)	Absolute		
Function	(RMDQ, 0-24)	≤ 4 month	ns (range of score	s: 0-24; Better in	dicated by lowe	r values)						
		very seriousª		no serious indirectness	Serious ^b	none	83	89	-	MD 0.9 higher (1.07 lower to 2.87 higher)	⊕OOO VERY LOW	CRITICAL
Function	(RMDQ, 0-24)	> 4 month	ns (range of score	s: 0-24; Better in	dicated by lowe	r values)						
		very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	83	90	-	MD 1.6 lower (3.51 lower to 0.31 higher)	⊕OOO VERY LOW	CRITICAL
Healthcar	e utilisation (¡	provider v	risits) >4 months (Better indicated	by lower values							
1		very serious ^a			no serious imprecision	none	83	90	-	MD 0.4 lower (1.55 lower to 0.75 higher)	⊕⊕OO LOW	IMPORTANT
Healthcar	e utilisation (l	low back p	pain medication fil	ls) > 4 months (E	Better indicated	by lower values)						
1		very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	83	90	-	MD 0.4 lower (3.01 lower to 2.21 higher)	⊕⊕OO LOW	IMPORTANT

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 48: Self-management (bed rest plus exercise) versus usual care for low back pain with or without sciatica

			Quality asse	essment			No of patients	No of patients Effect				Immoutono
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Self-management (bed rest + exercise) versus usual care	Control	Relative (95% CI)	Absolute	Quanty	Importance
Responde	er criteria (No	o pain) ≤ 4	months									
1	randomised	Seriousª	no serious	no serious	Serious ^b	none	47/63	71.7%	RR 1.04	29 more per 1000	⊕⊕00	CRITICAL

	trials		inconsistency	indirectness			(74.6%)		(0.84 to 1.29)	(from 115 fewer to 208 more)	LOW	
Respond	er criteria (No	pain) > 4	months									
1	randomised trials			no serious indirectness	very serious ^b	none	37/60 (61.7%)	64.8%	RR 0.95 (0.72 to 1.26)	32 fewer per 1000 (from 181 fewer to 168 more)	⊕000 VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 49: Self-management (bed rest plus exercise) versus bed rest for low back pain

Quality assessment						No of patients			Effect	Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Self-management (bed rest + exercise) versus bed rest	Control	Relative (95% CI)	Ansoluta		Importance
Responder criteria (No pain) ≤ 4 months												
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	47/63 (74.6%)	77.2%	RR 0.97 (0.79 to 1.18)	23 fewer per 1000 (from 162 fewer to 139 more)	⊕⊕⊕O MODERATE	CRITICAL
Respond	er criteria (No	pain) > 4	4 months									
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	37/60 (61.7%)	60.4%	RR 1.02 (0.76 to 1.37)	12 more per 1000 (from 145 fewer to 223 more)	⊕⊕OO LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 50: Self-management (bed rest plus exercise) versus self-management (exercise) for low back pain with or without sciatica

Quality assessment No of patients Effect Quality Importance

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Self-management (bed rest plus exercise) versus self- management (exercise)		Relative (95% CI)	Absolute		
Respond	ler criteria (N	lo pain) ≤	4 months									
1	randomised trials		no serious inconsistency		no serious imprecision	none	47/63 (74.6%)	74.2%	RR 1.01 (0.82 to 1.24)	7 more per 1000 (from 134 fewer to 178 more)	⊕⊕⊕O MODERATE	CRITICAL
Respond	ler criteria (N	lo pain) >	4 months								•	
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	37/60 (61.7%)	57.6%	RR 1.07 (0.8 to 1.44)	40 more per 1000 (from 115 fewer to 253 more)		CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 51: Self-management programme (exercise plus stretching plus booklet) versus manual therapy combination of techniques (manual mobilisation with manipulation excluded plus thermal plus electrotherapy) for low back pain without sciatica

			Quality ass	essment			ı	No of patients		Effect		
No of studies							Self-management (exercise+ stretching+ booklet)	Manual therapy combination of techniques (manual manipulation excluding mobilisation + thermal+ electrotherapy)	Relative (95% CI)		Quality	Importance
Function	ı (improveme	ent of OD	l) ≤ 4 months (fo	ollow-up mean	1 years; Bet	ter indicated by l	higher values)					
Function (improvement of ODI) ≤ 4 months (follow-up mean 1 years; Better indicated by higher values) 1 randomised trials Serious no serious inconsistency indirectness Serious none 35 33 - MD 1.10 lower (4.99 lower to 2.79 higher)											CRITICAL	
Function	ı (improveme	ent of OD	l) > 4 months (fo	ollow-up mean	1 years; Bet	ter indicated by	higher values)					

1	randomised trials			no serious indirectness	Serious ^b	none	32	32		MD 2.20 lower (6.76 lower to 2.36 higher)	
Healthca	are utilisatior	ı (visits t	o healthcare cer	ntres) (Better in	ndicated by I	ower values)					
	randomised trials			no serious indirectness	Serious ^b	none	32	32	ı	MD 0.30 higher (0.12 lower to 0.72 higher)	IMPORTANT

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 52: Self-management programme (exercise plus stretching plus booklet) versus manipulation therapy (bone-setting) for low back pain without sciatica

			Quality asse	essment			No of patie	ents		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Self-management (exercise+ stretching+ booklet)	Mobilisation (bone-setting)	Relative (95% CI)		Quanty	Importance
Disability	(ODI, 0-100)	≤ 4 month	ns (range of score	es: 0-100; Better	indicated by	y lower values)						
1	randomised trials	-100) ≤ 4 months (range of scores: 0-100; Better indicated by lower ised Serious ^a no serious inconsistency no serious indirectness Serious ^b none		none	35	43	-	MD 2.20 lower (6.52 lower to 2.12 higher)	⊕⊕OO LOW	CRITICAL		
Disability	(ODI, 0-100)	> 4 month	ns (range of score	es: 0-100; Better	r indicated by	y lower values)						
1	randomised trials			no serious indirectness	Serious ^b	none	32	44	-	MD 6.20 lower (10.78 to 1.62 lower)	⊕⊕OO LOW	CRITICAL
Healthcar	re utilisation (visits to l	nealthcare centre	s) (Better indica	ted by lower	values)						

1	randomised trials			no serious indirectness	Serious ^b	none	32	44	-	MD 0.10 higher (0.33 lower to 0.53 higher)		IMPORTANT
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^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Advice to stay active J.4.2

Table 53: Advice to stay active versus bed rest for back pain for low back pain with or without sciatica

			Quality asse	ssment			No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Advice to stay active versus bed rest	Control	Relative (95% CI)	Absolute	Quanty	importance
Function ((RMDQ, 0-24) :	≤ 4 months	(range of scores:	0-24; Better indic	ated by lowe	er values)						
1	randomised trials	- ,		no serious indirectness	Serious ^b	none	14	20	-	MD 2.7 higher (0.72 lower to 6.12 higher)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 54: Advice to stay active versus bed rest for back pain for low back pain without sciatica

			Quality ass	sessment			No of patie	nts		Effect	0174	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Advice to stay active	Bed rest	Relative (95% CI)		Quality	Importance
Days to full	l activity ≤ 4 m	onths (Bett	er indicated by low	er values)								

1				no serious indirectness	no serious imprecision	none	40	40	-	MD 5.23 lower (5.74 to 4.72 lower)	⊕⊕OO LOW	CRITICAL
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a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Bed rest J.4.3

Table 55: Bed rest versus usual care for low back pain with or without sciatica

			Quality asse	essment			No of patie	ents		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Bed rest versus usual care	Control	Relative (95% CI)	Absolute	Quanty	importance
Responde	er criteria (No _l	pain) ≤ 4 n	nonths									
	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	44/57 (77.2%)	71.7%	RR 1.08 (0.87 to 1.33)	57 more per 1000 (from 93 fewer to 237 more)	⊕⊕OO LOW	CRITICAL
Responde	er criteria (No _l	pain) > 4 n	nonths									
1	randomised trials		no serious inconsistency	no serious indirectness	very serious ^b	none	32/53 (60.4%)	64.8%	RR 0.93 (0.69 to 1.25)	45 fewer per 1000 (from 201 fewer to 162 more)	⊕OOO VERY LOW	CRITICAL
Function	(ODI, 0-100) ≤	4 months	(range of scores:	0-100; Better indi	cated by low	er values)		•				
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	67	67	-	MD 3.9 higher (0.1 to 7.7 higher)	⊕⊕OO LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the 95% CI crossed one MID, and downgraded by 2 increments if the 95% CI crossed both MIDs

Table 56: Bed rest versus usual care for low back pain with sciatica

				•								
			Quality as	sessment			No of patier	nts		Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Bed rest versus usual care	Control	Relative (95% CI)		Quality	Importance
Pain sever	ity (back pain	, VAS 0-10)) ≤ 4 months (range	of scores: 0-10;	Better indicated I	by lower values)						
1	randomised trials	very serious ^a			no serious imprecision	none	85	84	-	MD 0.3 lower (1.8 lower to 0.48 higher)	⊕⊕OO LOW	CRITICAL
Pain sever	rity (leg pain) ≤	≤ 4 months	(range of scores: ()-10; Better indica	ated by lower valu	ues)						
1	randomised trials	very seriousª			no serious imprecision	none	85	84	-	MD 2 higher (5.54 lower to 9.54 higher)	⊕⊕OO LOW	CRITICAL
Function (tion (ODI, 0-100) ≤ 4 months (range of scores: 0-100; Better indicated by lower values)											
1	randomised trials	very serious ^a			no serious imprecision	none	85	84	-	MD 0 higher (3.17 lower to 3.17 higher)	⊕⊕OO LOW	CRITICAL

a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Unsupervised exercise J.4.4

Table 57: Unsupervised exercise versus usual care for low back pain without sciatica

			Quality asse	ssment			No of patier	nts		Effect	Quality	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Unsupervised exercise	Usual care	Relative (95% CI)	Absolute	Quality	Importance
Disability (RMDQ, 0-24)	> 4 months	s (range of scores:	0-24; Better indic	cated by lowe	r values)		•			•	

1		very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	51	60	-	MD 1.65 lower (3.62 lower to 0.32 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	life (SF-36 Phy	ysical, 0-1	00) > 4 months (ran	ge of scores: 0-1	00; Better in	dicated by higher v	alues)					
1		very serious ^a	no serious inconsistency		very serious ^b	none	51	60	-	MD 2.08 lower (10.66 lower to 6.44 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	life (SF-36 Me	ntal, 0-100) > 4 months (rang	e of scores: 0-100	; Better indi	cated by higher val	ues)		•			
1		very serious ^a	no serious inconsistency		very serious ^b	none	51	60	-	MD 0.72 lower (7.38 lower to 8.22 higher)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 58: Unsupervised exercise versus usual care for low back pain with or without sciatica

			Quality asse	ssment			No of patients			Effect	Ouglitu	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Unsupervised exercise versus usual care	Control	Relative (95% CI)		Quality	Importance
Function (ODI, 0-100) ≤ 4 months (range of scores: 0-100; Better indicated by lower values)												
	randomised trials			no serious indirectness	Serious ^b	none	52	67	-	MD 2.6 higher (1.6 lower to 6.8 higher)	⊕⊕OO LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the 95% CI crossed one MID, and downgraded by 2 increments if the 95% CI crossed both MIDs

Table 59: Unsupervised exercise versus Alexander technique for low back pain without sciatica

Quality assessment	No of patients	Effect	Quality Importance

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Unsupervised exercise versus Alexander technique	Control	Relative (95% CI)	Absolute		
Quality o	f life (SF-36 P	hysical, 0	-100) > 4 months (range of scores:	0-100; Better i	ndicated by higher	rvalues)					
1	randomised trials	very serious ^a			no serious imprecision	none	102	119	-	MD 9.03 lower (17.09 to 0.96 lower)	⊕⊕OO LOW	CRITICAL
Pain seve	erity (Von Kor	ff, 0-10) >	4 months (range	of scores: 0-10;	Better indicated	by lower values)						
1	randomised trials	very serious ^a			no serious imprecision	none	102	119		MD 0.57 higher (0.32 lower to 1.46 higher)		CRITICAL
Quality o	f life (SF-36 M	lental, 0-1	00) > 4 months (ra	nge of scores: 0	-100; Better ind	licated by higher v	alues)	•				
1	randomised trials	very serious ^a			no serious imprecision	none	102	119	-	MD 3.38 lower (14.34 lower to 7.58 higher)	⊕⊕OO LOW	CRITICAL
Disability	Disability (RMDQ, 0-24) > 4 months (range of scores: 0-24; Better indicated by lower values)											
1	randomised trials	very serious ^a			no serious imprecision	none	102	119		MD 1.15 higher (0.78 lower to 3.07 higher)	⊕⊕OO LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 60: Unsupervised exercise versus exercise for low back pain with or without sciatica

			Quality ass	sessment			No of patients	5		Effect	Quality		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Unsupervised exercise versus exercise	Control	Relative (95% CI)	Absolute	Quality	Importance	
Pain seve	Pain severity (Back pain, VAS 0-10) ≤ 4 months (range of scores: 0-10; Better indicated by lower values)												

	1	1	1		1	1	1			1			
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	57	59	-	MD 1.32 higher (0.36 to 2.28 higher)	⊕⊕⊕O MODERATE	CRITICAL	
Pain sev	erity (Back pa	ain, VAS 0)-10) > 4 months (range of scores	s: 0-10; Better ii	ndicated by lower	values)						
2	randomised trials	very serious ^a	very serious ^b	no serious indirectness	no serious imprecision	none	77	79	1	MD 3.16 higher (2.55 to 3.77 higher)	⊕OOO VERY LOW	CRITICAL	
Number	of pain relaps	ses > 4 mo	onths (Better indi	cated by lower	values)								
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	20	20	1	MD 2.8 higher (1.95 to 3.65 higher)	⊕⊕OO LOW	CRITICAL	
Leg pain	ı ≤ 4 months (range of s	scores: 0-10; Bett	er indicated by	lower values)								
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	57	59	-	MD 1.64 higher (0.55 to 2.73 higher)	⊕⊕⊕O MODERATE	CRITICAL	
Leg pain > 4 months (range of scores: 0-10; Better indicated by lower values)													
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	57	59	-	MD 1.45 higher (0.41 to 2.49 higher)	⊕⊕⊕O MODERATE	CRITICAL	
Function	n (ODI, 0-100)	≤ 4 month	ns (range of score	es: 0-100; Bette	r indicated by lo	ower values)							
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	57	59	-	MD 6.5 higher (1.05 to 11.95 higher)	⊕⊕⊕O MODERATE	CRITICAL	
Function	n (ODI, 0-100)	> 4 month	ns (range of score	es: 0-100; Bette	r indicated by lo	ower values)				•			
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	57	59	-	MD 6.5 higher (0.94 to 12.06 higher)	⊕⊕⊕O MODERATE	CRITICAL	
Return to	o work > 4 mc	onths											
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	very serious ^c	none	40/70 (57.1%)	41/69 (59.4%)	RR 0.96 (0.73 to 1.27)	24 fewer per 1000 (from 160 fewer to 160 more)	⊕000 VERY LOW	CRITICAL	

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 2 increments because of heterogeneity, I² = 97%, p<0.00001 ^c Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

	Спопро	visca c/		nussage for to	w back pain	without sciatics						
			Quality as	sessment			No of patients			Effect	Ouglitu.	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Unsupervised exercise versus massage	Control	Relative (95% CI)	Absolute	Quality	Importance
Quality of	life (SF-36 P	hysical, 0-	100) > 4 months (I	range of scores:	0-100; Better in	dicated by higher	values)					
	randomised trials	- ,		no serious indirectness	no serious imprecision	none	51	64	-	MD 0.63 lower (12.03 lower to 10.77 higher)		CRITICAL
Quality of	life (SF-36 M	ental, 0-10	00) > 4 months (ra	nge of scores: 0	-100; Better indi	cated by higher va	alues)					
1	randomised trials	- ,		no serious indirectness	Serious ^a	none	51	64	-	MD 2.83 higher (8.06 lower to 13.72 higher)	⊕OOO VERY LOW	CRITICAL
Pain (McG	Gill, 0-78) ≤ 4 ı	months (B	etter indicated by	lower values)								
1	randomised trials	- ,		no serious indirectness	Serious ^a	none	12	12	-	MD 2.3 higher (2.31 lower to 6.91 higher)	⊕OOO VERY LOW	CRITICAL
Pain seve	rity (Von Kor	ff, 0-10) >	4 months (range o	of scores: 0-10; E	Better indicated	by lower values)						
	randomised trials	- ,		no serious indirectness	no serious imprecision	none	51	64	-	MD 0.6 lower (1.86 lower to 0.66 higher)	⊕⊕OO LOW	CRITICAL
Function	(RMDQ, 0-24)	> 4 mont	hs (range of score	s: 0-24; Better ir	ndicated by lowe	er values)						
1	randomised trials	- ,		no serious indirectness	Serious ^a	none	51	64	-	MD 1.2 lower (3.9 lower to 1.5 higher)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs ^b Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

J.4.5 Combinations of interventions – self-management adjunct

J.4.6 Low back pain without sciatica

Table 62: self-management (exercise prescription) + postural therapy (Alexander technique -6 lessons) plus versus Postural therapy (Alexander technique) - 6 lessons)

		14.0, 0	163301137									
			Quality ass	sessment			No of patients			Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alexander technique (6 lessons) + self-management (exercise prescription) versus Alexander technique (6 lessons)	Control	Relative (95% CI)	Absolute	Quality	Importance
Qualty o	life (SF-36 p	hysical c	component sumr	nary) >4 month	ns (follow-up 1	years; range of	scores: 0-100; Better indicated by	higher v	/alues)			
1	randomised trials			no serious indirectness	Serious ^b	none	57	58	-	MD 6.49 higher (2.03 lower to 15.01 higher)	⊕⊕OO LOW	CRITICAL
Qualty o	f life (SF-36 n	nental co	mponent summa	ary) >4 months	(follow-up 1 y	ears; range of sc	ores: 0-100; Better indicated by h	igher va	lues)			
1	randomised trials				no serious imprecision	none	57	58	1	MD 3.46 lower (11.41 lower to 4.49 higher)	⊕⊕⊕O MODERATE	CRITICAL
Pain (Vo	n Korff pain s	scale) >4	months (follow-	up 1 years; ran	ge of scores:	0-10; Better indic	ated by lower values)					
1	randomised trials			no serious indirectness	Serious ^b	none	57	58	-	MD 0.64 lower (1.59 lower to 0.31 higher)	⊕⊕OO LOW	CRITICAL
Function	(RMDQ, 0-24	4) >4 mor	nths (follow-up 1	years; range o	of scores: 0-24	; Better indicated	by lower values)					
1	randomised trials			no serious indirectness	Serious ^b	none	57	58	-	MD 1.54 lower (3.44 lower to 0.36 higher)	⊕⊕OO LOW	CRITICAL

Healthca	are utilisation	(primary	care contacts)	>4 months (foll	ow-up 1 years	; Better indicated	by lower values)							
1	randomised trials			no serious indirectness	no serious imprecision	none	57	58	-	MD 0.13 lower (0.45 lower to 0.19 higher)		IMPORTANT		
Healthca	Healthcare utilisation (prescriptions) >4months (follow-up 1 years; Better indicated by lower values)													
1	randomised trials			no serious indirectness	no serious imprecision	none	57	58	-	MD 0.06 lower (0.5 lower to 0.38 higher)	⊕⊕⊕O MODERATE	IMPORTANT		

Table 63: self-management (exercise prescription) + Postural therapy (Alexander technique - 24 lessons) versus Postural therapy (Alexander technique - 6 lessons)

	0 1033											
			Quality as:	sessment			No of patients			Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alexander technique (24 lessons) + self-management (exercise presctiption) versus Alexander technique (6 lessons)	Control	Relative (95% CI)	Absolute	Quality	Importance
Qualty of	ualty of life (SF-36 physical component summary) >4 months (follow-up 1 years; range of scores: 0-100; Better indicated by higher values)											
	randomised trials			no serious indirectness	Serious ^b	none	56	58	-	MD 7.39 higher (1.02 lower to 15.8 higher)	⊕⊕OO LOW	CRITICAL
Qualty of	f life (SF-36 n	nental co	mponent summa	ary) >4 months	(follow-up 1 y	ears; range of so	ores: 0-100; Better indicated by h	igher va	lues)			
	randomised trials				no serious imprecision	none	56	58	,	MD 0.89 higher (6.94 lower to 8.72 higher)	⊕⊕⊕O MODERATE	CRITICAL
Pain (Vo	n Korff pain s	scale) >4	months (follow-	up 1 years; ran	ge of scores:	0-10; Better indic	ated by lower values)					
1	randomised	Serious	no serious	no serious	Serious ^b	none	56	58	-	MD 1.19 lower	⊕⊕00	CRITICAL

^a Downgraded by one increment if the majority of the evidence was at high risk of bias.
^b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs.

	trials		inconsistency	indirectness						(2.13 to 0.25 lower)	LOW			
Function	Function (RMDQ, 0-24) >4 months (follow-up 1 years; range of scores: 0-24; Better indicated by lower values)													
1	randomised trials	Seriousª		no serious indirectness	Serious ^b	none	56	58	1	MD 2.78 lower (4.69 lower to 0.87 higher)	⊕⊕OO LOW	CRITICAL		
Healthca	lealthcare utilisation (primary care contacts) >4 months`` (follow-up 1 years; Better indicated by lower values)													
1	randomised trials				no serious imprecision	none	56	58	-	MD 0.11 higher (0.25 lower to 0.47 higher)		IMPORTANT		
Healthca	Healthcare utilisation (prescriptions) >4 months (follow-up 1 years; Better indicated by lower values)													
1	randomised trials				no serious imprecision	none	56	58	-	MD 0.04 higher (0.51 lower to 0.59 higher)	0000	IMPORTANT		

Table 64: self-management (exercise prescription) + Postural therapy (Alexander technique - 6 lessons) versus Postural therapy (Alexander technique -24 lessons)

	27 1033	•,											
			Quality as:	sessment			No of patients			Effect			
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alexander technique (6 lessons) + self-management (exercise prescription) versus Alexander technique (24 lessons)	Control	Relative (95% CI)	Absolute	Quality	Importance	
Qualty of	f life (SF-36 p	hysical o	component sumr	nary) >4 month	ıs (follow-up 1	years; range of	scores: 0-100; Better indicated by	higher v	/alues)				
	randomised trials				no serious imprecision	none	59	57	-	MD 3.3 lower (11.63 lower to 5.03 higher)	⊕⊕⊕O MODERATE	CRITICAL	
Qualty of	tualty of life (SF-36 mental component summary) >4 months (follow-up 1 years; range of scores: 0-100; Better indicated by higher values)												

^a Downgraded by one increment if the majority of the evidence was at high risk of bias.
^b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs.

1	randomised trials	Seriousª			no serious imprecision	none	57	61	-	MD 3.1 lower (11.42 lower to 5.22 higher)	⊕⊕⊕O MODERATE	CRITICAL
Pain (Vo	n Korff pain	scale) >4	months (follow-	-up 1 years; ran	ge of scores:	0-10; Better indic	ated by lower values)					
1	randomised trials	Seriousª			no serious imprecision	none	57	61	-	MD 0.26 higher (0.68 lower to 1.2 higher)		CRITICAL
Function	n (RMDQ, 0-2	4) > 4 mo	onths (follow-up	1 years; range	of scores: 0-24	4; Better indicated	d by lower values)					
1	randomised trials	Seriousª		no serious indirectness	Serious ^b	none	57	61	-	MD 1.16 higher (0.71 lower to 3.03 higher)	⊕⊕OO LOW	CRITICAL
Healthca	are utilisation	(primary	y care contacts)	>4 months (foll	ow-up 1 years	; Better indicated	by lower values)					
1	randomised trials	Seriousª			no serious imprecision	none	57	61	-	MD 0.09 lower (0.4 lower to 0.22 higher)	⊕⊕⊕O MODERATE	IMPORTANT
Healthca	are utilisation	(prescri	ptions) >4 month	ns (follow-up 1	years; Better i	ndicated by lowe	r values)					
1	randomised trials	Seriousª		no serious indirectness	Serious ^b	none	57	61	-	MD 0.49 lower (1.14 lower to 0.16 higher)	⊕⊕OO LOW	IMPORTANT

Table 65: self-management (exercise prescription) + Postural therapy (Alexander technique - 24 lessons) versus Postural therapy (Alexander technique - 24 lessons)

	Quality assessment						No of patients			Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Alexander technique (24 lessons) + self-management (exercise prescription) versus Alexander technique (24 lessons)	Control	Relative (95% CI)	Absolute	Quality	Importance

^a Downgraded by one increment if the majority of the evidence was at high risk of bias.
^b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs.

Qualty o	of life (SF-36 p	ohysical (component sum	mary) >4 montl	ns (follow-up 1	years; range of	scores: 0-100; Better indicated by	higher v	alues)			
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	56	61	-	MD 2.4 lower (10.62 lower to 5.82 higher)	⊕⊕⊕O MODERATE	CRITICAL
Qualty o	of life (SF-36 r	nental co	omponent summ	ary) >4 months	(follow-up 1 y	/ears; range of so	cores: 0-100; Better indicated by h	igher val	ues)			
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	56	61	-	MD 1.25 higher (6.96 lower to 9.46 higher)	⊕⊕⊕O MODERATE	CRITICAL
Pain (Vo	on Korff pain	scale) >4	months (follow-	up 1 years; rar	ige of scores:	0-10; Better indic	ated by lower values)					
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	56	61	-	MD 0.29 lower (1.21 lower to 0.63 higher)	⊕⊕⊕O MODERATE	CRITICAL
Functio	n (RMDQ, 0-2	4) >4 mo	nths (follow-up 1	years; range o	of scores: 0-24	; Better indicated	I by lower values)					
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	56	61	-	MD 0.08 lower (1.96 lower to 1.8 higher)	⊕⊕⊕O MODERATE	CRITICAL
Healthc	are utilisation	(primary	y care contacts)	> 4months (foll	ow-up 1 years	; Better indicated	l by lower values)	•				
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	56	61	-	MD 0.15 higher (0.2 lower to 0.5 higher)	⊕⊕OO LOW	IMPORTANT
Healthc	are utilisation	(prescri	ptions) >4 montl	ns (follow-up 1	years; Better	indicated by lowe	r values)					
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	61	57	-	MD 0.39 lower (1.12 lower to 0.34 higher)	⊕⊕OO LOW	IMPORTANT

^a Downgraded by one increment if the majority of the evidence was at high risk of bias ^b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs.

Table 66: self-management (exercise prescription) + Postural therapy (Alexander technique -24 lessons) versus Postural therapy (Alexander technique - 6 lessons) plus self-management (exercise prescription)

			Quality as:	sessment			No of patients			Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alexander technique (24 lessons) + self-management (exercise prescription) versus Alexander technique (6 lessons) + self-management (exercise prescription)	Control	Relative (95% CI)	Absolute	Quality	Importance
Qualty o	f life (SF-36	physical	component sum	nmary) >4 mon	ths (follow-up	1 years; range o	of scores: 0-100; Better indicated by h	nigher v	alues)			
1	randomised trials	Seriousª	no serious inconsistency		no serious imprecision	none	56	57	-	MD 0.9 higher (7.56 lower to 9.36 higher)	⊕⊕⊕O MODERATE	CRITICAL
Qualty o	f life (SF-36 ı	mental co	omponent sumn	nary) >4 month	s (follow-up 1	years; range of	scores: 0-100; Better indicated by hig	gher val	ues)			
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	56	57	-	MD 4.35 higher (3.97 lower to 12.67 higher)	⊕⊕OO LOW	CRITICAL
Pain (Vo	n Korff pain	scale) >4	months (follow	-up 1 years; ra	nge of scores	s: 0-10; Better inc	licated by lower values)					
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	56	57	-	MD 0.55 lower (1.49 lower to 0.39 higher)	⊕⊕OO LOW	CRITICAL
Function	n (RMDQ, 0-2	4) >4 mo	nths (follow-up	1 years; range	of scores: 0-2	24; Better indicat	ed by lower values)					
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	56	57	-	MD 1.24 lower (3.15 lower to 0.67 higher)	⊕⊕OO LOW	CRITICAL
Healthca	re utilisation	n (primar	y care contacts)	>4months (fol	low-up 1 year	s; Better indicate	ed by lower values)					
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	56	57	-	MD 0.24 higher (0.1 lower to 0.58 higher)	⊕⊕OO LOW	IMPORTANT

J.4.7

^a Downgraded by one ^b Downgraded by one					if the confidence interval crossed both N	MIDs.	
Low back pain v	vith or	without scia	tica				

none

Table 67:	Self-management	(home exercise)	plus electrotherapy	(laser)	compared with	electrotherany	(laser)
Table U/.	Jen-management	IIIOIIIE EXELUISEI	plus ciecti otilici apy	(lasel)	Compared with	ciccii otilici apy	(lasel)

no serious

imprecision

Healthcare utilisation (prescriptions) > 4 months (follow-up 1 years; Better indicated by lower values)

no serious

indirectness

randomised | Serious | no serious

trials

inconsistency

			Quality	assessment			No of patient	s		Effect	Quality	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Home exercise + laser	laser	Relative (95% CI)	Absolute	Quality	Importance
Pain (VAS	n (VAS 0-10) - ≤ 4 months (range of scores: 0-10; Better indicated by lower values)											
2		very serious ^a			no serious imprecision	none	44	41	,	MD 0.63 lower (1.24 to 0.01 lower)	⊕OOO VERY LOW	CRITICAL
Function (0	ODI, 0-100) ≤ 4	months (ra	ange of scores	0-100; Better indi	cated by lower va	lues)			•			
2		very serious ^a	,	no serious indirectness	Serious ^d	none	44	41	-	MD 2.82 lower (5.8 lower to 0.16 higher)	⊕OOO VERY LOW	CRITICAL

56

57

MD 0.1 higher

0.66 higher)

(0.46 lower to MODERATE

 $\oplus \oplus \oplus O$

IMPORTANT

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

^b Downgraded by two increments because of heterogeneity I²=86%, p=0.007

^c Downgraded by two increments because of heterogeneity I²=73%, p=0.06

^d Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 68: Self-management (unsupervised exercise) + electrotherapy (HILT laser) vs electrotherapy (HILT laser)

			Quality as	easemant		C. apy (No of patients			Effect		
			Quality as	36331116111			No or patients				0	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Self-management (unsupervised exercise) + electrotherapy (HILT laser) vs electrotherapy (HILT laser)	Control	Relative (95% CI)	Absolute	Quality	Importance
Pain seve	erity (VAS, 0-	10) ≤ 4 m	onths (range of s	cores: 0-10; Be	etter indicated I	by lower values)						
		- ,			no serious imprecision	none	28	20	-	MD 3.01 lower (3.66 to 2.36 lower)	⊕⊕OO LOW	CRITICAL
Function	(RMDQ, 0-24	.) ≤ 4 mor	nths (range of sco	ores: 0-24; Bette	er indicated by	lower values)						
		- 3			no serious imprecision	none	28	20	-	MD 1.85 lower (2.64 to 1.06 lower)	⊕⊕OO LOW	CRITICAL
Function	(MODI, 0-100)) ≤ 4 mor	nths (range of sc	ores: 0-100; Bet	tter indicated b	y lower values)						
		- ,			no serious imprecision	none	28	20	-	MD 3.91 lower (5.96 to 1.86 lower)	⊕⊕OO LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 69: Self-management (education) + exercise (biomechanical) vs exercise (biomechanical – motor control) for low back pain with or without sciatica

			Quality asses	ssment			No of patient	s		Effect				
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Self management plus exercise	Exercise	Relative (95% CI)	Absolute	Quality	Importance		
Pain sever	in severity (VAS, 0-10) (range of scores: 0-10; Better indicated by lower values)													

	randomised trials	- ,			very serious ^b	none	10	11	-	MD 0.7 higher (2.5 to 1.10 higher)	⊕OOO VERY LOW	CRITICAL
Function ((RMDQ, 0-24) (range of s	cores: 0-24; Better	indicated by lowe	er values)							
	randomised trials	- ,		no serious indirectness	Serious ^b	none	10	11	-	MD 1.64 higher (7.06 to 3.78 higher)	⊕OOO VERY LOW	CRITICAL

a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs.

J.5 Exercise therapies

J.5.1 Biomechanical Exercise

J.5.1.1 Individual biomechanical exercise

Table 70: Individual biomechanical exercise versus placebo/sham in low back pain with sciatica

			Quality ass	sessment			No of pati	ents		Effect	Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Individual biomechanical exercise	Placebo/sham	Relative (95% CI)	Absolute	quanty	mportanoc	
With scia	tica - Pain (V	AS 0-10) <	4 months (range	of scores: 0-10	Better indicate	ed by lower value	s)						
	randomised trials			no serious indirectness	Serious ^b	none	83	87	-	MD 0.8 lower (1.53 to 0.07 lower)	⊕⊕OO LOW	CRITICAL	
With scia	h sciatica - Pain (VAS 0-10) > 4 months (range of scores: 0-10; Better indicated by lower values)												
	randomised trials				no serious imprecision	none	82	88	-	MD 0.1 higher (0.58 lower to 0.78	⊕⊕⊕O MODERATE	CRITICAL	

higher)

Table 71: Individual biomechanical exercise versus usual care in low back pain with or without sciatica

			Quality as	sessment			No of patients	\$		Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Individual biomechanical exercise	Usual care	Relative (95% CI)	Absolute	Quality	Importance
Overall -	Quality of life	individua	I (SF-36/RAND-36	0-100) <4 month	ns - general hea	Ith (range of score	s: 0-100; Better indica	ated by h	nigher va	lues)		
2	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	28	29	-	MD 14.13 higher (5.56 to 22.7 higher)	⊕OOO VERY LOW	CRITICAL
Overall -	Quality of life	individua	I (SF-36/RAND-36	0-100) <4 month	s - vitality (rang	ge of scores: 0-100	; Better indicated by	higher v	alues)			•
2	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	28	29	-	MD 12.33 higher (3.4 to 21.25 higher)	⊕OOO VERY LOW	CRITICAL
Overall -	Quality of life	pain scor	re (SF-36/RAND-36	6 0-100) <4 mont	hs - bodily pain	(range of scores:	0-100; Better indicate	d by hig	her value	es)		•
2	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	28	29	-	MD 19.05 higher (12.5 to 25.61 higher)	⊕⊕OO LOW	CRITICAL
Overall -	Quality of life	individua	I (SF-36/RAND-36	0-100) <4 month	ıs - physical rol	e limitation (range	of scores: 0-100; Bet	ter indic	ated by h	igher values)		
2	randomised trials	very serious¹	no serious inconsistency	no serious indirectness	serious ²	none	28	29	-	MD 21.44 higher (10.21 to 32.75 higher)	⊕OOO VERY LOW	CRITICAL
Overall -	Quality of life	individua	I (SF-36/RAND-36	0-100) <4 month	s - emotional re	ole limitation (rang	e of scores: 0-100; Bo	etter indi	cated by	higher values)		•
2	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	28	29	-	MD 12.25 higher (1.34 to 23.16 higher)	⊕000 VERY LOW	CRITICAL
Overall -	Quality of life	individua	I (SF-36/RAND-36	0-100) <4 month	s - social funct	ioning (range of so	ores: 0-100; Better in	dicated	by highe	r values)		
2	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	28	29	-	MD 20.27 higher (11.27 to 29.27 higher)	⊕⊕OO LOW	CRITICAL
Overall -	Quality of life	individua	I (SF-36/RAND-36	0-100) <4 month	ns (unexplained	heterogeneity) - p	hysical functioning (r	ange of	scores: 0	-100; Better indicated	d by higher v	/alues)
2	randomised trials	very serious ¹	serious ³	no serious indirectness	very serious ²	none	28	29	-	MD 12.68 higher (7.94 lower to 33.3 higher)	⊕OOO VERY LOW	CRITICAL
Overall -	Quality of life	individua	I (SF-36/RAND-36	0-100) <4 month	s (unexplained	heterogeneity) - m	nental health (range o	fscores	0-100; E	Better indicated by hig	gher values)	
2	randomised trials	very serious ¹	very serious ⁴	no serious indirectness	very serious ²	none	28	29	-	MD 2.88 higher (14.38 lower to 20.15 higher)	⊕OOO VERY LOW	CRITICAL
Overall -	Pain (VAS 0-1	0) <4 mor	nths - Pain (follow	-up <4 months; r	range of scores	: 0-10; Better indic	ated by lower values)					
5	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	181	136	-	MD 0.74 lower (1.12 to 0.36 lower)	⊕⊕⊕O MODERATE	CRITICAL

a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

0	D=: (\/AC 0 4	10\ <4	otha Dain at want	(fallan, 44 ma		0 40: Datte		·=l=\				
Overall -	· · · · · · · · · · · · · · · · · · ·	, ' 		· ·		· · · · · ·	er indicated by lower v			1		
1	randomised	serious ¹	no serious	no serious	no serious	none	15	15	-	MD 1.61 lower (2.21	$\oplus \oplus \oplus O$	CRITICAL
	trials		inconsistency	indirectness	imprecision					to 1.01 lower)	MODERATE	
Overall -	Pain (VAS 0-1	l0) <4 mor	nths - Pain during	movement (follo	w-up <4 month	s; range of scores	: 0-10; Better indicated	d by low	er value	s)		
1	randomised	serious1	no serious	no serious	no serious	none	15	15	-	MD 2.07 lower (2.55	$\oplus \oplus \oplus O$	CRITICAL
	trials		inconsistency	indirectness	imprecision					to 1.59 lower)	MODERATE	
Overall -	Pain (VAS 0-1	10) <4 mor	nths - Pain- chair i	rise (follow-up <	months; range	of scores: 0-10; E	Setter indicated by low	er value	es)			
1	randomised	very	no serious	no serious	very serious ²	none	18	14	-	MD 0.4 lower (1.86	⊕000	CRITICAL
	trials	serious1	inconsistency	indirectness						lower to 1.066 higher)	VERY LOW	
Overall -	Pain (VAS 0-1	10) <4 mor	nths - Pain walking	g (follow-up <4 n	nonths; range o	f scores: 0-10; Bet	ter indicated by lower	values)				
1	randomised	verv	no serious	· ·	serious ²	none	18	14	_	MD 1.5 lower (3.38	⊕000	CRITICAL
	trials	serious ¹	inconsistency	indirectness			-			lower to 0.38 higher)		
Overall -	Pain (VAS 0-1	l0) <4 mor	nths - Pain stair cl	imb (follow-up <	4 months; range	e of scores: 0-10; I	Better indicated by low	ver value	es)			
1	randomised	verv	no serious	no serious	very serious ²	none	18	14	-	MD 0.3 higher (1.42	⊕000	CRITICAL
	trials	serious ¹	inconsistency	indirectness	- ,		-			lower to 2.02 higher)	VERY LOW	
Overall -	Pain (VAS 0-1	10) >4 mor	nths - 1 year (follo	w-up >4 months	; range of score	s: 0-10; Better ind	cated by lower values	5)		,		
1	randomised	verv	no serious	no serious	no serious	none	71	28	_	MD 0.08 lower (1.53	⊕⊕00	CRITICAL
	trials	serious1	inconsistency	indirectness	imprecision					lower to 1.37 higher)	LOW	
Overall -	Function (RM	IDQ/ODQ)	<4 months (follow	v-up <4 months;	Better indicated	by lower values)				,		
5	randomised	serious ¹	no serious	no serious	serious ²	none	150	103	-	SMD 1.31 lower (2.47	$\oplus \oplus OO$	CRITICAL
	trials		inconsistency	indirectness						to 0.15 lower)	LOW	
Overall -	Function (RM	DQ/ODQ	0-100) 4 months -	1 year (follow-up	>4 months; Be	tter indicated by lo	ower values)			· ·		
2	randomised	serious1	no serious	no serious	serious ²	none	101	58	_	SMD 0.32 lower (0.66	$\oplus \oplus OO$	CRITICAL
	trials		inconsistency	indirectness			-			lower to 0.01 higher)	LOW	
Overall -	Psychologica	l distress	(mental health in	ventory 24-142) (Better indicated	by lower values)						
1	randomised	very	no serious	no serious	serious ²	none	31	23	-	MD 11.3 lower (26.48	⊕OOO	
	trials	serious1	inconsistency	indirectness						lower to 3.88 higher)		
a Downar	radad by 1 inc	romont if t	ha majority of avid	Janca was at high	rick of high and	1 by 2 increments if	the majority of eviden	co wac a	t vory hi	ah risk of hias	I I	

a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias

Table 72: Individual biomechanical exercise versus usual care in low back pain with sciatica

			Quality asse	ssment			No of p	patients		Effect	Quality	Importance
No of studies					Other considerations	Individual biomechanica	Usual care	Relative (95% CI)	Absolute	,		

b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

c Heterogeneity, I2=84%, unexplained by subgroup analysis

d Heterogeneity, I2 = 80%, unexplained by subgroup analysis

							I exercise						
With sciatica - Pain (VAS 0-10) <4 months (Better indicated by lower values)													
		- 3			serious imprecision ^b	none	26	26	-	MD 1.70 lower (2.33 to 1.07 lower)	⊕000 VERY LOW	CRITICAL	

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 73: Individual biomechanical exercise versus usual care in low back pain without sciatica

			Quality asse	ssment			No of	patients	I	Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Individual biomechanica I exercise	Usual care	Relative (95% CI)	Absolute	Quanty	importance
Without	sciatica - Qua	lity of life (S	SF-36) <4 months	- Functional ca	apacity (Bette	r indicated by lov	ver values)					
1		very serious ^a		no serious indirectness	very serious ²	none	30	30	-	MD 1.1 lower (13.47 lower to 11.27 higher)	⊕OOO VERY LOW	CRITICAL
Without	Vithout sciatica - Quality of life (SF-36) <4 months - Pain (Better indicated by lower values)											
1		very serious ^a		no serious indirectness	serious ^b	none	30	30	-	MD 11.5 higher (2.25 to 20.75 higher)	⊕OOO VERY LOW	CRITICAL
Without	sciatica - Qua	lity of life (S	SF-36) <4 months	- General heal	th (Better indi	cated by lower v	alues)					
1	randomised trials	very serious ^a		no serious indirectness	very serious ^b	none	30	30	-	MD 6.9 higher (3.54 lower to 17.34 higher)	⊕OOO VERY LOW	CRITICAL
Without	sciatica - Qua	lity of life (S	SF-36) <4 months	- Vitality (Bette	er indicated b	y lower values)						
1	randomised trials	very serious ^a		no serious indirectness	no serious imprecision	none	30	30	-	MD 15.6 higher (6.35 to 24.85 higher)	⊕⊕OO LOW	CRITICAL

Mithou	t sciatica. Qua	lity of life /	SE 36) <4 month	e Social aspor	ets (Bottor ind	icated by lower v	alues)					
1		very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	30	30	-	MD 14.4 higher (3.27 to 25.53 higher)	⊕⊕OO LOW	CRITICAL
Vithou	t sciatica - Qua	ality of life (SF-36) <4 month	s - Emotional as	spects (Better	indicated by low	er values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	30	30	-	MD 19 higher (0.68 lower to 38.68 higher)	⊕OOO VERY LOW	CRITICAI
Withou	t sciatica - Qua	ality of life (SF-36) <4 month	s - physical (Be	tter indicated	by lower values)						
2	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	50	49	-	MD 13.54 higher (4.08 to 22.99 higher)	⊕⊕OO LOW	CRITICAL
Withou	t sciatica - Qua	ality of life (SF-36) <4 month	s - mental (Bett	er indicated b	y lower values)						
2	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	50	49	-	MD 12.63 higher (5.72 to 19.53 higher)	⊕⊕OO LOW	CRITICAI
Withou	t sciatica - Qua	lity of life (SF-36) 4 months	- 1 year - Funct	ional capacity	/ (Better indicated	d by lower valu	es)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	30	30	-	MD 5.4 higher (6.11 lower to 16.91 higher)	⊕OOO VERY LOW	CRITICAL
Withou	t sciatica - Qua	lity of life (SF-36) 4 months	- 1 year - Pain (Better indicat	ed by lower value	es)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	30	30	-	MD 8.5 higher (0.05 to 16.95 higher)	⊕OOO VERY LOW	CRITICAL
Withou	t sciatica - Qua	ality of life (SF-36) 4 months	- 1 year - Gener	ral health (Bet	ter indicated by I	ower values)	•	I .			
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b		30	30	-	MD 5.2 higher (5.57 lower to 15.97 higher)	⊕OOO VERY LOW	CRITICAL
Withou	t sciatica - Qua	lity of life (SF-36) 4 months	- 1 year - Vitalit	y (Better indi	cated by lower va	lues)	'		,		

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-	1	ı	T			1	ı			1		
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	30	30	-	MD 14 higher (4.39 to 23.61 higher)	⊕⊕OO LOW	CRITICAL
Without	sciatica - Qua	lity of life (SF-36) 4 months	- 1 year - Social	aspects (Bet	ter indicated by l	ower values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	30	30	-	MD 8.1 higher (4.55 lower to 20.75 higher)	⊕OOO VERY LOW	CRITICAL
Without	sciatica - Qua	lity of life (SF-36) 4 months	- 1 year - Emoti	onal aspects	(Better indicated	by lower values	s)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	30	30	-	MD 27.3 higher (9.55 to 45.05 higher)	⊕⊕OO LOW	CRITICAL
Without	sciatica - Qua	lity of life (SF-36) 4 months	- 1 year - Physic	cal (Better inc	licated by lower v	alues)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	30	30	1	MD 22.4 higher (3.4 to 41.4 higher)	⊕⊕OO LOW	CRITICAL
Without	sciatica - Qua	lity of life (SF-36) 4 months	- 1 year - Menta	l health (Bett	er indicated by lo	wer values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	30	30	-	MD 10.3 higher (0.02 to 20.58 higher)	⊕OOO VERY LOW	CRITICAL
Without	sciatica- Fund	tion (RMD)	Q) <4 months (rar	nge of scores: ()-23; Better in	dicated by lower	values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	18	14	-	MD 1.9 higher (1.46 lower to 5.26 higher)	⊕OOO VERY LOW	CRITICAL
Without	sciatica - Fun	ction (RMD	Q 0-24) <4 month	s (range of sco	res: 0-24; Be	tter indicated by I	ower values)					
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	43	43	-	MD 2.7 lower (4.4 to 1 lower)	⊕⊕⊕O MODERAT E	CRITICAL
Without	sciatica - Fun	ction (RMD	Q 0-24) 4 months	- 1 year (Better	r indicated by	lower values)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	43	43	-	MD 1.54 lower (3.1 lower to 0.03 higher)	⊕OOO VERY LOW	CRITICAL

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Withou	t sciatica - Fun	ction (RMD	Q 0-24) < 4 mont	hs (Better indic	ated by lowe	r values)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	237	181	-	MD 0.96 lower (1.95 lower to 0.04 higher)	⊕⊕OO LOW	CRITICAL
Withou	t sciatica - Fun	ction (RMD	Q 0-24) 4 months	s - 1 year (Bette	r indicated by	lower values)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	30	30	-		⊕000 VERY LOW	CRITICAL
Withou	t sciatica - Fun	ction (chan	ge score, ODI) <	4 months - Full	range of mot	ion (Better indica	ted by lower va	alues)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	10	7	-	MD 1.52 lower (2.174 to 0.866 lower)	⊕⊕OO LOW	CRITICAL
Withou	t sciatica - Fun	ction (chan	ge score, ODI) <	4 months - Lim	ited range of	motion (Better in	dicated by lowe	er values)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	7	7	-	MD 0.9 lower (1.536 to 0.264 lower)	⊕000 VERY LOW	CRITICAL
Withou	t sciatica - Pair	1 1 (VAS 0-10	') <4 months - Pa	in (VAS 0-10) <	4months (Bet	tter indicated by l	ower values)					
4	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	124	122	-	MD 1.14 lower (1.61 to 0.67 lower)	⊕OOO VERY LOW	CRITICAL
Withou	t sciatica - Pair	1 (VAS 0-10	') 4 months - 1 ye	ar - Pain (VAS ()-10) 4 month	s - 1 year (Better	indicated by lo	wer values)		·	1	
2	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	73	73	-	MD 1.05 lower (1.76 to 0.35 lower)	⊕000 VERY LOW	CRITICAL
Withou	t sciatica - Pair	n (0-85) <4 r	nonths (change	score) (range o	f scores: 0-85	; Better indicated	l by lower value	es)				
4	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	130	130	-	MD 0.00 higher (6.6 lower to 6.6 higher)	⊕⊕OO LOW	CRITICAL
Withou	t sciatica - Pair	n (VAS 0-85) >4 months - 1 y	ear (range of s	cores: 0-85; B	etter indicated by	y lower values)					

1		very serious ^a		no serious indirectness	no serious imprecision	none	137	134	-	MD 1 higher (4.48 lower to 6.48 higher)	⊕⊕OO LOW	CRITICAL
Without	sciatica - Pain	(change s	core VAS 0-10) <4	I months - Full	range of mot	ion (Better indica	ted by lower va	lues)				
1		very serious¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	10	7	-	MD 3.701 lower (5.642 to 1.76 lower)	⊕⊕OO LOW	CRITICAL
Without	sciatica - Pain	(change s	core VAS 0-10) <4	l months - Lim	ited range of	motion (Better inc	dicated by lowe	r values)				
1		very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	7	7	-	MD 2.3 lower (3.67 to 0.93 lower)	⊕⊕OO LOW	CRITICAL
without s	ciatica-adver	se events (morbidity)<4 mor	nths	•							
1	trials	very serious ¹	no serious inconsistency	no serious indirectness	serious²	none	3/20 (15%)	(0%)	RR 7 (0.38 to 127.32)		⊕OOO VERY LOW	IMPORTANT

Table 74: Individual biomechanical exercise versus self-management in low back pain with or without sciatica

			Quality ass	essment			No of p	patients		Effect	Ovelity	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Individual biomechanical exercise	Self-management (advice to stay active)	Relative (95% CI)		Quality	Importance
Overall -	Pain (VAS 0-	10) <4 mont	ths (range of sco	res: 0-10; Bette	r indicated by l	ower values)						
		,		no serious indirectness	Serious ^b	none	48	29	-	MD 0.7 lower (2 lower to 0.6 higher)	⊕OOO VERY LOW	CRITICAL
Overall -	Leg pain (VA	S 0-10) <4 r	months - Overall	with or without	sciatica (range	of scores: 0-10;	Better indicated by l	ower values)				
1	randomised	very	no serious	no serious	Serious ^b	none	48	29	-	MD 0.8 lower	⊕000	CRITICAL

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

	trials	serious ^a	inconsistency	indirectness						(2.2 lower to 0.6 higher)	VERY LOW	
Overall -	Pain (VAS 0-	10) 4 montl	ns - 1 year (range	of scores: 0-10	; Better indicat	ed by lower value	es)					
1		very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	45	26	-	MD 0.4 lower (1.7 lower to 0.9 higher)	⊕OOO VERY LOW	CRITICAL
Overall -	Leg pain (VA	S 0-10) 4 m	onths - 1 year (ra	inge of scores:	0-10; Better in	dicated by lower v	values)					
1		no serious risk of bias	very serious ^c	no serious indirectness	no serious imprecision	none	45	26	-	MD 1 lower (2.3 lower to 0.3 higher)	⊕⊕OO LOW	CRITICAL
Overall -	Function (RM	MDQ 0-24) <	4 months (range	of scores: 0-24	; Better indicat	ed by lower value	es)					
1		very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	48	29	-	MD 1 lower (4 lower to 2 higher)	⊕OOO VERY LOW	CRITICAL
Overall -	Function (RM	MDQ 0-24) 4	months - 1 year	(range of score	s: 0-24; Better	indicated by lowe	r values)					
1		very seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	45	26	-	MD 3 lower (6 lower to 0 higher)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs ^c Heterogeneity, I²=80%, unexplained by subgroup analysis

Table 75: Individual biomechanical exercise versus spinal manipulation (low-amplitude high-velocity thrust) in low back pain with sciatica

			Quality as:	sessment			No of pa	itients		Effect	0114	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Individual biomechanical exercise	SMT (low- amplitude high- velocity)	Relative (95% CI)	Absolute	Quality	Importance
With sciat	tica - Quality	of life (SF	F-36 0-100) <4 mo	nths- physical c	omponent (Bet	ter indicated by lo	ower values)					

1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	92	99	-	MD 1.7 higher (0.5 lower to 3.9 higher)		CRITICAL
With scia	tica - Quality	of life (SI	-36 0-100) <4 mc	onths- mental co	mponent (Bette	er indicated by lov	ver values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	92	99	1	MD 2 lower (3.91 to 0.09 lower)	⊕OOO VERY LOW	CRITICAL
With scia	tica - Quality	of life (SF	F-12 0-100) 4 mor	nths - 1 year - ph	ysical compon	ent (Better indicat	ed by lower values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	82	82	1	MD 2 higher (0.33 lower to 4.33 higher)	⊕OOO VERY LOW	CRITICAL
With scia	tica - Quality	of life (SF	F-12 0-100) 4 mor	nths - 1 year - me	ental componer	nt (Better indicate	d by lower values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	82	82	1	MD 1.3 lower (3.77 lower to 1.17 higher)	⊕⊕OO LOW	CRITICAL
With scia	tica - Pain (V	AS 0-10) <	<4 months (Bette	r indicated by lo	wer values)							
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	92	99	1	MD 0.3 lower (0.87 lower to 0.27 higher)	⊕⊕OO LOW	CRITICAL
With scia	tica - Pain (V	AS 0-10) 4	4 months - 1 year	r (Better indicate	d by lower valu	ies)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ²	none	82	82	1	MD 0.5 lower (1.17 lower to 0.17 higher)	⊕OOO VERY LOW	CRITICAL
With scia	tica - Functio	n (RMDQ	0-24) <4 months	(Better indicate	d by lower valu	es)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	92	99	-	MD 0.1 higher (1.22 lower to 1.42 higher)	⊕⊕OO LOW	CRITICAL
With scia	tica - Functio	n (RMDQ	0-24) 4 months -	- 1 year (Better in	ndicated by low	er values)						
1	randomised	very	no serious	no serious	no serious	none	82	82	-	MD 0.2 lower (1.72	⊕⊕00	CRITICAL

trials	coriousa	inconsistancy	indirectness	improcision			lower to 1.32	LOW	
uiais	serious	lilicorisistericy	illuli ecti less	imprecision			10Wei 10 1.32	LOVV	
							higher)		
							riigriei <i>)</i>		

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 76: Individual biomechanical exercise versus individual interferential exercise in low back pain with or without sciatica

			Quality as:	sessment			No of p	oatients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Individual biomechanical	Individual interferential therapy	Relative (95% CI)	Absolute	Quanty	Importance
Overall-P	ain (VAS 0-10)) <4 mon	ths (range of sco	res: 0-10; Better	indicated by lo	ower values)						
1	randomised trials				no serious imprecision	none	30	30	-	MD 1.2 lower (1.55 to 0.85 lower)	⊕⊕⊕O MODERATE	CRITICAL

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias

J.5.1.2 Group Biomechanical Exercise

Table 77: Group biomechanical exercise versus usual care in low back pain with or without sciatica

			Quality as	sessment			No of patients	5		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Group biomechanical exercise	Usual care	Relative (95% CI)	Absolute	Quality	Importance
Overall-P	ain (VAS) >4 r	nonths (r	ange of scores: 0-	10; Better indica	ted by lower va	lues)						
1		- ,	no serious inconsistency	no serious indirectness	serious ^b	none	64	63	-	MD 1.34 lower (1.9 to 0.78 lower)	⊕OOO VERY LOW	CRITICAL
Overall-P	ain (VAS) <4 r	nonths (r	ange of scores: 0-	10; Better indica	ted by lower va	lues)						
1		- ,	no serious inconsistency	no serious indirectness	serious ^b	none	64	63		MD 0.52 lower (1.12 lower to 0.08 higher)		CRITICAL
Overall - I	Pain <4 month	ns - stretc	hing (range of sco	ores: 0-10; Better	indicated by lo	wer values)			,			

1	randomised trials	very serious ^a	no serious inconsistency		no serious imprecision	none	62	60	-	MD 0.09 higher (0.8 lower to 0.98 higher)	⊕⊕OO LOW	CRITICAL	
Overall -	Pain (VAS 0-1	0) <4 mor	nths - core stabilit	y (Better indicate	ed by lower valu	es)							
1	randomised trials	serious ª	no serious inconsistency		no serious imprecision	none	20	20	-	MD 2.2 lower (2.96 to 1.44 lower)	⊕⊕⊕O MODERATE	CRITICAL	
Overall -	verall - Function (RMDQ 0-24) <4 months (Better indicated by lower values)												
1	randomised trials	serious ª	no serious inconsistency	no serious indirectness	serious ^b	none	20	20	-	MD 5.06 lower (8.65 to 1.47 lower)	⊕⊕OO LOW	CRITICAL	
Overall-N	SAID use >4 r	nonths (E	Better indicated by	lower values)									
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	30	30		MD 7.13 lower (14.5 lower to 0.24 higher)	⊕⊕OO LOW	IMPORTANT	

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 78: Group biomechanical exercise versus usual care in low back pain without sciatical

	. Group b		inical exercise	cisas asaai c	are iii iott ba	ck pain without	. sciatica		·		1	
			Quality as:	sessment			No of patients	5		Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Group biomechanical exercise	Usual care	Relative (95% CI)	Absolute	Quality	Importance
Without s	ciatica - Quali	ity of life o	composite scores	(SF-36 0-100) <4	months - Menta	er indicated by lower	r values)				
1	randomised trials	serious ^a			no serious imprecision	none	9	9	-	MD 9.04 higher (6.57 to 11.51 higher)	⊕⊕⊕O MODERATE	CRITICAL
Without s	ciatica - Quali	ity of life o	composite scores	(SF-36 0-100) <4	months - Physi	cal component (Be	etter indicated by low	er value	es)			
	randomised trials	serious ^a	no serious inconsistency		no serious imprecision	none	9	9	-	MD 8.3 higher (5.3 to 11.3 higher)	⊕⊕⊕O MODERATE	CRITICAL
Without s	ciatica - Quali	ity of life i	ndividual scores (SF-12) <4 month	s - general heal	th (Better indicate	d by lower values)		•			

1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ^b	none	20	14	-	MD 0.10 higher (0.51 lower to 0.71 higher)	⊕OOO VERY LOW	CRITICAL
Without	sciatica - Qua	lity of life i	ndividual scores	(SF-12) <4 month	ns - physical fun	ctioning (Better in	dicated by lower valu	ies)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	20	14	-	MD 0.1 higher (0.19 lower to 0.39 higher)	⊕000 VERY LOW	CRITICAL
Without	sciatica - Qua	lity of life i	individual scores	(SF-12) <4 month	ns - physical role	e limitation (Better	indicated by lower v	alues)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	20	14	ı	MD 0.2 higher (0.31 lower to 0.71 higher)	⊕000 VERY LOW	CRITICAL
Without	sciatica - Qua	lity of life i	individual scores	(SF-12) <4 month	ns - bodily pain (Better indicated b	y lower values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	20	14	-	MD 0.5 lower (1.11 lower to 0.11 higher)	⊕OOO VERY LOW	CRITICAL
Without	sciatica - Qua	lity of life i	individual scores	(SF-12) <4 month	ns - social functi	oning (Better indic	cated by lower values	s)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	20	14	-	MD 0.1 higher (0.31 lower to 0.51 higher)	⊕OOO VERY LOW	CRITICAL
Without	sciatica - Qua	lity of life i	individual scores	(SF-12) <4 month	ns - health perce	eption (Better indic	ated by lower values)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	20	14	-	MD 0.3 lower (0.84 lower to 0.24 higher)	⊕OOO VERY LOW	CRITICAL
Without	sciatica - Pain	(VAS 0-10)) <4 months (Bett	er indicated by I	ower values)							
2	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	29	23	-	MD 0.87 lower (1.27 to 0.46 lower)	⊕OOO VERY LOW	CRITICAL
Without	sciatica - Fun	ction (ODI	0-100) <4 months	(Better indicated	d by lower value	s)						
2	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	29	23	-	MD 13.97 lower (16.07 to 11.88 lower)	⊕000 VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 79: Group biomechanical exercise versus unsupervised exercise in low back pain with or without sciatica

			Quality asse	essment			No of pa	tients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Group biomechanical exercise	Unsupervised exercise	Relative (95% CI)	Absolute	Quanty	importance
Overall - F	Pain (VAS 0-10	0) <4 mon	ths (Better indicat	ed by lower valu	ies)							
		,		no serious indirectness	Serious ^b	none	83	87	1	MD 0.8 lower (1.53 to 0.07 lower)	⊕000 VERY LOW	CRITICAL
Overall - F	Pain (VAS 0-10	0) 4 montl	hs - 1 year (Better	indicated by low	er values)							
		,	no serious inconsistency	no serious indirectness	Serious ^b	none	71	70	-	MD 1.45 lower (2.2 to 0.7 lower)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Individual aerobic exercise J.5.1.3

Table 80: Individual aerobic exercise versus usual care in low back pain with or without sciatica

			Quality asse	ssment			No of patien	ts		Effect	Ovalita	l was a set a se a a
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Individual aerobic exercise	Usual care	Relative (95% Absolute CI)		Quanty	Importance
Overall - P	ain (VAS 0-10)	<4 month	s (Better indicated	by lower values)								
	randomised trials			no serious indirectness	serious	none	24	22	-	MD 0.3 lower (1.52 lower to 0.92 higher)	⊕⊕OO LOW	CRITICAL

Overall - F	unction (ALBF	PS 0-100) <	4 months (Better in	ndicated by lower	values)							
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	24	22	-	MD 1.8 lower (9.24 lower to 5.64 higher)	⊕⊕OO LOW	CRITICAL
Overall - F	unction (RMD	Q/ALBPS)	4 months - 1 year (Better indicated b	y lower value	es)						
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	24	22	-	MD 5.6 lower (14.36 lower to 3.16 higher)	⊕⊕OO LOW	CRITICAL

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 81: Individual aerobic exercise versus usual care in low back pain without sciatica

			Quality as	sessment			No of patien	ts		Effect	O. alita	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Individual aerobic exercise	Usual care	Relative (95% CI)	Absolute	Quality	Importance
Without s	hout sciatica - Quality of life (EuroQol weighted health index 0.59-1) 4 months - 1 year (Better indicated by lower values)											
	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	39	17	-	MD 0.06 lower (0.19 lower to 0.07 higher)		CRITICAL
Without s	ciatica - Quali	ty of life (F	EuroQol VAS 0-100)) 4 months - 1 ye	ear (Better indica	ted by lower value	es)					
	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	40	17	-	MD 9.6 higher (3.69 lower to 22.89 higher)	⊕⊕OO LOW	CRITICAL
Without s	ciatica - Pain	(VAS 0-10)	<4 months (deep	water running) (r	ange of scores:	0-10; Better indica	ated by lower valu	es)				
	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	25	24	-	MD 1.49 lower (2.35 to 0.63 lower)	⊕⊕OO LOW	CRITICAL
Without s	nout sciatica - Pain (VAS 0-10) <4 months (treadmill running) (range of scores: 0-100; Better indicated by lower values)											
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	19	18	-	MD 0.05 higher (1.62 lower to 1.72 higher)	⊕000 VERY LOW	CRITICAL

Without s	ciatica - Pain	(VAS 0-10)) 4 months - 1 year	(deep water run	ning) (range of s	scores: 0-10; Bette	r indicated by low	er value	s)			
1	randomised trials	Seriousª	no serious inconsistency		no serious imprecision	none	25	24	1	MD 2.6 lower (3.28 to 1.92 lower)	⊕⊕⊕O MODERATE	CRITICAL
Without s	sciatica - Pain ((VAS 0-10)	4 months - 1 year	(walking) (range	of scores: 0-10	; Better indicated b	y lower values)					
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	40	17	-	MD 0.3 lower (1.77 lower to 1.17 higher)	⊕⊕OO LOW	CRITICAL
Without s	ciatica - Func	tion (RMD	Q 0-24) <4 months	(Better indicated	by lower values	s)						
2	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	44	42	-	MD 2.6 lower (4.21 to 0.99 lower)	⊕⊕OO LOW	CRITICAL
Without s	ciatica - Psycl	hological (distress (BDI 0-63)	<4 months (Bette	er indicated by I	ower values)						
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	very serious ^b	none	19	18	-	MD 0.2 higher (5.57 lower to 5.97 higher)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 82: Individual aerobic exercise versus individual biomechanical exercise in low back pain with or without sciatica

			Quality asse	essment			No c	of patients		Effect	O. alifa	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Individual aerobic exercise	Individual biomechanical exercise	Relative al (95% Absolute CI)		Quality	Importance
Overall - I	Function (ODI	0-100) <4	months (Better in	ndicated by lowe	r values)							
1	randomised trials			no serious indirectness	Serious ^b	none	26	26	-	MD 3.5 higher (3.91 lower to 10.91 higher)	⊕⊕OO LOW	CRITICAL

a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias

b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 83: Individual aerobic exercise versus individual biomechanical exercise in low back pain with or without sciatica

14.0.0								ill with or withou				
			Quality asse	essment			No o	of patients		Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Individual aerobic exercise	Group biomechanical exercise	Relative (95% CI)	Absolute	Quality	Importance
Quality of	f life: SF-36, F	hysical C	omponent Score,	0-100 (follow-up	mean 6 moi	nths; Better indica	ted by higher va	ilues)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	16	14	-	MD 2.27 lower (8.67 lower to 4.13 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	f life: SF-36, N	lental Co	mponent Score, 0-	-100 (follow-up n	nean 6 montl	ns; Better indicate	d by higher valu	ies)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	16	14	-	MD 3.63 lower (11.94 lower to 4.68 higher)	⊕OOO VERY LOW	CRITICAL
Psycholo	gical distress	: HADS, A	Anxiety, 0-21 (follo	w-up mean 6 m	onths; Better	indicated by lowe	er values)					
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	16	14	-	MD 1.16 higher (1.54 lower to 3.86 higher)		CRITICAL
Psycholo	gical distress	: HADS, [Depression, 0-21 (follow-up mean	6 months; Be	etter indicated by	ower values)					
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	very serious ²	none	16	14	-	MD 0.32 higher (2.97 lower to 3.61 higher)		CRITICAL
Pain seve	erity: NRS ave	erage bacl	k pain <4 months,	0-10 (follow-up	mean 6 mont	hs; Better indicate	ed by lower valu	es)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	very serious ²	none	16	14	-	MD 0 higher (1.68 lower to 1.68 higher)	⊕000 VERY LOW	CRITICAL
Pain seve	erity: NRS ave	erage back	k pain >4 months,	0-10 (follow-up	mean 3 mont	hs; Better indicate	ed by lower valu	es)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	16	14	-	MD 1.1 higher (0.67 lower to 2.87 higher)	⊕OOO VERY LOW	CRITICAL

Pain severity: NRS average leg pain <4 months, 0-10 (follow-up mean 6 months; Better indicated by lower values)												
1		- ,		no serious indirectness	very serious ²	none	16	14		MD 0.07 higher (2.07 lower to 2.21 higher)		CRITICAL
Pain severity: NRS average leg pain >4 months, 0-10 (follow-up mean 3 months; Better indicated by lower values)												
1		- ,		no serious indirectness	very serious²	none	16	14		MD 0.04 lower (2.29 lower to 2.21 higher)		CRITICAL

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

Group aerobic exercise J.5.1.4

Table 84: Group aerobic exercise versus usual care in low back pain without sciaitca

Quality assessment								No of patients		Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Group aerobic exercise		Relative (95% CI)		Quality	Importance
Without sciatica - Quality of life (SF-36 mental component 0-100) <4 months (Better indicated by lower values)												
2		very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	59	50	-	MD 3.86 higher (2.19 to 5.53 higher)	⊕000 VERY LOW	CRITICAL
Without sciatica - Quality of life (SF-36 physical component 0-100) <4 months (Better indicated by lower values)												
2		very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	59	50	-	MD 2.26 higher (0.02 to 4.5 higher)	⊕000 VERY LOW	CRITICAL
Without sciatica - Quality of life (SF-36 physical functioning 0-100) <4 months (range of scores: 0-100; Better indicated by higher values)												

	_		•	,	•	_	,					1
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	10	10	-	MD 15.5 higher (4.55 lower to 35.55 higher)	⊕OOO VERY LOW	CRITICAL
Without	sciatica - Qualit	ty of life (SF	-36 physical role	limitation 0-100) <4	months (range	of scores: 0-100; Be	etter indicated	by high	er values	s)		
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	10	10	-	MD 17.5 higher (13.2 lower to 48.2 higher)	⊕OOO VERY LOW	CRITICAL
Without	sciatica - Pain (McGill Que	stionnaire 0-78) <	4 months (Better in	dicated by lowe	er values)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	21	19	-	MD 3.43 lower (9.9 lower to 3.04 higher)	⊕OOO VERY LOW	CRITICAL
Without	sciatica - Pain (VAS 0-10) <	<4 months (Better	indicated by lower	values)	•						
3	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	63	56	-	MD 1.13 lower (1.6 to 0.66 lower)	⊕OOO VERY LOW	CRITICAL
Without	sciatica - Pain (VAS 0-10) 4	I months - 1 year	(Better indicated by	lower values)							
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	47	36	-	MD 0.05 higher (1.07 lower to 1.16 higher)	⊕⊕OO LOW	CRITICAL
Without	sciatica - Funct	ion (ODI 0-1	100) <4 months (E	Better indicated by I	ower values)							
2	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	56	50	-	MD 2.99 lower (5.47 to 0.52 lower)	⊕OOO VERY LOW	CRITICAL
Without	sciatica - Funct	ion (ODQ 0	-100) 4 months - '	l year (Better indica	ted by lower va	lues)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	49	40	-	MD 1.84 lower (8.67 lower to 4.99 higher)	⊕OOO VERY LOW	CRITICAL
Without	sciatica - Psych	nological dis	stress (CESDS 0-	60) <4 months - witl	nout sciatica (B	etter indicated by lov	wer values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ²	none	21	19	-	MD 0.35 higher (2.64 lower to 3.34 higher)	⊕000 VERY	CRITICAL

						LOW	

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias

Table 85: Group aerobic exercise versus self-management in low back pain with or without sciatica

			Quality asse	essment			No	of patients		Effect	Quality	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Group aerobic exercise	Self-management (advice to stay active)	Relative (95% CI)	Absolute	Quality	Importance
Overall -	erall - Quality of life (SF-36 overall health rating 0-100) <4 months (Better indicated by lower values)											
1	randomised trials	very seriousª		no serious indirectness	Serious ^b	none	10	8	-	MD 19.4 higher (3.32 lower to 42.12 higher)	⊕OOO VERY LOW	CRITICAL
Overall -	Pain (0-10) <4	months (Better indicated b	y lower values)								
1	randomised trials	very serious ^a		no serious indirectness	Serious ^b	none	9	9	-	MD 1.85 lower (3.76 lower to 0.06 higher)	⊕OOO VERY LOW	CRITICAL
Overall -	Pain over pre	ceding we	eek (0-10) <4 mont	hs (range of sco	ores: 0-10; Be	etter indicated by	ower values)					
1	randomised trials	very serious ^a		no serious indirectness	Serious ^b	none	9	9	-	MD 1.2 lower (3.12 lower to 0.725 higher)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 86: Group aerobic exercise versus self-management in low back pain without sciatica

Quality assessment	No of patients	Effect	Quality	Importance
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^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Group aerobic exercise	Self-management (advice to stay active)	Relative (95% CI)	Absolute				
Without s	Vithout sciatica - Quality of life individual domain scores(SF-36 0-100) <4 months - Physical role limitation (range of scores: 0-100; Better indicated by lower values)													
1		- ,		no serious indirectness	Serious ^b	none	10	10	-	MD 17.8 higher (15.35 lower to 50.95 higher)	⊕OOO VERY LOW	CRITICAL		
Without s	Without sciatica - Quality of life individual domain scores(SF-36 0-100) <4 months - Physical functioning (range of scores: 0-100; Better indicated by lower values)													
1		- ,		no serious indirectness	Serious ^b	none	10	10		MD 17.3 higher (2.22 lower to 36.82 higher)		CRITICAL		

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 87: Group aerobic exercise versus group biomechanical exercise in low back pain without sciatica

			Quality asse	essment			No	of patients		Effect	0!!		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Group aerobic exercise	Group biomechanical exercise	Relative (95% Absolute CI)		Quality	Importance	
Without -	thout - Pain(VAS 0-10) <4 months (Better indicated by lower values)												
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	32	32	-	MD 1.1 higher (0.15 to 2.05 higher)	⊕⊕OO LOW	CRITICAL	
Without -	Pain (VAS 0-1	0) 4 mon	ths - 1 year (Better	indicated by lov	ver values)								
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	32	32	-	MD 0.4 higher (0.55 lower to 1.35 higher)	⊕⊕OO LOW	CRITICAL	
Without -	Function (OD	I 0-100) <	4 months (Better in	ndicated by lowe	r values)								
1	randomised	serious ¹	no serious	no serious	serious ²	none	32	32	-	MD 6.5 higher (1.27	⊕⊕00	CRITICAL	

	trials		inconsistency	indirectness						to 11.73 higher)	LOW		
Without -	Function (OD	l 0-100) 4	months - 1 year (E	Better indicated I	bv lower valu	ıes)							
1				no serious indirectness	serious ²	none	32	32	-	MD 4.5 higher (0.39 lower to 9.39 higher)	⊕⊕OO LOW	CRITICAL	
Overall - Pain (VAS 0-10) <4 months (Better indicated by lower values)													
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	47	44	-	MD 0.3 higher (0.58 lower to 1.18 higher)	⊕OOO VERY LOW	CRITICAL	
Overall - Pain (VAS 0-10) 4 months - 1 year (Better indicated by lower values)													
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	43	40	-	MD 0.3 higher (0.65 lower to 1.25 higher)	⊕OOO VERY LOW	CRITICAL	
Overall - I	Function (RMI	DQ 0-24) <	<4 months (Better	indicated by low	ver values)								
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ¹	none	47	44	-	MD 0.5 lower (2.52 lower to 1.52 higher)	⊕OOO VERY LOW	CRITICAL	
Overall - I	Function (RMI	DQ 0-24) 4	1 months - 1 year (Better indicated	by lower val	ues)							
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	43	40	-	MD 0.4 higher (1.63 lower to 2.43 higher)	⊕OOO VERY LOW	CRITICAL	

Table 88: Group aerobic exercise versus group biomechanical exercise in low back pain with or without sciatica

			Quality asse	essment			No	of patients		Effect	0	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Group aerobic exercise	Group biomechanical exercise	Relative (95% CI)	Absolute	Quality	Importance

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Overall -	Pain (VAS 0-1	0) <4 mon	ths (Better indicat	ed by lower valu	ies)									
1	randomised trials	very serious¹		no serious indirectness	serious²	none	47	44	-	MD 0.3 higher (0.58 lower to 1.18 higher)	⊕OOO VERY LOW	CRITICAL		
Overall -	Overall - Pain (VAS 0-10) 4 months - 1 year (Better indicated by lower values)													
1	randomised trials	very serious¹		no serious indirectness	serious²	none	43	40	-	MD 0.3 higher (0.65 lower to 1.25 higher)	⊕OOO VERY LOW	CRITICAL		
Overall -	Function (RMI	DQ 0-24) <	<4 months (Better	indicated by low	er values)									
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ¹	none	47	44	-	MD 0.5 lower (2.52 lower to 1.52 higher)	⊕OOO VERY LOW	CRITICAL		
Overall -	Function (RMI	DQ 0-24) 4	4 months - 1 year (Better indicated	by lower val	ues)								
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious²	none	43	40	-	MD 0.4 higher (1.63 lower to 2.43 higher)	⊕OOO VERY LOW	CRITICAL		

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Individual mind-body exercise J.5.1.5

Table 89: Individual mind-body exercise versus individual biomechanical exercise in low back pain with or without sciatica

			Quality ass	sessment			No of patients			Effect	Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Individual mind-body exercise versus individual biomechanical exercise		Relative (95% CI)	Absolute	Quanty	importance	
Overall-Fu	overall-Function (RMDQ) <4 months (range of scores: 0-23; Better indicated by lower values)												

1	randomised trials			no serious indirectness	Serious ^b	none	15	15	-	MD 5.18 lower (9.27 to 1.09 lower)	⊕⊕OO LOW	CRITICAL		
Tai Chi, d	overall-Pain (\	/AS 0-10)	<4 months (range	e of scores: 0-10	; Better indicat	ed by lower value	s)							
1		- ,			no serious imprecision	none	20	20	1	MD 0.7 lower (1.01 to 0.39 lower)	⊕⊕OO LOW	CRITICAL		
Yoga, ov	Yoga, overall-Pain (VAS 0-10) <4 months (range of scores: 0-10; Better indicated by lower values)													
1		- ,			no serious imprecision	none	15	15	-	MD 2.63 lower (3.48 to 1.24 lower)	⊕⊕OO LOW	CRITICAL		

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Group mind-body exercise J.5.1.6

Table 90: Group mind-body exercise versus usual care in low back pain with or without sciatica

				Jus usuai cai c		<u> </u>								
			Quality as:	sessment			No of patie	ents		Effect	Quality	Importance		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Group mind- body exercise	Usual care	ADSOILITE					
Overall - 0	rerall - Quality of life (EQ-5D 0-1) <4 months (Better indicated by lower values)													
2	randomised trials			no serious indirectness	Serious ^b	none	160	165	-	MD 0.06 higher (0.01 to 0.1 higher)	⊕⊕OO LOW	CRITICAL		
Overall Q	uality of life (l	EQ-5D 0-1) 4 months - 1 yea	ar (Better indicat	ed by lower val	ues)								
1	randomised trials				no serious imprecision	none	156	157	-	MD 0.02 higher (0.03 lower to 0.07 higher)	⊕⊕⊕O MODERATE	CRITICAL		
Overall - 0	Quality of life	(SF-12 0-1	100) <4 months - I	Physical compor	nent (Better indi	icated by lower va	ılues)							

			,	•		_	•			
2	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	160	166	-	MD 1.12 higher (1.1 ⊕⊕⊕O CRITICAL lower to 3.34 higher) MODERATE
Overall -	Quality of life	(SF-12 0-	100) <4 months -	Mental compone	ent (Better indic	ated by lower valu	ies)			
2	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	160	166	-	MD 2.05 higher (0.47 ⊕⊕⊕O CRITICAL lower to 4.56 higher) MODERATE
Overall -	Quality of life	(SF-12 0-	100) >4 months -	1 year (Better in	dicated by lowe	r values)				
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	156	157	-	MD 0.79 higher (1.49 ⊕⊕⊕⊖ CRITICAI lower to 3.07 higher) MODERATE
Overall -	Quality of life	(SF-12 0-	100) >4 months -	1 year (Better in	dicated by lowe	r values)				
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	156	157	-	MD 0.42 higher (2.16 ⊕⊕⊕⊖ CRITICAI lower to 3 higher) MODERATE
Overall -	Pain (VAS 0-1	0) <4 moi	nths - Hatha yoga	(range of score	s: 0-10; Better i	ndicated by lower	values)			
2	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	40	42	-	MD 0.88 lower (2.61 ⊕OOO CRITICAL lower to 0.85 higher) VERY LOW
Overall -	Pain (VAS 0-1	l0) <4 moi	nths - lyengar yog	ga (Better indica	ted by lower val	ues)				
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	43	47	-	MD 0.43 lower (1.21 ⊕OOO CRITICAL lower to 0.35 higher) VERY LOW
Overall -	Pain (VAS 0-1	0) 4 mont	hs - 1 year - Hath	a yoga (Better ir	ndicated by low	er values)	<u> </u>			
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	8	15	-	MD 0.6 lower (1.34 ⊕⊕OO CRITICAL lower to 0.14 higher) LOW
Overall -	Pain (VAS 0-1	(0) 4 mont	ths - 1 year - lyen	gar yoga (Better	indicated by lo	wer values)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	43	47	-	MD 1.08 lower (1.93 to 0.23 lower) ⊕OOO VERY LOW CRITICAL
Overall -	Pain (Aberde	en pain so	cale 0-100) <4 mo	nths (Better indi	cated by lower	values)				
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	156	157	-	MD 2.42 lower (5.21 ⊕⊕⊕O CRITICAI lower to 0.37 higher) MODERATE

Pain (Aberde	en pain so	cale 0-100) >4 mor	nths - 1 year (Be	etter indicated b	y lower values)						
randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	156	157	-	MD 0.72 lower (3.53 lower to 2.09 higher)	⊕⊕⊕O MODERATE	CRITICAL
Function (RM	DQ/ODI) <	<4 months - Yoga	(Better indicate	d by lower value	es)						
randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	255	261	-	SMD 0.34 lower (0.52 to 0.17 lower)	⊕⊕OO LOW	CRITICAL
Function (RM	DQ/ODI) 4	4 months - 1 year	(Better indicated	d by lower value	es)						
randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	207	219	-	SMD 0.3 lower (0.5 to 0.11 lower)	⊕⊕OO LOW	CRITICAL
Psychological	distress	(BDI 0-63) <4 mor	nths (Hatha) (Be	tter indicated by	lower values)		-				
randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	11	5	-	MD 10.18 lower (19.68 to 0.68 lower)	⊕000 VERY LOW	CRITICAL
Psychological	distress	(BDI 0-63) <4 mor	nths (Iyengar) (B	Setter indicated	by lower values)					'	
randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	43	47	-	MD 1.5 lower (3.94 lower to 0.94 higher)	⊕000 VERY LOW	CRITICAL
Psychologica	l distress	(BDI 0-63) 4 mon	ths - 1 year (Bet	ter indicated by	lower values)		•				
randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	43	47	-	MD 2.6 lower (4.7 to 0.5 lower)	⊕000 VERY LOW	CRITICAL
Responder cr	riteria (imi	provement in pair	n) <4 months			_	<u>'</u>	<u> </u>			
randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	37/80 (46.3%)	12/80 (15%)	RR 3.08 (1.74 to 5.47)	312 more per 1000 (from 111 more to 670 more)		IMPORTAN
Responder cr	iteria (imi	orovement in fund	ction) <4 months				<u>+</u>				
		no serious inconsistency	no serious indirectness	no serious imprecision	none	40/80 (50%)	19/80 (23.8%)	RR 2.11 (1.34 to 3.3)	264 more per 1000 (from 81 more to 546 more)	⊕⊕⊕O MODERATE	IMPORTAN
	randomised trials Function (RM randomised trials Function (RM randomised trials Psychological randomised trials Psychological randomised trials Psychological randomised trials Responder criandomised trials Responder criandomised trials	randomised trials Function (RMDQ/ODI) randomised trials Function (RMDQ/ODI) randomised trials Function (RMDQ/ODI) randomised trials Psychological distress randomised trials Responder criteria (important) randomised trials Responder criteria (important) Responder criteria (important)	randomised trials Function (RMDQ/ODI) <4 months - Yoga randomised trials Seriousa no serious inconsistency Function (RMDQ/ODI) 4 months - 1 year randomised trials Seriousa no serious inconsistency Psychological distress (BDI 0-63) <4 more randomised trials Seriousa no serious inconsistency Psychological distress (BDI 0-63) <4 more randomised trials Seriousa no serious inconsistency Psychological distress (BDI 0-63) <4 more randomised trials Seriousa no serious inconsistency Psychological distress (BDI 0-63) 4 more randomised trials Seriousa no serious inconsistency Responder criteria (improvement in pair randomised trials Seriousa no serious inconsistency Responder criteria (improvement in pair randomised trials Seriousa no serious inconsistency	randomised trials Serious ^a no serious inconsistency indirectness Function (RMDQ/ODI) <4 months - Yoga (Better indicate no serious inconsistency indirectness) Function (RMDQ/ODI) 4 months - 1 year (Better indicate no serious inconsistency indirectness) Function (RMDQ/ODI) 4 months - 1 year (Better indicate no serious inconsistency indirectness) Psychological distress (BDI 0-63) <4 months (Hatha) (Betrandomised trials Psychological distress (BDI 0-63) <4 months (Iyengar) (Betrandomised trials Psychological distress (BDI 0-63) <4 months (Iyengar) (Betrandomised trials Psychological distress (BDI 0-63) <4 months (Iyengar) (Betrandomised trials Psychological distress (BDI 0-63) 4 months - 1 year (Betrandomised trials Psychological distress (BDI 0-63) 4 months - 1 year (Betrandomised trials Psychological distress (BDI 0-63) 4 months - 1 year (Betrandomised trials Psychological distress (BDI 0-63) 4 months - 1 year (Betrandomised trials Responder criteria (improvement in pain) <4 months randomised Serious ^a no serious indirectness Responder criteria (improvement in function) <4 months randomised Serious ^a no serious indirectness	randomised trials Seriousa no serious indirectness no serious imprecision Function (RMDQ/ODI) <4 months - Yoga (Better indicated by lower value randomised trials Seriousa no serious indirectness Function (RMDQ/ODI) 4 months - 1 year (Better indicated by lower value randomised trials Function (RMDQ/ODI) 4 months - 1 year (Better indicated by lower value randomised trials Psychological distress (BDI 0-63) <4 months (Hatha) (Better indicated by randomised trials Psychological distress (BDI 0-63) <4 months (Hatha) (Better indicated by randomised trials Psychological distress (BDI 0-63) <4 months (Iyengar) (Better indicated Irandomised trials Psychological distress (BDI 0-63) <4 months (Iyengar) (Better indicated Irandomised trials Psychological distress (BDI 0-63) 4 months - 1 year (Better indicated by randomised very serious indirectness Psychological distress (BDI 0-63) 4 months - 1 year (Better indicated by randomised very serious indirectness Responder criteria (improvement in pain) <4 months randomised Seriousa no serious indirectness inprecision Responder criteria (improvement in function) <4 months randomised Seriousa no serious indirectness inprecision Responder criteria (improvement in function) <4 months	trials inconsistency indirectness imprecision Function (RMDQ/ODI) <4 months - Yoga (Better indicated by lower values) randomised trials Seriousa no serious inconsistency no serious indirectness Seriousb none randomised trials Seriousa no serious indirectness very seriousb none randomised trials Seriousa no serious indirectness very seriousb none randomised trials Seriousa no serious indirectness very seriousb none randomised trials Seriousa no serious indirectness very seriousb none randomised trials Seriousa no serious indirectness no serious indirectness randomised Seriousa no serious indirectness no serious indirectness randomised Seriousa no serious no serious indirectness indirectness randomised Seriousa no serious no serious indirectness indirectness randomised Seriousa no serious no serious indirectness no serious indirectness randomised Seriousa no serious no	randomised trials Function (RMDQ/ODI) <4 months - Yoga (Better indicated by lower values) randomised trials Seriousa no serious inconsistency randomised Seriousa no serious indirectness randomised trials Seriousa no serious indirectness randomised very seriousa no serious indirectness randomised very serious no serious indirectness randomised Serious no serious indirectness randomised Serious no serious indirectness randomised Serious no serious indirectness indirectness randomised Serious no serious indirectness indirectness randomised Serious no serious indirectness indirectness indirectness randomised Serious no serious indirectness	randomised brials Serious no serious no serious indirectness no serious indirectness no serious indirectness no serious indirectness imprecision none 156 157 Function (RMDQ/ODI) <4 months - Yoga (Better indicated by lower values) randomised Serious no serious no serious no serious none 255 261 Function (RMDQ/ODI) 4 months - 1 year (Better indicated by lower values) randomised Serious no serious no serious indirectness Serious none 207 219 randomised Serious no serious no serious indirectness very serious none 11 5 randomised Serious no serious indirectness very serious none 11 5 Psychological distress (BDI 0-63) <4 months (Iyengar) (Better indicated by lower values) randomised Serious no serious indirectness very serious none 43 47 randomised Serious no serious indirectness Serious none 43 47 randomised Very no serious indirectness Serious none 43 47 randomised Very no serious no serious indirectness Serious none 43 47 Responder criteria (improvement in pain) <4 months randomised Serious no serious no serious none 37/80 12/80 randomised Serious no serious no serious none indirectness indirectness indirectness indirectness indirectness indirectness indirectness none 12/80 Responder criteria (improvement in function) <4 months randomised Serious no serious no serious none 19/80 1	randomised Serious® no serious inconsistency indirectness imprecision none 156 157 - Function (RMDQ/ODI) <4 months - Yoga (Better indicated by lower values) randomised Serious® no serious indirectness Serious® none 255 261 - Function (RMDQ/ODI) 4 months - 1 year (Better indicated by lower values) Function (RMDQ/ODI) 4 months - 1 year (Better indicated by lower values) randomised Serious® no serious inconsistency indirectness Serious® none 207 219 - randomised Serious® no serious indirectness indirectness inconsistency indirectness inconsistency indirectness inconsistency indirectness very serious® none 11 5 - Psychological distress (BDI 0-63) <4 months (Hatha) (Better indicated by lower values) randomised Serious® no serious inconsistency indirectness very serious® none 11 5 - Psychological distress (BDI 0-63) <4 months (Hyengar) (Better indicated by lower values) randomised Serious® no serious inconsistency indirectness very serious® none 43 47 - Psychological distress (BDI 0-63) 4 months - 1 year (Better indicated by lower values) randomised very no serious inconsistency indirectness Serious® none 43 47 - Responder criteria (improvement in pain) <4 months randomised Serious® no serious inconsistency indirectness imprecision none 37/80 (46.3%) (15%) (1.74 to 5.47) Responder criteria (improvement in function) <4 months randomised Serious® no serious no serious no serious no serious none and serious inconsistency indirectness none 40/80 19/80 RR 2.11	randomised Serious no serious indirectness no serious indirectness Function (RMDQ/ODI) <4 months - Yoga (Better indicated by lower values) randomised Serious no serious indirectness no serious indirectness randomised Serious no serious indirectness randomised Serious no serious indirectness Responder criteria (improvement in pain) <4 months randomised Serious no serious indirectness indi	trandomised Serious* no serious indirectness imprecision none 156 157 - MD 0.72 lower (3.53 ⊕⊕⊕⊕ MODERATE Function (RMDQ/ODI) <4 months - Yoga (Better indicated by lower values) randomised Serious* no serious inconsistency indirectness serious inconsistency indirectness indirectness serious inconsistency indirectness serious inconsistency indirectness indirectness serious inconsistency indirectness indirectness serious inconsistency indirectness indirectness inconsistency indirectness indirectness inconsistency indirectness inconsiste

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1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	5	9	-	MD 0.73 lower (2.49 lower to 1.03 higher)	⊕OOO VERY LOW	IMPORTANT			
Overall -	Healthcare ut	ilisation -	Practice nurse vi	sits <4 months (Better indicated	by lower values)									
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	5	9	-	MD 0.11 lower (0.44 lower to 0.22 higher)	⊕000 VERY LOW	IMPORTANT			
Overall -	Healthcare ut	ilisation -	physiotherapist v	visits <4 months	(Better indicate	ed by lower values)								
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	5	9	-	MD 0.33 lower (1.33 lower to 0.67 higher)	⊕OOO VERY LOW	IMPORTANT			
Overall -	Healthcare ut	ilisation -	Medication use <	4 months (Viniye	oga)										
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	4/5 (80%)	6/9 (66.7%)	RR 1.2 (0.63 to 2.27)	133 more per 1000 (from 247 fewer to 847 more)	⊕000 VERY LOW	IMPORTANT			
Overall -	verall - Healthcare utilisation - Medication use <4 months (Hatha)														
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	2/15 (13.3%)	11/15 (73.3%)	RR 0.18 (0.05 to 0.68)	601 fewer per 1000 (from 235 fewer to 697 fewer)	⊕⊕OO LOW	IMPORTANT			
Overall -	Healthcare ut	ilisation -	Reduced or stop	ped medication	<4 months										
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	14/20 (70%)	6/24 (25%)	RR 2.8 (1.32 to 5.93)	450 more per 1000 (from 80 more to 1000 more)	⊕⊕OO LOW	IMPORTANT			
Overall -	Healthcare ut	ilisation -	Reduced or stop	ped medication	>4 months - 1 y	ear		•							
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	10/20 (50%)	15/22 (68.2%)	RR 0.73 (0.43 to 1.24)	184 fewer per 1000 (from 389 fewer to 164 more)	⊕000 VERY LOW	IMPORTANT			
Without s	sciatica - Pain	(VAS 0-1	0) <4 months (Be	ter indicated by	lower values)										
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	20	22	-	MD 1.1 lower (2.18 to 0.02 lower)	⊕OOO VERY LOW	CRITICAL			
Without s	sciatica - Pain	(VAS 0-1	0) >4 months - 1 չ	ear (Better indic	ated by lower v	values)									

trials serious inconsistency indirectness ochous indirectness					no serious indirectness	Serious ^b	none	20	22	-	MD 1.4 lower (2.4 to 0.4 lower)	0000	CRITICAL
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^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias

Table 91: Group mind-body exercise versus usual care in low back pain without sciatica

			Quality asses	ssment			No of patien	ts		Effect	Quality	Immontono	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Group mind-body exercise	Usual care	Relative (95% CI)	Absolute	Quality	Importance	
Without sc	Vithout sciatica - Pain (VAS 0-10) <4 months (Better indicated by lower values)												
1		,	no serious inconsistency	no serious indirectness	Serious ^b	none	20	22	-	MD 1.1 lower (2.18 to 0.02 lower)	⊕OOO VERY LOW	CRITICAL	
Without sc	iatica - Pain (V	'AS 0-10) >	4 months - 1 year (E	Better indicated by	/ lower value	es)							
		,	no serious inconsistency	no serious indirectness	Serious ^b	none	20	22	-	MD 1.4 lower (2.4 to 0.4 lower)	⊕OOO VERY LOW	CRITICAL	

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias

Table 92: Group mind-body exercise versus self-management in low back pain without sciatica

			Quality ass	sessment			No	of patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Group mind- body exercise	Self management (advice to stay active)	Relative (95% CI)	Absolute	Quanty	Importance

^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Function	unction (RMDQ 0-24) <4 months - without sciatica (Better indicated by lower values)														
2	randomised	very	no serious inconsistency	no serious indirectness	no serious imprecision	none	81	44	-	MD 2.78 lower (3.76 to 1.81 lower)	⊕⊕OO LOW	CRITICAL			
Without	Nithout - Function (RMDQ 0-24) 4 months - 1 year - without sciatica (Better indicated by lower values)														
2		very serious ^a	Serious ^b	no serious indirectness	Serious ^c	none	83	81	-	MD 2.60lower (4.34 to 0.85 lower)	⊕OOO VERY LOW	CRITICAL			
Without	ithout - Responder criteria (improvement in function) 4 months - 1 year - without sciatica														
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^c	none	0/81 (0%)	0%	RR 1.67 (1.17 to 2.38)	-	⊕⊕OO LOW	IMPORTANT			
Healthca	re utilisation	- medicat	ion use >4 mont	ns - 1 year - with	nout sciatica										
1		very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	7/34 (20.6%)	17/29 (58.6%)	RR 0.35 (0.17 to 0.73)	381 fewer per 1000 (from 158 fewer to 487 fewer)	⊕⊕OO LOW	IMPORTANT			

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Heterogeneity, I²=88%, unexplained by subgroup analysis. ³ Downgraded by 1 increment if the confidence interval crossed 1 MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 93: Group mind-body exercise versus group mixed exercise in low back pain without sciatica

	. C. Cup.		dy chereise vei	Sas S. Sab	ACG CACICIOC	m lett back pa		- Cia ti Ga				
			Quality as:	sessment			No of pa	atients		Effect	Ovality	Immortono
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Group mind- body exercise	Group mixed exercise	Relative (95% CI)	Absolute	Quality	Importance
Without s	ciatica - Fun	ction (RM	DQ 0-24) <4 mont	hs (Better indica	ated by lower v	alues)						
	randomised trials	Serious ^a		no serious indirectness	Serious ^b	none	117	111	-	MD 0.89 lower (2.32 lower to 0.55 higher)		CRITICAL

Without s	Nithout sciatica - Function (RMDQ 0-24) 4 months - 1 year (Better indicated by lower values)														
			no serious	no serious indirectness	no serious imprecision	none	117	112		MD 0.72 lower (1.68 lower to 0.24 higher)	⊕⊕⊕O MODERATE	CRITICAL			
Without	Vithout sciatica - Responder criteria (improvement in function) <4 months														
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	0/81 (0%)	0%	RR 1.06 (0.87 to 1.29)	-	⊕⊕OO LOW	IMPORTANT			
Without s	sciatica - Hea	Ithcare ut	ilisation - medica	tion use 4 mont	hs - 1 year - He	althcare utilisation	n - medication	use 4 months	s - 1 year						
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	7/34 (20.6%)	16/32 (50%)		295 fewer per 1000 (from 65 fewer to 400 fewer)	⊕⊕OO LOW	IMPORTANT			

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs ^c Heterogeneity, I²=55%, unexplained by subgroup analysis.

Table 94: Group mind-body exercise versus individual biomechanical exercise in low back pain with or without sciatica

			Quality as:	sessment			No of patients			Effect	0	I
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Group mind-body exercise versus individual biomechanical exercise	Control	Relative (95% CI)		Quality	Importance
Overall-Pa	ain (VAS) - <4	months (range of scores:	0-10; Better indi	cated by lower	values)						
	randomised trials				no serious imprecision	none	30	30	-	MD 1.5 lower (1.96 to 1.04 lower)	⊕⊕⊕⊕ HIGH	CRITICAL
Overall-P	ain (VAS) - >4	months ((range of scores:	0-10; Better indi	cated by lower	values)		•			•	
	randomised trials				no serious imprecision	none	30	30	-	MD 2 lower (2.47 to 1.53 lower)	⊕⊕⊕⊕ HIGH	CRITICAL

a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias

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Table 95: Individual mixed exercise versus waiting list in low back pain with sciatica

			Quality asses	ssment			No of patier	nts		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Individual mixed exercise	Waiting list	Relative (95% CI)			
Pain (VAS,	0-10) < 4 mon	ths (range	of scores: 0-10; Bet	ter indicated by lo								
1	randomised trials			no serious indirectness	Serious⁵	none	15	15	-	MD 2.34 lower (4.02 to 0.66 lower)	⊕⊕OO LOW	CRITICAL
With sciation	ca - Leg pain (VAS 0-10)	< 4 months (Better i	ndicated by lower	values)							
1	randomised trials			no serious indirectness	Serious ^b	none	15	15	-	MD 3 lower (5.06 to 0.94 lower)	⊕⊕OO LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

Table 96: Individual mixed exercise versus unsupervised exercise in low back pain with or without sciatica

			Quality as	sessment			No of	patients		Effect	Quality	Immontono
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Individual mixed exercise	Unsupervised exercise	Relative (95% CI)	Absolute	Quality	Importance
Overall - F	Pain (VAS 0-10)) 4 month	ıs - 1 year (range o	of scores: 0-10; B	etter indicated l	by lower values)						
		- ,			no serious imprecision	none	20	20	-	MD 4.65 lower (5.44 to 3.86 lower)		CRITICAL

a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias

Table 97: Individual mixed exercise versus individual biomechanical exercise in low back pain with or without sciatica

Table 37	· illaiviac	iai iiiixca i	CACICISC VCISUS	illaiviadai bi	Officerialing	car exercise iii	iow back pain with or	WILLIOU	it sciati	ca		
			Quality asses	ssment			No of patients			Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Individual mixed exercise versus biomechanical	Control	Relative ntrol (95% Absolute CI)		Quality	Importance
Overall-fu	ınction (ODI)	<4 months (range of scores: ()-100; Better ind	icated by lov	ver values)						
1				no serious indirectness	Seriousª	none	31	32	-	MD 2.8 lower (5.52 to 0.08 lower)	⊕⊕⊕O MODERATE	CRITICAL
Overall-Pain (VAS 0-10) <4 months (range of scores: 0-10; Better indicated by lower values)												
				no serious indirectness	Serious ^a	none	31	32	-	MD 0.3 lower (0.83 lower to 0.23 higher)	⊕⊕⊕O MODERATE	CRITICAL

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias

J.5.1.8 Group mixed exercise

Table 98: Group mixed exercise versus usual care in low back pain with or without sciatica

			Quality as	sessment			No of patie	nts		Effect	Ovality.	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Group mixed exercise	Usual care	I 195% I ANSOINTE		Quality	Importance
Overall - Pain (VAS 0-10) <4 months (Better indicated by lower values)												
	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious⁵	none	84	78	-	MD 1.15 lower (1.8 to 0.49 lower)	⊕⊕OO LOW	CRITICAL

verall-P	ain (VAS) <4	l months -	Pain at flexion	(range of score	es: 0-10: Better i	indicated by low	er values)					
	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	21	17	-	MD 5.21 lower (5.48 to 4.94 lower)	⊕⊕⊕O MODERATE	CRITICA
verall-P	ain (VAS) <4	l months -	Pain at rest (ra	inge of scores:	0-10; Better ind	icated by lower	values)					
	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	21	17	-	MD 4.05 lower (4.31 to 3.79 lower)	⊕⊕⊕O MODERATE	CRITICA
erall -	Pain (VAS 0-	-10) 4 mon	ths - 1 year (Be	etter indicated b	y lower values)							
	randomised trials	very serious ^a	very serious ^c	no serious indirectness	very serious ^b	none	49	43	-	MD 2.55 lower (6.73 lower to 1.64 higher)	⊕OOO VERY LOW	CRITICA
verall -	Pain (von Ko	orff 0-100)	<4 months [me	an difference fr	om control] (Be	etter indicated by	/ lower values)					
	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	14	13	-	MD 0.88 lower (2.26 lower to 0.5 higher)	⊕⊕OO LOW	CRITICA
/erall -	Pain (von Ko	orff 0-100)	4 months - 1 ye	ear - Pain (von h	Corff 0-100) (Bet	ter indicated by	lower values)					
	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	14	13	-	MD 0.15 higher (1.34 lower to 1.63 higher)	⊕OOO VERY LOW	CRITICA
erall -	Function (RI	MDQ 0-24)	<4 months (Be	etter indicated b	y lower values)							
	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	84	78	-	MD 2.02 lower (3.48 to 0.55 lower)	⊕⊕OO LOW	CRITICA
/erall -	Function (RI	MDQ 0-24)	4 months - 1 y	ear (Better indic	cated by lower v	values)						
	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	29	23	-	MD 0.57 lower (3.45 lower to 2.31 higher)	⊕OOO VERY LOW	CRITICA
/erall -	Function (RI	MDQ 0-24)	<4 months [me	ean difference fi	rom control) (Be	etter indicated b	y lower values)					
	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	14	13	-	MD 1.91 lower (5.41 lower to 1.6 higher)	⊕⊕OO LOW	CRITICA
erall -	Function (RI	MDQ 0-24)	4 months - 1 y	ear [mean differ	ence from cont	rol] (Better indic	ated by lower va	alues)				

1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	14	13	-	MD 3 lower (6.88 lower to 0.88 higher)	⊕⊕OO LOW	CRITICAL		
Overall-	SF-36 (0-100) <4 month	ns - Physical (ra	ange of scores:	0-100; Better in	dicated by highe	er values)							
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	21	17	-	MD 1 lower (2.1 lower to 0.1 higher)	⊕⊕OO LOW	CRITICAL		
Overall-	Overall- SF-36 (0-100) <4 months - Mental (range of scores: 0-100; Better indicated by higher values)													
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	21	17	-	MD 4.5 higher (2.89 to 6.11 higher)	⊕⊕⊕O MODERATE	CRITICAL		
Overall -	Overall - Psychological distress (BDI 0-63) (Better indicated by lower values)													
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	52	50	-	MD 2.09 lower (3.86 to 0.32 lower)	⊕⊕OO LOW	CRITICAL		

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs ^c Heterogeneity, I²=97% unexplained by subgroup analysis

Table 99: Group mixed exercise versus usual care in low back pain with sciatica

						ann with solut						
			Quality as	sessment			No of patie	nts		Effect	O. alife	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Group mixed exercise	Usual care	Relative (95% CI)	Absolute	Quality	Importance
With scia	tica - Pain (\	VAS/NRS 0										
	randomised trials	Serious ^a	no serious inconsistency		no serious imprecision	none	27	26	-	MD 2.59 lower (3.11 to 2.07 lower)	⊕⊕⊕O MODERATE	CRITICAL
With sciatica - Pain (VAS/NRS 0-10) <4 months - Pain on movement (range of scores: 0-10; Better indicated by lower values)												
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	27	26	-	MD 2.47 lower (3 to 1.94 lower)	⊕⊕⊕O MODERATE	CRITICAL

	randomised ve	,		no serious indirectness	Serious ^b	none	25	25	-	MD 0.7 lower (1.48 lower to 0.08 higher)	⊕OOO VERY LOW	CRITICAL	
Nith sciatica - Pain (NRS 0-10) 4 months - 1 year (Better indicated by lower values)													
	randomised ve	,		no serious indirectness	very serious ^b	none	23	21	-	MD 2.3 lower (3.17 to 1.43 lower)	⊕OOO VERY LOW	CRITICAL	
Vith sciatica Function (RMDQ 0-24) <4 months (Better indicated by lower values)													
	randomised ve	,		no serious indirectness	Serious ^b	none	23	21	-	MD 1.2 higher (0.43 to 1.97 higher)	⊕OOO VERY LOW	CRITICAL	
With sciatica - Function (RMDQ 0-24) 4 months - 1 year (Better indicated by lower values)													
Vith so	iatica - Function	n (RMDQ	U-24) 4 months	s - Tyear (Delle	i iliulcateu by i	ower values,							

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 100: Group mixed exercise versus usual care in low back pain without sciatica

			Quality as	sessment			No of patie	nts		Effect	Quality	Immoutonee
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Group mixed exercise	Usual care	Relative (95% Absolute CI)		Quality	Importance
Without	sciatica - Qu	ality of life	(SF-36 0-100) <	<4 months - gen	eral health (Bet	ter indicated by	lower values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious⁵	none	16	20	-	MD 3.8 higher (2.31 lower to 9.91 higher)	⊕OOO VERY LOW	CRITICAL
Without s	sciatica - Qu	ality of life	(SF-36 0-100) <	<4 months - vital	lity (Better indi	cated by lower v	alues)					

		,										
1	randomised trials	very seriousª	no serious inconsistency	no serious indirectness	very serious ^b	none	16	20	-	MD 0.1 higher (9.47 lower to 9.67 higher)	⊕OOO VERY LOW	CRITICAL
Without	sciatica - Qu	ality of life	(SF-36 0-100)	<4 months - phy	sical functionir	ng (Better indica	ted by lower val	ues)				
1	randomised trials	very seriousª	no serious inconsistency	no serious indirectness	very serious ^b	none	16	20	-	MD 0.5 higher (5.88 lower to 6.88 higher)	⊕OOO VERY LOW	CRITICAL
Without	sciatica - Qu	ality of life	score (SF-36 ()-100) <4 month	s - Pain (Better	indicated by lov	ver values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	16	20	-	MD 2.1 higher (6.92 lower to 11.12 higher)	⊕OOO VERY LOW	CRITICAL
Without	sciatica - Qu	ality of life	(SF-36 0-100)	<4 months - phy	sical role limita	ation (Better ind	cated by lower v	/alues)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	16	20	-	MD 12.7 higher (53.17 lower to 78.57 higher)	⊕OOO VERY LOW	CRITICAL
Without	sciatica - Qu	ality of life	(SF-36 0-100)	<4 months - em	otional role limi	tation (Better in	dicated by lower	values)			
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	16	20	-	MD 7.4 higher (12.66 lower to 27.46 higher)	⊕⊕OO LOW	CRITICAL
Without	sciatica - Qu	ality of life	(SF-36 0-100)	<4 months - soc	ial functioning	(Better indicate	d by lower value	s)				
1	randomised trials		no serious	no serious indirectness	very serious ^b	none	16	20	-	MD 1.2 lower (11.2 lower to 8.8 higher)	⊕OOO VERY LOW	CRITICAL
Without s	sciatica - Qu	ality of life	(SF-36 0-100)	<4 months - me	ntal health (Bet	ter indicated by	lower values)	<u> </u>	!			
1	randomised trials		no serious inconsistency	no serious indirectness	very serious ^b	none	16	20	-	MD 0.9 lower (6.94 lower to 5.14 higher)	⊕OOO VERY LOW	CRITICAL
Without	sciatica - Pa	in (VAS 0-1	10) <4 months (Better indicated	l by lower value	es)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	16	13	-	MD 0.95 lower (1.1 to 0.8 lower)	⊕⊕OO LOW	CRITICAL
Without	sciatica - Pa	in (VAS 0-1	10, change sco	re) <4 months (I	Better indicated	by lower values	<u> </u>					
1	randomised trials	very serious ²	no serious inconsistency	no serious indirectness	very serious ¹	none	30	29	-	MD 4.9 lower (15.73 lower to 5.93 higher)	⊕OOO VERY LOW	CRITICAL

Without s	sciatica - Fu	nction (OD	I/RMDQ, chang	e score) <4 moi	nths (Better ind	licated by lower	r values)							
2	randomised trials	very serious²		no serious indirectness	serious ¹	none	46	42	-	SMD 0.66 lower (1.09 to 0.22 lower)	⊕000 VERY LOW	CRITICAL		
Without	Without sciatica - Psychological distress (HADS 0-21) <4 month - anxiety score (Better indicated by lower values)													
1	randomised trials	very serious²		no serious indirectness	very serious ¹	none	16	13	-	MD 0.55 lower (2.21 lower to 1.11 higher)	⊕OOO VERY LOW	CRITICAL		
Without	sciatica - Ps	ychologica	I distress (HAD	S 0-21) <4 mon	th - depression	score (Copy) (Better indicated	by lowe	r values)					
1	randomised trials	very serious²		no serious indirectness	serious ¹	none	16	13	-	MD 0.99 lower (2.39 lower to 0.41 higher)	⊕000 VERY LOW	CRITICAL		

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 101: Group mixed exercise versus self-management in low back pain without sciatica

			Quality as:	sessment			No	of patients		Effect	Quality	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Group mixed exercise	Self-management (advice to stay active)	Relative (95% CI)	Absolute	Quality	Importance
Without	sciatica - Res	ponder c	riteria (improvem	ent in function)) 4 months - 1 y	year						
	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	0/81 (0%)	0/44 (0%)	RR 1.58 (1.1 to 2.27)	-	⊕⊕OO LOW	IMPORTANT
								0%		-		
Without	sciatica - Fun	ction (RM	IDQ 0-24) <4 mor	nths (Better indi	cated by lower	r values)						
	randomised trials		no serious inconsistency		no serious imprecision	none	81	44	-	MD 1.99 lower (2.96 to 1.02 lower)	⊕⊕⊕O MODERATE	CRITICAL
Without	sciatica - Fun	ction (RM	IDQ 0-24) 4 mont	hs - 1 year - wit	hout sciatica (Better indicated b	y lower valu	es)				

	randomised trials			no serious indirectness	Serious ^b	none	83	81	-	MD 1.65 lower (2.72 to 0.57 lower)	⊕⊕OO LOW	CRITICAL		
Without	Vithout sciatica - Healthcare utilisation - medication use 4 months - 1 year													
	randomised trials			no serious indirectness	very serious ^b	none	16/32 (50%)	17/29 (58.6%)		88 fewer per 1000 (from 270 fewer to 205 more)		IMPORTANT		

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 102: Group mixed exercise versus cognitive therapy in low back pain without sciatica

			Quality asse	essment			No of p	atients		Effect	Ouglitu	Immontono
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Group mixed exercise	Placebo/sham	Relative (95% CI)	Absolute	Quanty	Importance
Without s	ciatica - Pain (VAS 0-10)	<4 months (Better	indicated by low	er values)							
1	randomised trials		no serious inconsistency	no serious indirectness	very serious ^b	none	10	11	-	MD 1.8 lower (5.16 lower to 1.56 higher)	⊕OOO VERY LOW	CRITICAL
Without s	ciatica - Pain (VAS 0-10)	4 months - 1 year	(Better indicated	by lower valu	ıes)						
1	randomised trials		no serious inconsistency	no serious indirectness	very serious ^b	none	14	13	-	MD 1.3 lower (4.4 lower to 1.8 higher)	⊕OOO VERY LOW	CRITICAL
Without s	ciatica - Funct	ion (RMDC	Q 0-24) <4 months	- without sciatica	(Better indic	ated by lower valu	es)					
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	10	11	1	MD 4.9 lower (9.08 to 0.72 lower)	⊕⊕OO LOW	CRITICAL
Without s	ciatica - Psych	nological d	istress (BDI 0-63)	<4 months (Bette	r indicated by	y lower values)						
1	randomised	Serious ^a	no serious	no serious	Serious ^b	none	10	11	-	MD 6.3 lower (18.7	⊕⊕00	CRITICAL

	trials	inconsistency	indirectness			lower to 6.1 higher)	LOW	

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 103: Group mixed exercise versus cognitive behavioural approaches in low back pain with or without sciatica

			Quality as	sessment			No o	of patients		Effect	Quality	Immoutonce
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Group mixed exercise	cognitive behavioural approaches	Relative (95% CI)	Absolute	Quality	Importance
With/with	out sciatica -	Pain (VA	S 0-10) <4 month	s (Better indicat	ed by lower val	ues)						
	randomised trials		no serious inconsistency	no serious indirectness	serious ^b	none	52	55	-	MD 0.56 lower (1.48 lower to 0.36 higher)	⊕⊕OO LOW	CRITICAL
With/with	out sciatica -	Pain (VA	S 0-10) >4 month	s (Better indicat	ted by lower val	ues)						
	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	51	52	-	MD 0.09 lower (1.02 lower to 0.84 higher)	⊕OOO VERY LOW	CRITICAL
With/with	out sciatica -	Function	ı (RMDQ) <4 mon	ths (Better indic	ated by lower v	alues)						
	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	52	55	-	MD 0.62 lower (2.4 lower to 1.16 higher)	⊕OOO VERY LOW	CRITICAL
With/with	out sciatica -	Function	ı (RMDQ) >4 mon	ths (Better indic	ated by lower v	alues)						
	randomised trials		no serious inconsistency	no serious indirectness	very serious ^b	none	51	52	-	MD 0.46 lower (2.28 lower to 1.36 higher)	⊕OOO VERY LOW	CRITICAL
With/with	out sciatica -	Psycholo	ogical distress (B	DI 0-63) <4 mon	ths (Better indi	cated by lower val	ues)					
	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	52	55	-	MD 0.55 higher (1.46 lower to 2.56	⊕OOO VERY LOW	CRITICAL

										higher)		
With/with	out sciatica -	Psychol	ogical distress (B	SDI 0-63) >4 mon	ths (Better indi	cated by lower val	ues)					
1	randomised trials		no serious inconsistency	no serious indirectness	very serious ^b	none	51	52	-	MD 1.15 higher (0.9 lower to 3.2 higher)		CRITICAL
With/with	out sciatica -	HC use (general practice	- visits) >4 mont	ths (Better indic	ated by lower valu	ues)					
1	randomised trials	serious ª	no serious inconsistency	no serious indirectness	very serious ^b	none	52	52	-	MD 0.30 lower (2.27 lower to 1.67 higher)	⊕000 VERY LOW	IMPORTANT
With/with	out sciatica -	HC use (specialist care - \	visits) >4 month	s (Better indica	ted by lower value	es)					
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	52	52	-	MD 0.58 higher (0.35 lower to 1.51 higher)	⊕⊕OO LOW	IMPORTANT
With/with	out sciatica -	HC use (radiography - vis	its) >4 months (Better indicated	d by lower values)						
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	52	52	-	MD 0.10 lower (0.24 lower to 0.04 higher)	⊕⊕⊕O MODERATE	IMPORTANT
With/with	out sciatica -	HC use (occupational phy	/sician - visits) >	•4 months (Bett	er indicated by lov	wer values)					
1	randomised trials	serious ª	no serious inconsistency	no serious indirectness	no serious imprecision	none	52	52	-	MD 0.14 lower (0.42 lower to 0.14 higher)	⊕⊕⊕O MODERATE	IMPORTANT
With/with	out sciatica -	HC use (psychologist - vi	sits) >4 months	(Better indicate	d by lower values)					
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	52	52	-	MD 0.28 higher (0.64 lower to 1.2 higher)	⊕000 VERY LOW	IMPORTANT
With/with	out sciatica -	HC use (therapist -sessio	ns) >4 months (Better indicated	l by lower values)						
1	randomised trials		no serious inconsistency	no serious indirectness	very serious ^b	none	52	52	-	MD 4.62 lower (10.23 lower to 0.99 higher)		IMPORTANT

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

NICE **J.5.2**2016 Combinations – exercise therapy adjunct

Low back pain without sciatica population

Table 104: Exercise (biomechanical) plus Electrotherapy (TENS) compared with Electrotherapy (TENS)

			Quality as:	sessment			No of patients	•		Effect	Quality	Importance	
No of studies	ies Design bias Inconsistency		Indirectness	Imprecision	Other considerations	Exercise (biomech) + TENS	TENS	Relative (95% CI)		Quanty	Importance		
Pain (Borg	n (Borg verbal pain rating scale 0-10) - <4 months (follow-up 8 weeks; measured with: Borg; range of scores: 0-10; Better indicated by lower values)												
	randomised trials	, ,			no serious imprecision	none	21	23	-	MD 0.16 lower (0.21 to 0.11 lower)	⊕⊕OO LOW	CRITICAL	
Function (Oswestry inde	x 0-100) - <	4 months (follow-u	p 8 weeks; measu	red with: ODI; ra	nge of scores: 0-50	; Better indicated by	y lowe	er values				
1	randomised trials	- ,			no serious imprecision	none	21	23	-	MD 3.2 lower (4.4 to 2 lower)	⊕⊕OO LOW	CRITICAL	

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 105: Exercise (biomechanical + aerobic) + electrotherapy (PENS) compared to sham electrotherapy (PENS)

			Quality as:	sessment			No of patients	S		Effect	Ovelity			
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Exercise (biomech + aerobic) + PENS	sham PENS	Relative (95% CI)	Absolute	Quality	Importance		
SF-36 (0-1	SF-36 (0-100) - <4 months: Mental component summary score (follow-up 6 weeks; measured with: SF-36; range of scores: 0-100; Better indicated by higher values)													
1	randomised trials	- ,			no serious imprecision	none	45	48	-	MD 0.2 lower (4.72 lower to 4.32 higher)	⊕⊕OO LOW	CRITICAL		

SF-36 (0-1	l00) - >4 mont	ths: Menta	I component sum	mary score (folio	w-up 6 months;	measured with: SF	F-36; range of scores:	0-100; B	etter ind	icated by higher value	es)	
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	45	48	-	MD 1.4 lower (6.52 lower to 3.72 higher)	⊕⊕OO LOW	CRITICAL
SF-36 (0-1	l00) - <4 mont	hs: Physic	cal component su	mmary score (fo	llow-up 6 weeks	measured with: S	F-36; range of scores	: 0-100; E	Better inc	dicated by higher valu	es)	
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	45	48	-	MD 2 lower (12.11 lower to 8.11 higher)	⊕⊕OO LOW	CRITICAL
SF-36 (0-1	l00) - >4 mont	hs: Physic	cal component su	mmary score (fo	llow-up 6 months	s; measured with:	SF-36; range of score	s: 0-100;	Better in	ndicated by higher val	ues)	
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	45	48	-	MD 0.7 lower (10.87 lower to 9.47 higher)	⊕⊕OO LOW	CRITICAL
Pain (McC	Gill) - <4 mont	hs (follow	-up 6 weeks; mea	sured with: McGi	II; range of score	es: 0-78; Better ind	icated by lower value	s)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	45	48	-	MD 1.8 lower (4.79 lower to 1.19 higher)	⊕000 VERY LOW	CRITICAL
Pain (McC	Gill) - >4 mont	hs (follow	-up 6 months; me	asured with: McC	Gill; range of sco	res: 0-78; Better in	dicated by lower valu	es)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	45	48	-	MD 0.5 lower (3.84 lower to 2.84 higher)	⊕⊕OO LOW	CRITICAL
Function	(Roland Morr	ris) - <4 mo	onths (follow-up 6	weeks; measure	ed with: RMDQ; r	ange of scores: 0-	24; Better indicated by	/ lower v	alues)			
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	45	48	-	MD 0.1 higher (1.62 lower to 1.82 higher)	⊕⊕OO LOW	CRITICAL
Function	(Roland Morri	is) - >4 mo	nths (follow-up 6	months; measur	ed with: RMDQ;	range of scores: 0	-24; Better indicated b	y lower v	/alues)			
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	45	48	-	MD 0.9 higher (0.93 lower to 2.73 higher)	⊕000 VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 106: Exercise (biomechanical + aerobic) + electrotherapy (PENS) compared to electrotherapy (PENS)

			Quality as	sessment		·	No of patients	1		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Exercise (biomech + aerobic) + PENS	PENS	Relative (95% CI)	Absolute		·
SF-36 (0-1	100) - <4 mont	hs: Menta	l component summ	nary score (follow	v-up 6 weeks; me	easured with: SF-3	6; range of scores: 0-	100; B	etter indi	cated by higher values	s)	
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	45	47	-	MD 1.8 lower (6.58 lower to 2.98 higher)	⊕OOO VERY LOW	CRITICAL
SF-36 (0-1	100) - >4 mont	hs: Menta	l component summ	nary score (follow	v-up 6 months; n	neasured with: SF	-36; range of scores: 0	-100; I	Better inc	dicated by higher value	es)	
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	45	47	-	MD 1.6 higher (4.37 lower to 7.57 higher)	⊕OOO VERY LOW	CRITICAL
SF-36 (0-1	l00) - <4 mont	hs: Physic	cal component sur	nmary score (foll	ow-up 6 weeks; ı	measured with: SF	-36; range of scores:	0-100;	Better in	dicated by higher valu	ies)	
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	45	47	-	MD 5 higher (4.58 lower to 14.58 higher)	⊕000 VERY LOW	CRITICAL
SF-36 (0-1	l00) - >4 mont	hs: Physic	cal component sur	nmary score (foll	ow-up 6 months;	measured with: S	F-36; range of scores	0-100	; Better i	ndicated by higher val	lues)	
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	45	47	-	MD 10.3 higher (0.78 to 19.82 higher)	⊕000 VERY LOW	CRITICAL
Pain (McC	Gill) - <4 montl	hs (follow-	up 6 weeks; meas	ured with: McGill	; range of scores	s: 0-78; Better indi	cated by lower values)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	45	47	-	MD 1.2 lower (4.76 lower to 2.36 higher)	⊕000 VERY LOW	CRITICAL
Pain (McC	Gill) - >4 mont	hs (follow-	up 6 months; mea	sured with: McGi	II; range of score	es: 0-78; Better ind	licated by lower value	s)				
1	randomised	very	no serious	no serious	no serious	none	45	47	-	MD 0.4 lower (3.75	⊕⊕OO	CRITICAL

	trials	serious ^a	inconsistency	indirectness	imprecision ^b					lower to 2.95 higher)	LOW			
Function	(Roland Morr	ris) - <4 mo	onths (follow-up 6	weeks; measured	I with: RMDQ; ra	nge of scores: 0-24	4; Better indicated by I	ower	values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	45	47	-	MD 0 higher (1.86 lower to 1.86 higher)	⊕⊕OO LOW	CRITICAL		
Function	Function (Roland Morris) - >4 months (follow-up 6 months; measured with: RMDQ; range of scores: 0-24; Better indicated by lower values)													
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	45	47	-	MD 0 higher (1.74 lower to 1.74 higher)	⊕⊕OO LOW	CRITICAL		

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 107: Group exercise (mixed: biomechanical + aerobic) + self management (education) + manual therapy (manipulation) compared to individual exercise (biomechanical) + self management (education) + manual therapy (manipulation)

			Quality ass	essment			No of p	atients		Effect				
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Group exercise (biomech + aerob) + education + manipulation	individual exercise (biomech) + education + manipulation	Relative (95% CI)	Absolute	Quality	Importance		
Analges	Analgesic use - <4 months (follow-up mean 8 weeks)													
1	randomised trials	,		no serious indirectness	Serious ^b	none	13/33 (39.4%)	20.7%	RR 1.9 (0.83 to 4.36)	186 more per 1000 (from 35 fewer to 696 more)	⊕000 VERY LOW	IMPORTANT		

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 108: Exercise (aerobic) + psychological intervention (behavioural therapy) compared to psychological intervention (behavioural therapy)

Quality assessment No of patients Effect Quality Ir	Quality assessment	No of patients	Effect	Quality Ir	mportance
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No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Exercise (aerobic) + behavioural therapy	behavioural therapy	Relative (95% CI)	Absolute			
Pain (Mc	ain (McGill) - <4 months (follow-up 8 weeks; measured with: McGill; range of scores: 0-78; Better indicated by lower values)												
1		- ,		no serious indirectness	Serious ^b	none	18	18	-	MD 2.93 lower (10.62 lower to 4.76 higher)	⊕OOO VERY LOW	CRITICAL	

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 109: Exercise (aerobic) + psychological intervention (cognitive behavioural approaches) + self management (education) compared to psychological intervention (cognitive behavioural approaches) + self management (education)

	psychio	iogicai i	incorrection (c	oginare ser	aviourara	pproudines, : s	en management (eut	ica ciori,				
			Quality ass	essment			No of pati	ents		Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Exercise (aerobic) + cognitive behavioural approaches + education	cognitive behavioural approaches + education	Relative (95% CI)		Quality	Importance
Pain (0-1	00 NRS conv	erted to (0-10) - <4 months	(follow-up 3 m	onths; meas	ured with: NRS;	range of scores: 0-10; Bet	ter indicated by lowe	er values)			
	randomised trials	,	no serious inconsistency		very serious ^b	none	15	12	1	MD 0.35 lower (2.34 lower to 1.64 higher)	⊕OOO VERY LOW	CRITICAL
Function	(Roland Mor	rris 0-24)	- <4 months (follo	ow-up 3 months	s; measured	with: RMDQ; ran	ge of scores: 0-24; Better	indicated by lower v	alues)			
	randomised trials	,	no serious inconsistency	no serious indirectness	Serious ^b	none	15	12	-	MD 2.1 higher (1.41 lower to 5.61 higher)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 110: Exercise (biomechanical – pilates) + self management (education) compared to self-management (education)

			Quality asse	ssment			No of p	patients		Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Pilates + education +	self- management	Relative (95% CI)	Absolute	Quality	Importance
Pain (NRS	6 0-10) - <4 mo	nths (follo	w-up 6 weeks; me	asured with: NRS	; range of so	cores: 0-10; Better	indicated by lov	wer values)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	43	43	-	MD 2.1 lower (3.07 to 1.13 lower)	⊕OOO VERY LOW	CRITICAL
Pain (NRS 0-10) - >4 months (follow-up 6 months; measured with: NRS; range of scores: 0-10; Better indicated by lower values)												
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	43	43	-	MD 0.8 lower (1.75 lower to 0.15 higher)	⊕000 VERY LOW	CRITICAL
Function ((Roland Morri	s 0-24) - <4	l months (follow-u	o 6 weeks; measu	red with: RN	IDQ; range of scor	res: 0-24; Better	indicated by lo	wer valu	es)		
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	43	43	-	MD 3.5 lower (5.48 to 1.52 lower)	⊕000 VERY LOW	CRITICAL
Function ((Roland Morri	s 0-24) - >4	l months (follow-u	o 6 months; meas	sured with: R	RMDQ; range of sco	ores: 0-24; Bette	er indicated by	lower val	ues)		
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	43	43	-	MD 2.2 lower (4.35 to 0.05 lower)	⊕000 VERY	CRITICAL

						LOW	ĺ
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^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Low back pain with sciatica population J.5.2.2

Table 111: Exercise (biomechanical) + self-management (unsupervised exercise) compared to TENS + laser + massage + self-management (unsupervised exercise)

		-	Quality as	sessment			No of patients			Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Exercise (biomech) + self- management (unsupervised exercise)		Relative (95% CI)	Absolute	Quality	Importance
Overall -	rall - Pain (VAS 0-10) <4 months (Better indicated by lower values)											
	randomised trials			no serious indirectness	no serious imprecision	none	20	20	-	MD 3.19 lower (3.95 to 2.43 lower)	⊕⊕⊕O MODERATE	CRITICAL
Overall -	Function (rev	ised ODI	0-100) < 4 month	s (Better indica	ted by lower va	lues)						
	randomised trials			no serious indirectness	no serious imprecision	none	20	20	-	MD 18.21 lower (23.07 to 13.35 lower)	⊕⊕⊕O MODERATE	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

NICE. 2016 Low back pain with/without sciatica population

Table 112: Exercise plus orthoses compared to orthoses

			Quality asse	ssment			No of pat	ients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Exercise + orthoses	orthoses	Relative (95% CI) Absolute		,	
Responder criteria (remission of pain) - >4 months												
		,			very serious ^b	none	6/24 (25%)	25%	RR 1 (0.38 to 2.66)	0 fewer per 1000 (from 155 fewer to 415 more)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 113: Exercise plus self-management (education) compared to self-management

			Quality as	sessment			No of p	oatients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Exercise + education	self- management	Relative (95% CI)	Absolute		
Number i	lumber improving on Disability index - >4 months											
1		very serious ^a			no serious imprecision	none	17/46 (37%)	6.8%	RR 5.42 (1.71 to 17.22)	301 more per 1000 (from 48 more to 1000 more)	⊕⊕OO LOW	CRITICAL
Number i	mproving on	Quality of	f life index - >4 mo	onths								
1		very seriousª			no serious imprecision	none	45/46 (97.8%)	27.3%	RR 3.59 (2.21 to 5.82)	707 more per 1000 (from 330 more to 1000 more)	⊕⊕OO LOW	CRITICAL

a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 114: Exercise plus self-management (mixed modality – home exercise plus education) compared to usual care

Tubic II	-TI EXCITING	pius sen	management	(IIIIXCU IIICUUII	ity Home CA	croise plas caa	cation) compared to us	 	1			
			Quality as	sessment			No of patients			Effect	Ouality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Exercise + home exercise + relaxation + education	usual care	Relative (95% CI)	Absolute	Quanty	importance
Function ((Roland Morri	s 0-24) - <	4 months (Better i	ndicated by lowe	r values)							
1		very serious ^a	no serious inconsistency		no serious imprecision	none	100	109	-	MD 0.8 lower (1.33 to 0.27 lower)	⊕⊕OO LOW	CRITICAL
Function ((Roland Morri	s 0-24) - >	4 months (Better i	ndicated by lowe	r values)							
1		very serious ^a	no serious inconsistency		no serious imprecision	none	100	109	-	MD 2.3 lower (2.87 to 1.73 lower)	⊕⊕OO LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 115: Exercise plus self management (mixed modality – home exercise + education) compared to self-management (education)

			Quality as:	sessment			No of patients			Effect	Ouglitu	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Exercise + home exercise + relaxation + education	education	Relative		Quality	Importance
Function (Roland Morris 0-24) - <4 months (Better indicated by lower values)												
		,			no serious imprecision	none	100	139	-	MD 0 higher (0.48 lower to 0.48 higher)		CRITICAL

Function	unction (Roland Morris 0-24) - >4 months (Better indicated by lower values)														
		- ,			no serious imprecision	none	100	139		MD 0.4 lower (1.05 lower to 0.25 higher)		CRITICAL			

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 116: Exercise (biomechanical) + self-management (home exercise) compared to self-management (self-care advice based on the Back Book)

			Quality as	sessment			No of pat	ients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Exercise (biomech) + home exercise	self- management	Relative (95% CI)	Absolute	Quality	Importance
Quality of	life (15D 0 to	1) - <4 m	onths (Better ind	icated by lower	values)							
	randomised trials			no serious indirectness	Serious ^b	none	43	40	-	MD 0.01 higher (0.02 lower to 0.04 higher)	⊕⊕OO LOW	CRITICAL
Quality of	life (15D 0 to	1) - >4 m	onths (Better ind	icated by lower	values)							
	randomised trials	Serious ^a		no serious indirectness	Serious ^b	none	43	40	-	MD 0.02 higher (0.01 lower to 0.05 higher)	⊕⊕OO LOW	CRITICAL
Pain (0-10	00 VAS conve	erted to 0-	10) - <4 months (I	Better indicated	by lower value	s)						
	randomised trials	Seriousª		no serious indirectness	Serious ^b	none	43	40	-	MD 0.4 lower (1.45 lower to 0.65 higher)	⊕⊕OO LOW	CRITICAL
Pain (0-10	00 VAS conve	erted to 0-	10) - >4 months (I	Better indicated	by lower value	s)						
	randomised trials	Seriousª		no serious indirectness	Serious ^b	none	43	40	-	MD 1 lower (2.02 lower to 0.02 higher)	⊕⊕OO LOW	CRITICAL
Function	(Roland Morr	ris 18 item	ı) - <4 months (Be	etter indicated by	y lower values)							

1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	43	40	-	MD 0 higher (1.94 lower to 1.94 higher)	⊕⊕⊕O MODERATE	CRITICAL
Function	(Roland Mor	ris 18 iten	n) - >4 months (B	etter indicated b	y lower values)							
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	43	40	-	MD 1 lower (3.15 lower to 1.15 higher)	⊕⊕OO LOW	CRITICAL
							2 increments if the n			vas at very high risk o	of bias	

Table 117: Exercise (biomechanical – core stability) + manual therapy (massage) compared to manual therapy (massage)

			Quality as:	sessment			No of patients			Effect			
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Exercise (biomechanical - core stability) + manual therapy (massage) vs manual therapy (massage)	Control	Relative (95% CI)	Absolute	Quality	Importance	
Pain sev	erity (VAS, 0	-10) < 4 m	nonths (Better in	dicated by lowe	er values)								
		- ,			no serious imprecision	none	46	46	-	MD 1.39 lower (1.9 to 0.88 lower)	⊕⊕OO LOW	CRITICAL	
Function	unction (ODI, 0-100) < 4 months (Better indicated by lower values)												
		,		no serious indirectness	Serious ^b	none	46	46	1	MD 5.19 lower (6.46 to 3.92 lower)	⊕OOO VERY LOW	CRITICAL	

Respon	der criteria (p	ain free i	nterval > 30 days	;)						
1		,		no serious indirectness	Serious ^b	none	43/43 (100%)	100%	0 fewer per 1000 (from 40 fewer to 50 more)	IMPORTANT

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 118: Exercise (core stability) + manual therapy (manipulation) compared to self-management (advice to stay active) + manual therapy (manipulation)

Risk of bias life (SF-12 (very serious ^a	Inconsistency 0-100) <4 months	Indirectness	Imprecision	Other considerations	Exercise (core stability) + manipulation	Self management (advice to stay active)	Relative (95%	Absolute	Quality	Importance
d very		s - Physical (Bet	tter indicated b			+ manipulation	CI)			
,	no serious			y lower values)						
Scrious			no serious imprecision	none	12	13	-	MD 9.3 higher (3.12 to 15.48 higher)	⊕⊕OO LOW	CRITICAL
life (SF-12 (0-100) <4 months	s - Mental (Bette	er indicated by	lower values)						
d very serious ^a		no serious indirectness	very serious ^b	none	12	13	-	MD 2.6 higher (5.51 lower to 10.71 higher)	⊕OOO VERY LOW	CRITICAL
life (SF-12 (0-100) 4 months -	- 1 year - Physic	cal (Better indi	cated by lower val	lues)					
d very serious ^a		no serious indirectness	serious ^b	none	12	13	-	MD 3.4 higher (1.94 lower to 8.74 higher)	⊕OOO VERY LOW	CRITICAL
:(very serious ^a life (SF-12 of very serious ^a	d very no serious inconsistency life (SF-12 0-100) 4 months of very serious inconsistency	d very serious inconsistency indirectness life (SF-12 0-100) 4 months - 1 year - Physic very serious inconsistency indirectness no serious indirectness	d very serious inconsistency no serious very serious no serious serious inconsistency no serious serious very very very very very very very very	serious ^a inconsistency indirectness life (SF-12 0-100) 4 months - 1 year - Physical (Better indicated by lower value) d very serious ^a inconsistency indirectness serious ^b none	d very no serious no serious indirectness very serious none 12 life (SF-12 0-100) 4 months - 1 year - Physical (Better indicated by lower values) d very no serious no serious serious none 12	d very serious inconsistency indirectness very serious none 12 13 life (SF-12 0-100) 4 months - 1 year - Physical (Better indicated by lower values) d very serious inconsistency inconsistency indirectness serious none 12 13	d very serious inconsistency indirectness very serious none 12 13 - life (SF-12 0-100) 4 months - 1 year - Physical (Better indicated by lower values) d very serious inconsistency indirectness serious indirectness indirectness serious indirectness in	d very serious inconsistency indirectness very serious none 12 13 - MD 2.6 higher (5.51 lower to 10.71 higher) life (SF-12 0-100) 4 months - 1 year - Physical (Better indicated by lower values) d very serious inconsistency inconsistency indirectness serious indirectness indir	d very serious inconsistency indirectness very serious none 12 13 - MD 2.6 higher (5.51 lower to 10.71 higher) VERY LOW life (SF-12 0-100) 4 months - 1 year - Physical (Better indicated by lower values) d very serious inconsistency indirectness serious indirectness serious indirectness indirectness serious indirectness indirec

1		very seriousª	no serious inconsistency	no serious indirectness	serious ^b	none	12	13	-	MD 8.3 higher (0.59 to 16.01 higher)	⊕000 VERY LOW	CRITICAL
Overall -	Pain (McGill	- sensor	y, 0-33) <4 month	s (Better indica	ated by lower v	alues)						
1		very seriousª	no serious inconsistency	no serious indirectness	serious ^b	none	12	13	-	MD 3.5 lower (6.9 to 0.1 lower)	⊕OOO VERY LOW	CRITICAL
Overall -	Pain (McGill	- sensor	y, 0-33) 4 months	- 1 year (Bette	r indicated by I	ower values)						
1		very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	12	13	-	MD 2.3 lower (5.48 lower to 0.88 higher)	⊕000 VERY LOW	CRITICAL
Overall -	Pain (McGill	- affectiv	e, 0-12) <4 montl	ns (Better indic	ated by lower v	/alues)						
1		very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	12	13	-	MD 1.9 lower (4.97 lower to 1.17 higher)	⊕OOO VERY LOW	CRITICAL
Overall -	Pain (McGill	- affectiv	e, 0-12) 4 months	s - 1 year (Bette	r indicated by	lower values)						
1		very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	12	13	-	MD 0.6 lower (1.74 lower to 0.54 higher)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 119: Mixed exercise (biomechanical + aerobic) + Alexander technique compared to Alexander technique

	Quality assessment					No of patie		Effect				
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Mixed exercise + Alexander technique	Alexander	Relative (95% CI)	Absolute	Quality	Importance
Overall - F	Function (RM	DQ 0-24) <	4 months (Better	indicated by low	er values)						•	

	randomised trials				very serious ^b	none	15	15		MD 1.28 higher (2.8 lower to 5.36 higher)		CRITICAL
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^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 120: Exercise (individual biomechanical) + self management compared to self management

very serious ¹	nths (Better indicate inconsistency	no serious indirectness	no serious imprecision	Other considerations none	Individual biomechanical exercise + self management	Self management 256	Relative (95% CI)	Absolute MD 1.36 lower (2.15 to 0.57 lower)	⊕⊕OO LOW	CRITICAL
very serious ¹ 1) - > 4 more very	no serious inconsistency	no serious indirectness	no serious imprecision	none	225	256	-	(2.15 to 0.57		CRITICAL
serious ¹ 1) - > 4 mor	inconsistency nths (Better indic	indirectness	imprecision	none	225	256	-	(2.15 to 0.57		CRITICAL
very	no serious		/alues)							
		no sorious						1		
	inconsistency	indirectness	no serious imprecision	none	216	248	-	MD 0.39 lower (1.24 lower to 0.46 higher)	⊕⊕OO LOW	CRITICAL
- < 4 montl	hs (Better indicat	ted by lower val	lues)							
very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	204	239	-	MD 0.46 lower (0.85 to 0.07 lower)	⊕⊕OO LOW	CRITICAL
- > 4 montl	hs (Better indicat	ted by lower val	lues)							
very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	200	235	-	MD 0.69 lower (1.18 to 0.2 lower)	⊕⊕OO LOW	CRITICAL
	very serious¹ -> 4 mont very serious¹	very no serious inconsistency -> 4 months (Better indicativery no serious inconsistency)	very no serious inconsistency no serious indirectness -> 4 months (Better indicated by lower value) very no serious no serious inconsistency inconsistency indirectness	serious¹ inconsistency indirectness imprecision -> 4 months (Better indicated by lower values) very	very no serious inconsistency no serious indirectness no serious imprecision -> 4 months (Better indicated by lower values) very no serious no serious inconsistency no serious indirectness imprecision	very serious¹ no serious indirectness no serious imprecision none 204 -> 4 months (Better indicated by lower values) very serious¹ no serious inconsistency no serious indirectness no serious imprecision none 200	very serious¹ no serious inconsistency no serious indirectness no serious imprecision none 204 239 -> 4 months (Better indicated by lower values) very no serious no serious no serious none 200 235	very serious¹ no serious inconsistency no serious indirectness none 204 239 - -> 4 months (Better indicated by lower values) very serious¹ no serious inconsistency no serious indirectness none 200 235 -	very serious¹ no serious inconsistency no serious indirectness no serious imprecision none 204 239 - MD 0.46 lower (0.85 to 0.07 lower) -> 4 months (Better indicated by lower values) very serious¹ no serious inconsistency no serious indirectness no serious imprecision none 200 235 - MD 0.69 lower (1.18 to 0.2 lower)	very serious¹ no serious inconsistency no serious indirectness none 204 239 - MD 0.46 lower (0.85 to 0.07 LOW lower) ⊕⊕OO LOW -> 4 months (Better indicated by lower values) very serious¹ no serious inconsistency no serious indirectness none 200 235 - MD 0.69 lower (1.18 to 0.2 lower) LOW

1		- ,	no serious inconsistency		no serious imprecision	none	191	227	-	MD 2.41 higher (1.13 to 3.69 higher)	⊕⊕OO LOW	CRITICAL
Quality o	f life (SF36 0-	100) - > 4	months: Physica	al component (E	Setter indicated	by lower values)						
1		- ,	no serious inconsistency		no serious imprecision	none	194	221	-	MD 1.55 higher (0.14 lower to 3.24 higher)	⊕⊕OO LOW	CRITICAL
Quality o	f life (SF36 0-	100) - < 4	months: Mental	component (Be	tter indicated b	y lower values)						
1		- ,			no serious imprecision	none	191	227	-	MD 0.75 higher (1.04 lower to 2.54 higher)	⊕⊕OO LOW	CRITICAL
Quality o	f life (SF36 0-	100) - > 4	months: Mental	component (Be	tter indicated b	y lower values)						
1		- ,	no serious inconsistency		no serious imprecision	none	194	221	-	mean 0 higher (1.86 lower to 2.52 higher)	⊕⊕OO LOW	CRITICAL
Function	(Von Korff di	sability, ()-100) - < 4 montl	ns (Better indica	ted by lower va	alues)						
1		,	no serious inconsistency		no serious imprecision	none	205	239	-	MD 0.5 lower (0.94 to 0.06 lower)	⊕⊕OO LOW	CRITICAL
Function	(Von Korff di	sability, ()-100) - > 4 montl	ns (Better indica	ited by lower va	alues)						
1		- ,	no serious inconsistency		no serious imprecision	none	202	235	-	MD 0.46 lower (0.91 to 0.01 lower)	⊕⊕OO LOW	CRITICAL

J.6 Postural therapies

J.6.1 Single interventions

Table 121: Alexander technique (6 lessons) versus usual care for low back pain and sciatica at > 4 months - 1 year (without sciatica)

			<u> </u>	<u> </u>								
			Quality as	sessment			No of patients			Effect	Quality	Importanc
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alexander technique (6 lessons) versus usual care	Contro I	Relative (95% CI)	Absolute	Quality	e
SF-36 phy	ysical (1 year)	(range o	f scores: 0-100; B	Setter indicated I	oy higher values	s)						
	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	58	60	-	MD 2.04 higher (5.58 lower to 9.66 higher)	⊕⊕OO LOW	CRITICAL
SF-36 me	ntal (1 year) (range of	scores: 0-100; Be	tter indicated by	higher values)							
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	58	60	-	MD 4.1 higher (3.27 lower to 11.47 higher)	⊕⊕OO LOW	CRITICAL
Von Korff	pain scale (1	year) (ra	nge of scores: 0-	10; Better indica	ted by lower va	lues)						
	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	58	60	-	MD 0.44 lower (1.31 lower to 0.43 higher)	⊕⊕OO LOW	CRITICAL
Roland M	orris Disabili	ty scale (1 year) (range of s	scores: 0-28; Bet	tter indicated by	lower values)						
	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	58	60	1	MD 1.44 lower (3.34 lower to 0.46 higher)	⊕⊕OO LOW	CRITICAL
Primary c	are contacts	(Better in	dicated by lower	values)								
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	58	60	-	MD 0.05 higher (0.25 lower to 0.35 higher)	⊕⊕⊕O MODERAT E	IMPORTAN T

Prescript	ions (Better in	ndicated I	oy lower values)						
1	randomised trials			 no serious imprecision	none	58	60	MD 0.21 lower (0.72 lower to 0.3 higher)	IMPORTAN T

Low back pain and sciatica in over 16s Quality assessment

Table 122: Alexander technique (10 sessions) versus usual care (overall population)

			Quality asse	essment			No of patients Effect				Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alexander texhnique (10 lessons) versus usual care	Control	Relative (95% CI)	Absolute	Quanty	importance
Overall - I	unction (RMI	OQ 0-24) <	4 months [mean d	ifference from c	ontrol] (Bette	er indicated by low	ver values)					
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	15	13	-	MD 1.38 lower (4.82 lower to 2.07 higher)	⊕OOO VERY LOW	CRITICAL
Overall - I	Pain (von Korf	ff 0-100) <	4 months [mean d	ifference from co	ontrol] (Bette	r indicated by low	er values)					
1	randomised trials		no serious inconsistency	no serious indirectness	serious ^b	none	15	13	-	MD 0.63 lower (1.99 lower to 0.73 higher)	⊕⊕OO LOW	CRITICAL
Overall - I	Function (RMI	OQ 0-24) 4	months - 1 year [mean difference	from control] (Better indicated	by lower values)					
1	randomised trials		no serious inconsistency	no serious indirectness	serious ^b	none	15	13	-	MD 2.86 lower (6.53 lower to 0.81 higher)	⊕⊕OO LOW	CRITICAL
Overall - I	Pain (von Korf	f 0-100) 4	months - 1 year [r	nean difference	from control]	(Better indicated	by lower values)					
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	15	13	-	MD 0.09 higher (1.35 lower to 1.52 higher)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 123: Alexander technique (24 lessons) versus usual care for low back pain and sciatica at > 4 months - 1 year (without sciatica)

			Quality as	sessment			No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alexander technique (24 lessons) versus usual care	Control	Relative (95% CI)	Absolute	Quanty	Importance
SF-36 phy	/sical (1 year)	(range o	f scores: 0-100; E	Better indicated I	by higher value	s)						
1	randomised trials		no serious inconsistency	no serious indirectness	serious ^b	none	61	60	-	MD 11.83 higher (4.42 to 19.24 higher)	⊕⊕OO LOW	CRITICAL
SF-36 me	ntal (1 year) (range of	scores: 0-100; Be	tter indicated by	higher values)							
1 -	randomised trials		no serious inconsistency	no serious indirectness	serious ^b	none	61	60	-	MD 3.74 higher (3.56 lower to 11.04 higher)	⊕⊕OO LOW	CRITICAL
Von Korff	pain scale (1	year) (ra	nge of scores: 0-	10; Better indica	ted by lower va	alues)		•				
1	randomised trials		no serious inconsistency	no serious indirectness	serious ^b	none	61	60	ı	MD 1.34 lower (2.2 to 0.48 lower)	⊕⊕OO LOW	CRITICAL
Roland M	orris Disabili	ty scale (1	1 year) (range of s	scores: 0-28; Be	tter indicated b	y lower values)						
	randomised trials		no serious inconsistency	no serious indirectness	serious ^b	none	61	60	-	MD 4.14 lower (6.01 to 2.27 lower)	⊕⊕OO LOW	CRITICAL
Primary c	are contacts	(Better in	dicated by lower	values)								
	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	61	60	1	MD 0.01 higher (0.28 lower to 0.3 higher)	⊕⊕⊕O MODERATE	IMPORTANT
Prescript	ions (Better in	ndicated l	oy lower values)									

1	randomised trials			no serious indirectness	serious ^b	none	61	60	-	MD 0.22 higher (0.48 lower to 0.92 higher)	⊕⊕OO LOW	IMPORTANT
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^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 124: Alexander technique (6 lessons) versus self-management (exercise prescription)at > 4 months - 1 year (without sciatica)

			<u> </u>			<u> </u>	· · · · · · · · · · · · · · · · · · ·					
			Quality as	sessment			No of patients			Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alexander technique (6 lessons) versus exercise prescription		Relative (95% CI)	Absolute	Quality	Importance
SF-36 ph	ysical (1 year)) (range o	of scores: 0-100;	Better indicated	by higher valu	ies)						
	randomised trials		no serious inconsistency	no serious indirectness	serious ^b	none	58	51	-	MD 4.12 higher (5.17 lower to 13.41 higher)	⊕⊕OO LOW	CRITICAL
SF-36 me	ental (1 year) (range of	scores: 0-100; B	etter indicated b	y higher value	s)						
	randomised trials		no serious inconsistency		no serious imprecision	none	58	51	-	MD 3.38 higher (5.2 lower to 11.96 higher)	⊕⊕⊕O MODERATE	CRITICAL
Von Korf	f pain scale (1	l year) (ra	ange of scores: 0	-10; Better indic	ated by lower	values)						
	randomised trials		no serious inconsistency		no serious imprecision	none	58	51	-	MD 0.13 lower (1.15 lower to 0.89 higher)	⊕⊕⊕O MODERATE	CRITICAL
Roland M	lorris Disabili	ty scale (1 year) (range of	scores: 0-28; B	etter indicated	by lower values)						
	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	58	51	-	MD 0.21 higher (1.76 lower to 2.18 higher)	⊕⊕⊕O MODERATE	CRITICAL
Primary o	are contacts	(Better in	ndicated by lower	r values)								

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		atica in
		over
		16s

1	randomised trials			 no serious imprecision	none	58	51	-	MD 0.02 lower (0.38 lower to 0.34 higher)	0000	IMPORTANT
Prescript	tions (Better i	ndicated	by lower values)								
1	randomised trials			 no serious imprecision	none	58	51	-	MD 0.24 lower (0.76 lower to 0.28 higher)		IMPORTANT

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 125: Alexander technique (24 lessons) versus self-management (exercise prescription)at > 4 months - 1 year (without sciatica)

			Quality as	sessment			No of patients Effect				Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alexander technique (24 lessons) versus exercise prescription		Relative (95% CI)	Absolute	Quality	Importance
SF-36 ph	ysical (1 year) (range o	of scores: 0-100;	Better indicated	by higher valu	es)						
1	randomised trials		no serious inconsistency	no serious indirectness	serious ^b	none	61	51	-	MD 13.91 higher (4.79 to 23.03 higher)	⊕⊕OO LOW	CRITICAL
SF-36 me	ental (1 year)	(range of	scores: 0-100; B	etter indicated b	y higher values	s)						
1	randomised trials				no serious imprecision	none	61	51	-	MD 3.02 higher (5.91 lower to 11.95 higher)	⊕⊕⊕O MODERATE	CRITICAL
Von Korf	f pain scale (1 year) (ra	ange of scores: 0	-10; Better indic	ated by lower v	values)						
1	randomised trials		no serious inconsistency	no serious indirectness	serious ^b	none	61	51	-	MD 1.03 lower (2.04 to 0.02 lower)	⊕⊕OO LOW	CRITICAL
Roland M	lorris Disabili	ty scale (1 year) (range of	scores: 0-28; B	etter indicated	by lower values)						

1	randomised trials			no serious indirectness	serious ^b	none	61	51	-	MD 2.49 lower (4.43 to 0.55 lower)	⊕⊕OO LOW	CRITICAL
Primary	care contacts	(Better in	ndicated by lower	r values)								
1	randomised trials				no serious imprecision	none	61	51	-	MD 0.06 lower (0.41 lower to 0.29 higher)		IMPORTANT
Prescrip	tions (Better i	ndicated	by lower values)									
1	randomised trials			no serious indirectness	serious ^b	none	61	51	-	MD 0.19 higher (0.52 lower to 0.9 higher)	⊕⊕OO LOW	IMPORTANT

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 126: Alexander technique (24 lessons) versus Alexander technique (6 lessons) at > 4 months - 1 year (without sciatica)

			Quality as	sessment			No of patients			Effect	Quality			
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alexander technique (24 lessons) versus Alexander technique (6 lessons)		Relative (95% CI)	Absolute	Quality	Importance		
SF-36 ph	F-36 physical (1 year) (range of scores: 0-100; Better indicated by higher values)													
	randomised trials			no serious indirectness	serious ^b	none	61	58	-	MD 9.79 higher (18.08 to 1.5 higher)	⊕⊕OO LOW	CRITICAL		
SF-36 me	SF-36 mental (1 year) (range of scores: 0-100; Better indicated by higher values)													
	randomised trials				no serious imprecision	none	61	58	-	MD 0.36 lower (7.47 higher to 8.19 lower)	⊕⊕⊕O MODERATE	CRITICAL		
Von Korf	Von Korff pain scale (1 year) (range of scores: 0-10; Better indicated by lower values)													

1	randomised trials		no serious inconsistency	no serious indirectness	serious ^b	none	61	58	-	MD 0.9 lower (0.03 higher to 1.83 lower)	⊕⊕OO LOW	CRITICAL			
Roland N	Roland Morris Disability scale (1 year) (range of scores: 0-28; Better indicated by lower values)														
1	randomised trials		no serious inconsistency	no serious indirectness	serious ^b	none	61	58	-	MD 2.7 lower (0.83 to 4.57 lower)	⊕⊕OO LOW	CRITICAL			
Primary (Primary care contacts (Better indicated by lower values)														
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	61	58	-	MD 0.04 lower (0.29 higher to 0.37 lower)	⊕⊕⊕O MODERATE	IMPORTANT			
Prescrip	Prescriptions (Better indicated by lower values)														
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	61	58	-	MD 0.43 higher (1.07 higher to 0.21 lower)	⊕⊕⊕O MODERATE	IMPORTANT			

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 127: Alexander technique (6 lessons) versus massage at > 4 months - 1 year (without sciatica)

	. r. Alexand	ici tetili	inque (o lesson	3) VCI 3U3 IIIU3	suge at > + II	ionitiis I year	(Without Sciatica)					•		
			Quality as:	sessment			No of patients			Effect	.			
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alexander technique (6 lessons) versus massage	Control	Relative II (95% Absolute CI)	Quality	Importance			
SF-36 phy	F-36 physical (1 year) (range of scores: 0-100; Better indicated by higher values)													
	randomised trials			no serious indirectness	serious ^b	none	58	64	-	MD 3.49 higher (4.96 lower to 11.94 higher)	⊕⊕OO LOW	CRITICAL		
SF-36 me	SF-36 mental (1 year) (range of scores: 0-100; Better indicated by higher values)													

1	randomised trials	serious ^a	no serious inconsistency		no serious imprecision	none	58	64	-	MD 6.21 higher (1.58 lower to 14 higher)	⊕⊕⊕O MODERATE	CRITICAL			
Von Korf	Von Korff pain scale (1 year) (range of scores: 0-10; Better indicated by lower values)														
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	58	64	-	MD 0.73 lower (1.67 lower to 0.21 higher)	⊕⊕OO LOW	CRITICAL			
Roland M	coland Morris Disability scale (1 year) (range of scores: 0-28; Better indicated by lower values)														
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	58	64	-	MD 0.99 lower (2.84 lower to 0.86 higher)		CRITICAL			
Primary o	care contacts	(Better in	dicated by lower	values)											
1	randomised trials	serious ^a	no serious inconsistency		no serious imprecision	none	58	64	-	MD 0.19 lower (0.6 lower to 0.22 higher)		IMPORTANT			
Prescript	ions (Better i	ndicated l	by lower values)												
1	randomised trials	serious ^a	no serious inconsistency		no serious imprecision	none	58	64	-	MD 0.13 lower (0.63 lower to 0.37 higher)		IMPORTANT			

Table 128: Alexander technique (24 lessons) versus massage at > 4 months - 1 year (without sciatica)

				,			(Without Sciatica)						
			Quality as	sessment			No of patients			Effect	Quality	Immortonos	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alexander technique (24 lessons) versus massage	Control	Relative (95% Absolute CI)		Quanty	Importance	
SF-36 phy	SF-36 physical (1 year) (range of scores: 0-100; Better indicated by higher values)												
1	randomised trials	serious ^a		no serious indirectness	serious ^b	none	61	64	-	MD 13.28 higher (5.02 to 21.54 higher)	⊕⊕OO LOW	CRITICAL	

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

SF-36 mental (1 year) (range of scores: 0-100; Better indicated by higher values)															
SF-36 me	ental (1 year) (range of	scores: 0-100; Be	tter indicated by	/ higher values)	1						1			
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	61	64	ı	MD 5.85 higher (2.32 lower to 14.02 higher)	⊕⊕OO LOW	CRITICAL			
Von Korff pain scale (1 year) (range of scores: 0-10; Better indicated by lower values)															
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	61	64	ı	MD 1.63 lower (2.56 to 0.7 lower)	⊕⊕OO LOW	CRITICAL			
Roland M	Roland Morris Disability scale (1 year) (range of scores: 0-28; Better indicated by lower values)														
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	61	64	-	MD 3.69 lower (5.51 to 1.87 lower)	⊕⊕OO LOW	CRITICAL			
Primary o	are contacts	(Better in	dicated by lower	values)											
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	61	64	ı	MD 0.23 lower (0.63 lower to 0.17 higher)		IMPORTANT			
Prescript	ions (Better i	ndicated	by lower values)												
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	61	64	-	MD 0.3 higher (0.39 lower to 0.99 higher)	⊕⊕OO LOW	IMPORTANT			

Table 129: Alexander technique (10 sessions) versus mixed exercise (overall population)

			Quality asse	essment			No of patients			Effect	Quality	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alexander technique (10 lessons) versus mixed exercise	Control	Relative (95% CI)	Absolute	Quality	Importance
Overall - Function (RMDQ 0-24) <4 months (Better indicated by lower values)												
1	randomised	Serious ^a	no serious	no serious	very	none	15	14	-	MD 0.12 higher (3.06	⊕000	CRITICAL

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

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trials	inconsistency	indirectness	serious ^b			lower to 3.3 higher)	VERY	
	•					,	LOW	

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Combined interventions (postural therapy adjunct) J.6.2

Table 130: Combined intervention Postural therapy + MBR versus MBR only (< 4 months)

			Quality as:	sessment			N	lo of patients		Effect				
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Combined intervention	MBR programme 3 elements: physical + psychological + education	Relative (95% CI)	Absolute	Quality	Importance		
Back pai	ack pain severity (NRS, 0-10) < 4 months (follow-up 2 years; Better indicated by lower values)													
1	randomised trials	serious ^a			no serious imprecision	none	77	77	-	MD 0.1 higher (0.3 lower to 0.5 higher)	⊕⊕⊕O MODERATE	CRITICAL		
Leg pain	severity (NR	S, 0-10) ·	< 4 months (follo	w-up 2 years; E	Better indicated	d by lower values)							
1	randomised trials	serious ^a			no serious imprecision	none	77	77	-	MD 0.2 higher (0.34 lower to 0.74 higher)	⊕⊕⊕O MODERATE	CRITICAL		
Function	(ODI, 0-100)	< 4 mont	ths (follow-up 2 y	ears; Better in	dicated by low	er values)								
1	randomised trials	serious ^a			no serious imprecision	none	77	77	-	MD 2.8 lower (4.63 to 0.97 lower)	⊕⊕⊕O MODERATE	CRITICAL		

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias

Table 131: Alexander technique (6 lessons) + self-management (exercise prescription) versus usual care (without sciatica)

				•		•				•				
			Quality ass	essment			No of patients			Effect				
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alexander techniques (6 lessons) + self management (exercise prescription)	Usual care	Relative (95% CI)	Absolute	Quality	Importance		
Function	unction (RMDQ 0-24) - Function (RMDQ 0-24) (Better indicated by lower values)													
1	randomised trials			no serious indirectness	serious ²	none	71	72	-	MD 2.98 lower (4.88 to 1.08 lower)	⊕⊕OO LOW	CRITICAL		
Pain (Vor	in (Von Korff scale 0-10) - Pain (Von Korff scale 0-10) (Better indicated by lower values)													
1	randomised trials			no serious indirectness	serious ²	none	71	72	-	MD 1.08 lower (1.96 to 0.2 lower)	⊕⊕OO LOW	CRITICAL		
Quality o	f life: SF-36 n	nental - Q	uality of life: SF-3	6 mental (Bette	r indicated b	y lower values)								
1	randomised trials	serious ¹		no serious indirectness	very serious²	none	71	72	-	MD 0.64 higher (6.79 lower to 8.07 higher)	⊕OOO VERY LOW	CRITICAL		
Quality o	f life: SF-36 p	hysical -	Quality of life: SF	-36 physical (Be	etter indicate	d by lower values								
1	randomised trials	serious¹		no serious indirectness	serious²	none	71	72	-	MD 8.53 higher (0.86 to 16.2 higher)	⊕⊕OO LOW	CRITICAL		

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 132: Alexander technique (24 lessons) + self-management (exercise prescription) versus usual care (without sciatica)

Quality assessment							No of patients			Effect	Quality	Importance
No of	Design	Risk of	Inconsistency	Indirectness	Imprecision	Other	Alexander techniques (24	Usual	Relative	Absolute		

studies

bias

							(exercise prescription)		CI)			
unction	(RMDQ 0-24)) - Functi	on (RMDQ 0-24) (Better indicated	d by lower valu	es)						
	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	71	72	-	MD 4.22 lower (6.13 to 2.31 lower)	⊕⊕⊕O MODERATE	CRITICAL
Pain (Voi	n Korff scale	0-10) - Pa	ain (Von Korff sc	ale 0-10) (Better	indicated by le	ower values)						
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	71	72	ı	MD 1.63 lower (2.49 to 0.77 lower)	⊕⊕OO LOW	CRITICAL
Quality o	f life: SF-36 r	nental - C	Quality of life: SF	-36 mental (Bett	er indicated by	lower values)						
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	71	72	-	MD 4.99 higher (2.31 lower to 12.29 higher)	⊕⊕OO LOW	CRITICAL
Quality o	f life: SF-36 p	hysical -	Quality of life: S	F-36 physical (I	Better indicated	d by lower values)						
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious²	none	71	72	-	MD 9.43 higher (1.88 to 16.98 higher)	⊕⊕OO LOW	CRITICAL

considerations lessons) + self management care

(95%

Table 133: Alexander technique (10 sessions) + mixed exercise versus usual care (overall population)

			Quality ass	essment			No of patients			Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alexander technique (10 lessons) + mixed exercise versus usual care	Control Relative (95% Absolute CI)			Quality	Importance
Overall - I	Function (RM	DQ 0-24)	<4 months [mean	difference from	control] (Bet	ter indicated by l	ower values)					
1	randomised trials			no serious indirectness	very serious ^b	none	15	13		MD 0.75 lower (4.21 lower to 2.72 higher)		CRITICAL

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments

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Overall -	Overall - Pain (von Korff 0-100) <4 months [mean difference from control] (Better indicated by lower values)													
1	randomised trials		no serious inconsistency	no serious indirectness	serious ^b	none	15	13		MD 1.27 lower (2.63 lower to 0.1 higher)	⊕⊕OO LOW	CRITICAL		
Overall -	Overall - Function (RMDQ 0-24) 4 months - 1 year [mean difference from control] (Better indicated by lower values)													
1	randomised trials		no serious inconsistency	no serious indirectness	serious ^b	none	15	13		MD 2.51 lower (6.21 lower to 1.19 higher)		CRITICAL		
Overall -	Pain (von Kor	ff 0-100)	4 months - 1 year	[mean differenc	e from contro	ol] (Better indicate	ed by lower values)							
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	15	13		MD 0.59 lower (2.04 lower to 0.86 higher)		CRITICAL		

Low back pain and sciatica in over 16s Quality assessment

Table 134: Combined interventions: Alexander technique (10 sessions) + mixed exercise versus mixed exercise (overall)

			Quality asse	essment			No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alexander technique (10 sessions) + mixed exercise versus mixed exercise	Control	Relative (95% CI)	Absolute	Quanty	importance
Overall -	Function (RM	DQ 0-24) ·	<4 months (Better	indicated by lo	wer values)							
1	randomised trials			no serious indirectness	very serious ^b	none	15	14	-	MD 0.45 higher (3.4 lower to 4.3 higher)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Orthotics J.7

Table 135: Back belts versus usual care (low back pain population)

			Quality asse	essment			No of pat	ients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Belts/corsets	Usual care	Relative (95% CI)	Absolute		
Function (follow-up 3 m	onths; me	asured with: EIFE	L (French version	of RMDQ); r	ange of scores: 0-	24; Better ind	icated by	lower values)			
	randomised trials	,	no serious inconsistency	no serious indirectness	serious ^b	none	98	92	1	MD 1.5 lower (2.8 to 0.2 lower)	⊕OOO VERY LOW	CRITICAL
Pain seve	rity (follow-up	3 months	; measured with: F	ain visual analog	ue scale; rar	nge of scores: 0-10); Better indica	ated by lo	wer values)			
	randomised trials	,	no serious inconsistency	no serious indirectness	serious ^b	none	98	92	1	MD 0.95 lower (1.54 to 0.36 lower)	⊕OOO VERY LOW	CRITICAL
Responde	r criteria (pair	n complete	ely improved) (folio	ow-up ≤4 months)								
	randomised trials	- ,	no serious inconsistency		very serious ^b	none	5/30 (16.7%)	3/29 (10.3%)	RR 1.61 (0.42 to 6.14)	63 more per 1000 (from 60 fewer to 532 more)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 136: Corset versus usual care (low back pain population)

			ar care (low back	- paritir paparata	•,							
			Quality asse	ssment			No of patient	s		Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Corsets/belts v. usual care	Control	Relative (95% CI)	Absolute	Quality	Importance

Change ir	function (all c	orsets) (fo	ollow-up 2 weeks; E	Better indicated by	/ higher valu	ies)						
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	69	58	-	MD 8.48 higher (3.59 to 13.38 higher)	⊕⊕OO LOW	CRITICAL
Change ir	function - Ine	xtensible (orthotics (follow-up	2 weeks; Better i	ndicated by	higher values)						
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	37	29	-	MD 11.6 higher (4.47 to 18.73 higher)	⊕⊕OO LOW	CRITICAL
Change ir	ı function - Ext	ensible or	thotics (follow-up 2	2 weeks; Better in	dicated by h	igher values)						
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	32	29	-	MD 5.7 higher (1.03 lower to 12.43 higher)	⊕⊕OO LOW	CRITICAL
Change ir	pain (all corse	ets) (follov	v-up 2 weeks; Bette	er indicated by hig	her values)							
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	69	68	-	MD 0.9 higher (0.09 lower to 1.89 higher)	⊕⊕OO LOW	CRITICAL
Change ir	pain - Inexten	sible orth	otics (follow-up 2 w	eeks; Better indic	ated by high	her values)						
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	37	39	-	MD 0.9 higher (0.47 lower to 2.27 higher)	⊕⊕OO LOW	CRITICAL
Change ir	pain - Extensi	ible orthot	ics (follow-up 2 we	eks; Better indica	ted by highe	er values)						
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	32	29	-	MD 0.9 higher (0.53 lower to 2.33 higher)	⊕⊕OO LOW	CRITICAL

Table 137: Belts/corsets versus manipulation (low back pain population)

			Quality asse	ssment			No of p	patients		Effect	Quality	Importance
No of studies							Belts/corsets	Manipulation	Relative (95% CI)	Absolute	Š	

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Function	(follow-up.3 v	veeks; mea	asured with: Revis	sed ODI; range o	f scores: 0-1	00; Better indicate	d by lower val	ues)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	12	26	-	MD 10.85 higher (1.77 to 19.93 higher)	⊕OOO VERY LOW	CRITICAL		
Pain seve	Pain severity (follow-up 3 weeks; measured with: Pain visual analogue scale 1-10; range of scores: 0-100; Better indicated by lower values)													
1	randomised trials		no serious inconsistency	no serious indirectness	serious ^b	none	25	65	-	MD 0.82 higher (0.43 lower to 2.65 higher)	⊕⊕OO LOW	CRITICAL		
Responde	er criteria (im	proved pai	n) (follow-up ≤4 m	onths)										
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	27/93 (29%)	44/98 (44.9%)	RR 0.65 (0.44 to 0.95)	157 fewer per 1000 (from 22 fewer to 251 fewer)	⊕OOO VERY LOW	CRITICAL		

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 138: Belt/corset versus massage (low back pain population)

			Quality asses	ssment			No of pat	ients		Effect	Overlife	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Belts/corsets		Relative (95% CI)	Absolute	Quality	Importance
Function (1	ollow-up 3 we	eks; meas	ured with: ODI; rang	ge of scores: 0-10	cated by lower val	ues)						
	randomised trials	- ,		no serious indirectness	serious ^b	none	12	15	-	MD 11.67 lower (23.69 lower to 0.35 higher)	⊕OOO VERY LOW	CRITICAL
Pain sever	ity (follow-up 3	3 weeks; m	neasured with: Pain	visual analogue s	cale; range o	of scores: 0-100; B	etter indicated	l by lowe	values)			
	randomised trials	serious ^a		no serious indirectness	serious ^b	none	25	32	-	MD 0.13 higher (1.24 lower to 1.5 higher)	⊕⊕OO LOW	CRITICAL

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias

Table 139: Corset versus non-opioid analgesic (low back pain population)

			Quality asse	essment			No of patien	ts		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Corsets versus paracetamol	Control	Relative (95% CI)			
Responde	er criteria (imp	roved pai	n) (follow-up ≤4 m	onths)								
1		- ,		no serious indirectness	very serious ^b	none	27/93 (29%)	33/100 (33%)	RR 0.88 (0.58 to 1.34)	40 fewer per 1000 (from 139 fewer to 112 more)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 140: Foot orthotics versus placebo (low back pain and sciatica population)

			Quality as	sessment			No of	f patients		Effect	Quality		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Foot orthotics	Placebo/sham	Relative (95% CI)	Absolute	Quality	Importance	
Function (follow-up 4 w	eeks; mea	sured with: ODI; ra	ange of scores: 0	-100; Better indi	cated by lower val	ues)						
	randomised trials		no serious inconsistency		no serious imprecision	none	29	22	-	MD 12.95 lower (17.88 to 8.02 lower)	⊕⊕⊕O MODERATE	CRITICAL	
Pain sever	in severity (follow-up 4 weeks; measured with: Pain visual analogue scale; range of scores: 0-100; Better indicated by lower values)												
	randomised trials		no serious inconsistency		no serious imprecision	none	29	22	-	MD 3.47 lower (4.43 to 2.51 lower)	⊕⊕⊕O MODERATE	CRITICAL	

a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias

^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 141: Rocker sole shoes versus placebo/sham (flat sole shoes) (low back pain population)

Table 14	FI. NUCKEI 3	OIE SIIUE	es versus piacer	oo, silalii (liat s	sole silves/ (ic	ow back pain po						
			Quality as	sessment			No of patien	ts		Effect	Quality	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Foot orthotics versus usual care	Control	Relative (95% CI)	Absolute	Quanty	Importance
Function:	≤4 months (fo	llow-up 6	weeks; measured	with: Roland Mo	rris disability qu	estionnaire; range	e of scores: 0-24; B	etter ind	icated by	lower values)		
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	50	50	-	MD 1.2 lower (3.07 lower to 0.67 higher)	⊕⊕OO LOW	CRITICAL
Function	>4 months - 1	year (foll	ow-up 12 months;	range of scores:	0-24; Better ind	icated by lower va	lues)					
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	44	49	1	MD 0.8 lower (2.8 lower to 1.2 higher)	⊕⊕OO LOW	CRITICAL
Pain ≤4 m	onths (follow	-up 6 wee	ks; range of score	s: 0-10; Better in	dicated by lower	r values)						
1	randomised trials	seriousª	no serious inconsistency	no serious indirectness	serious ^b	none	50	50	-	MD 0.30 lower (1.2 lower to 0.6 higher)	⊕⊕OO LOW	CRITICAL
Pain >4 m	onths - 1 yea	r (follow-u	ıp 12 months; rang	je of scores: 0-10); Better indicate	ed by lower values)					
1	randomised trials	seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	44	49	ı	MD 0 higher (1.25 lower to 1.25 higher)	⊕⊕⊕O MODERATE	CRITICAL
Anxiety ≤₄	4 months (foll	ow-up 6 v	veeks; range of sc	ores: 0-21; Bette	r indicated by lo	wer values)						
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	50	50	1	MD 1.3 higher (0.62 lower to 3.22 higher)	⊕⊕OO LOW	CRITICAL
Anxiety >	4 months - 1 y	ear (follo	w-up 12 months; E	Setter indicated b	y lower values)							
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	44	49	1	MD 0.3 higher (1.59 lower to 2.19 higher)	⊕⊕⊕O MODERATE	CRITICAL
Depression	on ≤4 months	(follow-up	o 6 weeks; range o	f scores: 0-21; B	etter indicated b	y lower values)						
1	randomised	serious ^a	no serious	no serious	serious ^b	none	50	50	-	MD 0.9 higher (0.81	⊕⊕OO	CRITICAL

	trials		inconsistency	indirectness						lower to 2.61 higher)	LOW	
Depression >4 months - 1 year (follow-up 12 months; Better indicated by lower values)												
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	44	49	-	MD 0.8 higher (0.94 lower to 2.54 higher)	⊕⊕OO LOW	CRITICAL
EQ-5D ≤4 months (follow-up 6 weeks; range of scores: 0-1; Better indicated by higher values)												
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	49	50	-	MD 0.1 lower (0.24 lower to 0.04 higher)	⊕⊕OO LOW	CRITICAL
EQ-5D >4 months - 1 year (range of scores: 0-1; Better indicated by higher values)												
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	44	49	-	MD 0.10 lower (0.24 lower to 0.4 higher)	⊕⊕OO LOW	CRITICAL

Low back pain and sciatica in over 16s Quality assessment

Table 142: Foot orthotics versus usual care (low back pain and sciatica population)

			Quality asses	ssment			No of pa	tients		Effect	Quality	Importance		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Foot orthotics	Usual care	Relative (95% CI)	Absolute	Quality	Importance		
Function (f	unction (follow-up 6 weeks; measured with: ODI; range of scores: 0-50; Better indicated by lower values)													
1	randomised trials	very serious ^a		no serious indirectness	serious ^b	none	23	25	-	MD 8 lower (14 to 2 lower)	⊕000 VERY LOW	CRITICAL		
Pain sever	ity (follow-up r	nean 6 wee	ks; measured with:	Pain visual analog	gue scale; ra	nge of scores: 0-10	; Better indic	ated by	ower val	ues)				
1	randomised trials	very serious ^a		no serious indirectness	serious ^b	none	23	25	-	MD 1.3 lower (2.69 lower to 0.09 higher)	⊕OOO VERY LOW	CRITICAL		

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias

Table 143: Foot orthotics versus usual care (non-randomised study) (low back pain and sciatica population)

			Quality assess	ment			No of pat	tients		Effect	Ovality	
No of studies	Design Risk of bias Inconsistency		Inconsistency	Indirectness	Imprecision	Other considerations	Foot orthotics	Usual care	Relative (95% CI)	Absolute	Quality	Importance
Function (f	ollow-up 8 weeks	; measured	d with: ODI; range of	scores: 0-100; Be	tter indicated	d by lower values)						
	observational studies	,		no serious indirectness	Serious ^b	none	30	34	-	MD 6.9 lower (12.2 to 1.6 lower)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Combinations of interventions – orthotics adjunct J.7.1

Low back pain with or without sciatica

Table 144: Orthotics (corset) plus electrotherapy plus massage plus traction compared with electrotherapy plus mixed modality manual therapy (massage plus traction)

Quality	Quality assessment								Effect			
No of studie s	Design	Risk of bias	Inconsistenc y	Indirectness	Imprecision	Other	Corset + electrotherapy + massage + traction	Electrotherapy + massage + traction	Relativ e (95% CI)	Absolute	Quality	Importance
Pain (0-	100 VAS conver	ted to 0-10	scale) - ≤4 mont	hs (Better indicat	ted by lower va	lues)						
1: he 2006	Randomised trials	Very serious ^a	No serious inconsistency	No serious indirectness	No serious imprecision	None	29	29	-	MD 1.02 lower (1.7 to 0.33	LOW	CRITICAL

^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Quality	assessment					No of patients		Effect				
No of studie s	Design	Risk of bias	Inconsistenc y	Indirectness	Imprecision	Other	Corset + electrotherapy + massage + traction	Electrotherapy + massage + traction	Relativ e (95% CI)	Absolute	Quality	Importance
Function	l n (Japanese Ort	hopaedics .	L Academic Assoc	iation) lumbar d	 isease grade (0	-29) - ≤4 r	 nonths (Better indica	ted by lower values))	lower)		
1: he 2006	Randomised trials	Very serious ^a	No serious inconsistency	No serious indirectness	No serious imprecision	None	29	29	-	MD 3.17 higher (1.5 to 4.84 higher)	LOW	CRITICAL

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and by 2 increments if the majority of evidence was at very high risk of bias

J.8 Manual therapies

J.8.1 Soft tissue techniques

Table 145: Soft tissue techniques (massage) versus sham in low back pain without sciatica

TUDIC IT	5. 5616 61554		ques (Illassage)	versus situiti ii	riott back par	ii Witiioat Sciati	<u></u>						
			Quality ass	sessment			No of patie	ents		Effect			
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Massage versus sham	Control	Relative (95% Absolute CI)		Quality	Importance	
Pain (VAS	0-10) <4 mont	hs (range	of scores: 0-10; Be	tter indicated by	lower values)								
		very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	36	36	-	MD 1.01 lower (2.03 lower to 0.02 higher)	⊕000 VERY LOW	CRITICAL	
Pain (McG	Pain (McGill score 0-78) <4 months (range of scores: 0-78; Better indicated by lower values)												

3		- 3		no serious indirectness	Serious ^b	none	74	72	-	MD 4.73 lower (7.56 to 1.9 lower)	⊕OOO VERY LOW	CRITICAL	
Function	Function (Quebec Disability Score 0-100) <4 months (range of scores: 0-100; Better indicated by lower values)												
3		very seriousª			no serious imprecision	none	74	72	-	MD 4.3 lower (8.28 to 0.32 lower)	⊕⊕OO LOW	CRITICAL	

Low back pain and sciatica in over 16s Quality assessment

Table 146: Soft tissue techniques (massage) versus usual care in low back pain without sciatica

			Quality as	sessment			No of patier	nts		Effect	0		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Massage versus usual care	Control	Relative (95% CI)	Absolute	Quality	Importance	
Pain (Von	in (Von Korff scale 0-10) <4 months (Better indicated by lower values)												
	randomised trials		no serious inconsistency		no serious imprecision	none	120	103	-	MD 0.41 lower (0.91 lower to 0.09 higher)	⊕⊕⊕O MODERATE	CRITICAL	
Pain (Von	ain (Von Korff scale 0-10) >4 months (Better indicated by lower values)												
	randomised trials		no serious inconsistency		no serious imprecision	none	120	111	1	MD 0.01 lower (0.65 lower to 0.63 higher)	⊕⊕⊕O MODERATE	CRITICAL	
Quality of	life composit	e scores (SF-36- Physical co	mponent 0-100)	<4 months (rang	e of scores: 0-100	; Better indicated	by high	er values	s)			
	randomised trials	very serious ^a	Serious ^b		no serious imprecision	none	247	226	-	MD 0.53 lower (1.62 lower to 0.56 higher)	⊕000 VERY LOW	CRITICAL	
Quality of	life composit	e scores (SF-36 - Mental con	nponent 0-100) <4	4 months (range	of scores: 0-100;	Better indicated b	y highe	r values)				
	randomised trials	- ,	no serious inconsistency	no serious indirectness	Serious ^c	none	247	226	-	MD 2.43 higher (0.71 to 4.14 higher)	⊕000 VERY LOW	CRITICAL	

^a Downgraded by one increment if the majority of the evidence was at high risk of bias, and downgraded by two increments if the majority of the evidence was at very high risk of bias ^b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

Quality of	Quality of life composite scores (SF-36 - Physical component 0-100) > 4 months (range of scores: 0-100; Better indicated by higher values)													
2	randomised trials	very	no serious inconsistency	no serious	no serious imprecision	none	247	227	-	MD 0.08 higher (1.15 lower to 1.31 higher)	⊕⊕OO LOW	CRITICAL		
Quality of	Quality of life composite scores (SF-36- Mental component 0-100) > 4 months (range of scores: 0-100; Better indicated by higher values)													
2	randomised trials	,	no serious inconsistency		no serious imprecision	none	247	227	-	MD 0.41 higher (1.66 lower to 2.48 higher)	⊕⊕OO LOW	CRITICAL		
Function	Function (RMDQ 0-24) <4 months (range of scores: 0-24; Better indicated by lower values)													
2	randomised trials	- ,	no serious inconsistency	no serious indirectness	Serious ^c	none	247	226	-	MD 2.27 lower (3.07 to 1.47 lower)	⊕OOO VERY LOW	CRITICAL		
Function	(RMDQ 0-24)	> 4 months	s (range of scores:	: 0-24; Better indi	cated by lower v	ralues)								
2	randomised trials	- ,	no serious inconsistency	no serious indirectness	Serious ^c	none	247	227	-	MD 0.35 lower (1.22 lower to 0.51 higher)	⊕OOO VERY LOW	CRITICAL		

^a Downgraded by one increment if the majority of the evidence was at high risk of bias, and downgraded by two increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment because of heterogeneity, I2=42%, p=0.19) ^c Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

Table 147: Soft tissue techniques (massage) versus acupuncture in low back pain without sciatica

TUDIC IT	7 . 5016 61556	40 (00)	ques (massage	, versus acupa	netare in low	Back pain With	Jul Jeiutica					
			Quality as:	sessment			No of patient	s		Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Massage versus acupuncture	Control	Relative (95% Absolute CI)		Quality	Importance
Function (RMDQ 0-24) <	<4 months	(range of scores:	0-24; Better indic	ated by lower va	alues)						
1	randomised trials	- 3	no serious inconsistency	no serious indirectness	Serious ^b	none	77	89	-	MD 1.6 lower (3.44 lower to 0.24 higher)	⊕OOO VERY LOW	CRITICAL
Function (Function (RMDQ 0-24) > 4 months (range of scores: 0-24; Better indicated by lower values)											

1 randomised very no serious no serious no serious indirectness imprecision	none 76	90 -	MD 1.2 lower (3.12 lower to 0.72 higher) LOW	CRITICAL
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^a Downgraded by one increment if the majority of the evidence was at high risk of bias, and downgraded by two increments if the majority of the evidence was at very high risk of bias

Table 148: Soft tissue techniques (massage) versus self-management in low back pain without sciatica

			Quality as	sessment			No of patients			Effect	Ovality	
No of studies	I Design I Inconsistency I indirectness I imprecision					Other considerations	Massage versus self- management	Control	Relative (95% CI)	Absolute	Quanty	Importance
Function ((RMDQ 0-24)	<4 months	(range of scores:	0-24; Better indi	cated by lower v	alues)						
	randomised trials	,		no serious indirectness	Serious ^b	none	77	83	-	MD 2.5 lower (4.35 to 0.65 lower)	⊕OOO VERY LOW	CRITICAL
Function (RMDQ 0-24) > 4 months (range of scores: 0-24; Better indicated by lower values)												
	randomised trials	- ,			no serious imprecision	none	76	83	-	MD 0.4 higher (1.43 lower to 2.23 higher)	⊕⊕OO LOW	CRITICAL

^a Downgraded by one increment if the majority of the evidence was at high risk of bias, and downgraded by two increments if the majority of the evidence was at very high risk of bias

J.8.2 Traction

Table 149: Traction versus sham in low back pain with or without sciatica (mixed population)

			Quality asse	essment			No of pati	ients		Effect	Quality	Importance
o of udies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Traction versus sham	Control	Relative (95% CI)	Absolute		•

^b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

^b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

Pain VAS	S (0-10) <4 ma	nths (mecha	nical traction) (Be	etter indicated b	y lower values)							
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^a	none	77	73	-	MD 0.56 higher (0.46 lower to 1.58 higher)	⊕⊕⊕O MODERATE	CRITICAL
Pain VAS	S (0-10) <4 mo	nths (inversi	on traction) (Bett	er indicated by I	ower values)							
1	randomised trials	Serious ^b	no serious inconsistency	no serious indirectness	no serious imprecision	none	14	15	-	MD 1.59 lower (2.44 to 0.74 lower)	⊕⊕⊕O MODERATE	CRITICAL
Pain VAS	S (0-10) > 4 me	onths (range	of scores: 0-10; I	Better indicated	by lower values	s)						
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	no serious imprecision	none	76	72	-	MD 0.37 higher (0.84 lower to 1.58 higher)	⊕⊕⊕⊕ HIGH	CRITICAL
Function	(RMDQ 0-24)	<4 months (range of scores:	0-24; Better indi	cated by lower	values)						
1	randomised trials	Serious ^b	no serious inconsistency	no serious indirectness	no serious imprecision	none	77	73	-	MD 0.10 higher (1.8 lower to 2 higher)	⊕⊕⊕O MODERATE	CRITICAL
Function	(RMDQ 0-24)	> 4 months	range of scores:	0-24; Better ind	icated by lower	values)						
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	no serious imprecision	none	76	72	-	MD 0.7 higher (1.1 lower to 2.5 higher)	⊕⊕⊕⊕ HIGH	CRITICAL
Healthca	re utilisation	- other medic	cal treatments so	ught <4 months								
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^a	none	26/77 (33.8%)	18/73 (24.7%)	RR 1.37 (0.82 to 2.28)	91 more per 1000 (from 44 fewer to 316 more)	0000	IMPORTANT
								0%	·	-		
Healthca	re utilisation	other medic	cal treatments so	ught > 4 months				, ,				
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	very serious ^a	none	34/76 (44.7%)	30/72 (41.7%)	RR 1.07 (0.74 to 1.55)	29 more per 1000 (from 108 fewer to 229 more)	⊕⊕OO LOW	IMPORTANT
					1			0%		-		

^a Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs ^b Downgraded by one increment if the majority of the evidence was at high risk of bias, and downgraded by two increments if the majority of the evidence was at very high risk of bias

Table 150: Traction versus sham in low back pain without sciatica

			ann in low back p									
			Quality as:	sessment			No of pa	itients		Effect	Quality	I m n a utan a a
No of studies	Design Inconsistency Indirectness Imprecision							Absolute	Quality	Importance		
Pain VAS (0-10) <4 month	ns (range o	of scores: 0-10; Bette	er indicated by low	ver values)							
	randomised trials	Serious ^a			no serious imprecision	none	29	31	-	MD 0.4 lower (1.76 lower to 0.96 higher)	⊕⊕⊕O MODERATE	CRITICAL

a Downgraded by one increment if the majority of the evidence was at high risk of bias, and downgraded by two increments if the majority of the evidence was at very high risk of bias

Table 151: Traction versus usual care in low back pain with or without sciatica (mixed population)

			Quality asses	sment			No of p	atients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Imprecision	Other considerations	Traction		Relative (95% CI)	Absolute	Quanty	importance	
Pain VAS (0	ain VAS (0-10) <4 months (range of scores: 0-10; Better indicated by lower values)											
		very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	20	19	-	MD 0.5 higher (0.57 lower to 1.57 higher)	⊕OOO VERY LOW	CRITICAL
Function (ODI, 0-100) <4 months (range of scores: 0-24; Better indicated by lower values)												
		very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	20	19	-	MD 4 higher (2.78 lower to 10.78 higher)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by one increment if the majority of the evidence was at high risk of bias, and downgraded by two increments if the majority of the evidence was at very high risk of bias ^b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

Table 152: Traction versus usual care in low back pain with sciatica

			Quality asse	essment			No of patier	nts		Effect	Ovelite	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Traction versus usual care	Control	Relative (95% CI)	Absolute	Quanty	importance
Quality of	Life (SF-36 - 0	eneral hea	alth 0-100) <4 mont	hs (range of score	es: 0-100; Be	tter indicated by h	igher values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	18	18	-	MD 21.91 higher (6.82 to 37 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	Life (SF-36 - P	hysical fu	nction 0-100) <4 m	onths (range of so	ores: 0-100;	Better indicated by	y higher values)					
1	randomised trials	very seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	18	18	-	MD 14.91 higher (1.22 lower to 31.04 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	Life (SF-36 - P	hysical ro	le limitation 0-100)	<4 months (range	of scores: 0	-)-100; Better indica	ted by higher valu	ies)				
1	randomised trials	very seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	18	18	-	MD 26.88 higher (1.46 to 52.3 higher)	⊕000 VERY LOW	CRITICAL
Quality of	Life (SF-36 - E	odily pain	0-100) <4 months	(range of scores:	0-100; Better	r indicated by high	er values)	•				
1	randomised trials	very seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	18	18	-	MD 16.07 higher (3.91 to 28.23 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	Life (SF-36 - V	itality 0-10	00) <4 months (rang	ge of scores: 0-10	0; Better indi	cated by higher va	lues)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	18	18	-	MD 20.67 higher (3.08 to 38.26 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	Life (SF-36 - S	ocial func	tion 0-100) <4 mon	ths (range of scor	res: 0-100; Be	etter indicated by h	igher values)					
1	randomised	very	no serious	no serious	Serious ^b	none	18	18	-	MD 18.55 higher (0.43 to	⊕000	CRITICAL

VERY

LOW

36.67 higher)

NICE, 2016

trials

seriousa

inconsistency

indirectness

Quality of	Life (SF-36 - N	lental heal	th 0-100) <4 month	s (range of scores	s: 0-100; Bet	ter indicated by hig	her values)							
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	18	18	-	MD 20.65 higher (2.17 to 39.13 higher)	⊕OOO VERY LOW	CRITICAL		
Quality of	Life (SF-36 - E	motional r	ole limitation 0-100)) <4 months (rang	ge of scores:	: 0-100; Better indic	ated by higher va	lues)						
1	randomised trials	very seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	18	18	-	MD 36.87 higher (9.13 to 64.61 higher)	⊕OOO VERY LOW	CRITICAL		
Function (Function (ODI 0-100) <4 months (range of scores: 0-100; Better indicated by lower values)													
2	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	49	51	-	MD 5.98 higher (0.82 lower to 12.77 higher)	⊕⊕OO LOW	CRITICAL		
Pain VAS	(0-10) <4 mont	ths (weight	tbath traction) (rang	ge of scores: 0-10	; Better indi	cated by lower valu	es)							
1	randomised trials	very seriousª	no serious inconsistency	no serious indirectness	very serious ^b	none	18	18	-	MD 2.98 lower (4.51 to 1.45 lower)	⊕OOO VERY LOW	CRITICAL		
Pain VAS	(0-10) <4 mont	ths (mecha	nical traction) (ran	ge of scores: 0-10); Better indi	cated by lower valu	ies)							
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	31	33	-	MD 0.2 higher (1 lower to 1.4 higher)	⊕OOO VERY LOW	CRITICAL		

^a Downgraded by one increment if the majority of the evidence was at high risk of bias, and downgraded by two increments if the majority of the evidence was at very high risk of bias ^b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

Table 153: Traction versus exercise (biomechanical) in low back pain with or without sciatica (mixed population)

			Quality asses	ssment			No of patients	•		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Traction versus biomechanical	Control	Relative (95% CI)	Absolute	•	·

							exercise							
Healthca	Healthcare utilisation - visited other healthcare practitioners > 4 months													
1	randomised trials			no serious indirectness	Serious ^a	none	41/107 (38.3%)	45/84 (53.6%)	RR 0.72 (0.52 to 0.98)	150 fewer per 1000 (from 11 fewer to 257 fewer)	⊕⊕⊕O MODERATE	CRITICAL		

a Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

J.8.3 Manipulation/mobilisation

Table 154: Manipulation/mobilisation versus sham in low back pain without sciatica

			Quality ass	essment			No of patients			Effect	O life	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation/mobilisation versus sham	Control	Relative (95% CI)	Absolute	Quality	Importance
Quality o	of life (Euroqu	ol health sta	te 0-100) < 4 moi	nths (Better ind	icated by high	er values)						
	randomised trials			no serious indirectness	Serious ^a	none	89	85	-	MD 4.4 higher (0.42 lower to 9.22 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality o	of life (Euroqu	l health sta	te 0-100) > 4 moi	nths (Better ind	icated by high	er values)						
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	85	81	-	MD 2.5 higher (2.43 lower to 7.43 higher)	⊕⊕⊕⊕ HIGH	CRITICAL
Quality of life (SF-12/SF36 - Physical composite score0-100) <4 months (Better indicated by higher values)												
	randomised trials			no serious indirectness	Serious ^a	none	89	85	-	MD 4.1 higher (1.29 to 6.91 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality o	of life (SF-12/S	SF36- Menta	al composite sco	re 0-100) <4 mo	nths (Better in	dicated by higher	r values)					

1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^a	none	89	85	-	MD 2.4 lower (5.64 lower to 0.84 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality of	of life (SF-12/	SF36- Pain	subscale 0-100)	<4 months (Bet	ter indicated b	y higher values)						
1	randomised trials	very serious ^b	no serious inconsistency	no serious indirectness	Serious ^a	none	69	67	-	MD 0.11 higher (0.48 lower to 0.7 higher)	⊕000 VERY LOW	CRITICAL
Quality of	of life (SF-12/	SF36 - Phys	sical function sul	oscale0-100) <4	months (Bette	er indicated by hig	her values)					
1	randomised trials	very serious ^b	no serious inconsistency	no serious indirectness	no serious imprecision	none	69	67	-	MD 0.01 lower (0.18 lower to 0.16 higher)	⊕⊕OO LOW	CRITICAL
Quality of	of life (SF-12	0-100) > 4 m	nonths (Better in	dicated by high	er values)							
1			no serious inconsistency	no serious indirectness	no serious imprecision	none	85	81	-	MD 1.9 higher (1.51 lower to 5.31 higher)	⊕⊕⊕ HIGH	CRITICAL
Quality of	of life (SF-12	- Physical c	omposite score	0-100) 4 month:	s - 1 year - Mer	ntal composite sco	ore (Better indicated by highe	r values)				
1		no serious risk of bias	no serious inconsistency	no serious indirectness	no serious imprecision	none	85	81	-	MD 0.7 lower (4.46 lower to 3.06 higher)	⊕⊕⊕⊕ HIGH	CRITICAL
Pain (VA	AS 0-10) <4 m	onths (Bett	er indicated by lo	ower values)								
5	randomised trials	Serious ^b	no serious inconsistency	no serious indirectness	no serious imprecision	none	265	268	-	MD 0.30 lower (0.56 to 0.04 lower)	⊕⊕⊕O MODERATE	CRITICAL
Pain (VA	AS 0-10) > 4 m	onths (Bett	ter indicated by I	ower values)								
2	randomised	no serious	no serious inconsistency	no serious indirectness	no serious imprecision	none	111	118	-	MD 0.20 lower (0.67 lower to 0.26 higher)	⊕⊕⊕⊕ HIGH	CRITICAL
Function	n (ODI 0-100)	<4 months	(Better indicated	by lower value	es)							
4	randomised trials	Serious ^b	no serious inconsistency	no serious indirectness	Serious ^a	none	180	194	-	MD 3.91 lower (6.47 to 1.34	⊕⊕OO LOW	CRITICAL

		1					T				1	
										lower)		
Function	(Von Korff, (0-100) < 4 m	nonths (range of	scores: 0-100; I	Better indicate	d by lower values)					
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^a	none	89	85	-	MD 7.2 lower (13.82 to 0.58 lower)	⊕⊕⊕O MODERATE	CRITICAL
Function	(ODI 0-100)	> 4 months	(Better indicated	l by lower value	es)							
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^a	none	26	37	-	MD 2.53 lower (8.85 lower to 3.79 higher)	⊕⊕⊕O MODERATE	CRITICAL
Function	(Von Korff, (0-100) > 4 m	nonths (range of	scores: 0-100; I	Better indicate	d by lower values)					
1		no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^a	none	85	81	-	MD 5.6 lower (12.45 to 1.25 lower)	⊕⊕⊕O MODERATE	CRITICAL

Table 155: Manipulation/mobilisation versus sham in low back pain with sciatica

		No of patients		Effect	Quality	Importance						
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation/mobilisation	Sham	Relative (95% CI)	Absolute		
Quality o	Quality of life (SF-36 0-100 - Physical functioning) >4 months (range of scores: 0-100; Better indicated by higher values)											
		no serious risk of bias		no serious indirectness	serious	none	48	48	-	MD 6.9 higher (1.23 lower to 15.03 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality o	f life (SF-36 0)-100 - Phys	sical role limitati	on) >4 months	(range of score	es: 0-100; Better i	ndicated by higher values)					
		no serious risk of bias			no serious imprecision	none	48	48	-	MD 2 higher (13.04 lower to 17.04 higher)	⊕⊕⊕⊕ HIGH	CRITICAL

^a Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs ^b Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Quality o	f life (SF-36 0)-100 - Bod	ily pain) >4 mon	ths (range of so	ores: 0-100; B	etter indicated by	higher values)	_				
		no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^a	none	48	48	-	MD 1.9 higher (3.33 lower to 7.13 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality o	f life (SF-36 0)-100 - Gen	eral health) >4 n	nonths (range o	f scores: 0-100); Better indicated	by higher values)					
		no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^a	none	48	48	-	MD 3.7 lower (11.09 lower to 3.69 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality o	f life (SF-36 0)-100 - Vita	lity) >4 months (range of scores	s: 0-100; Better	indicated by high	ner values)					
		no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^a	none	48	48	-	MD 5.6 higher (0.52 lower to 11.72 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality o	f life (SF-36 0)-100 - Soc	ial functioning)	>4 months (ran	ge of scores: 0	-100; Better indica	ated by higher values)					
		no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^a	none	48	48	-	MD 5.7 higher (0.31 lower to 11.71 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality o	f life (SF-36 0)-100 - Emo	otional role limita	ation) >4 month	s (range of sco	ores: 0-100; Better	r indicated by higher value	es)				
		no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^a	none	48	48	-	MD 7.2 higher (9.72 lower to 24.12 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality o	f life (SF-36 0)-100 - Men	tal health) >4 me	onths (range of	scores: 0-100;	Better indicated	by higher values)	•				
		no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^a	none	48	48	-	MD 3.3 higher (3.04 lower to 9.64 higher)	⊕⊕⊕O MODERATE	CRITICAL
Respond	er criteria (>3	30% VAS p	ain - Local back	pain) > 4 montl	าร					- · ·		
		no serious risk of bias	no serious inconsistency	no serious indirectness	very serious ^a	none	15/48 (31.3%)	6.3%	RR 5 (1.55 to 16.16)	252 more per 1000 (from 35 more to 955 more)	⊕⊕OO LOW	IMPORTANT
Respond	er criteria (>3	30% VAS p	ain - Radiating p	pain) > 4 months	<u> </u>							

1		no serious risk of bias	no serious inconsistency	no serious indirectness	very serious ^a	none	29/48 (60.4%)	20.8%	RR 2.9 (1.6 to 5.27)	395 more per 1000 (from 125 more to 888 more)	⊕⊕OO LOW	IMPORTANT
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^a Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 156: Manipulation/mobilisation versus usual care in low back pain with or without sciatica (mixed population)

	Quality assessment						No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation/mobilisation versus usual care	Control	Relative (95% CI)	Absolute		
Pain (VAS 0-10) < 4 months (Better indicated by lower values)												
	randomised trials			no serious indirectness	very serious ^b	none	0	-	-	MD 0.03 higher (0.55 lower to 0.61 higher)	⊕OOO VERY LOW	CRITICAL
Pain (VAS 0-10) > 4 months (Better indicated by lower values)												
	randomised trials		no serious inconsistency		no serious imprecision	none	0	-	-	MD 0.22 higher (0.25 lower to 0.69 higher)	⊕⊕⊕O MODERATE	CRITICAL
Function	(RMDQ 0-24) <4 mon	ths (high velocit	y thrust) (Bette	r indicated by	lower values)						
		- ,	no serious inconsistency	no serious indirectness	Serious ^b	none	96	49	-	MD 1.5 lower (3.1 lower to 0.1 higher)	⊕000 VERY LOW	CRITICAL
Function	(RMDQ 0-24) <4 mon	ths (spinal adjus	sting - mobilisa	tion) (Better in	dicated by lower	values)	<u> </u>				
	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	169	170	-	MD 0.75 higher (0.29 lower to 1.79 higher)	⊕⊕OO LOW	CRITICAL
Function	(RMDQ 0-24) <4 mon	ths (traction gap	manipulation)	(Better indica	ted by lower valu	es)					
1	randomised	very	no serious	no serious	no serious	none	15	14	-	MD 3.31 lower	⊕⊕00	CRITICAL

	trials	seriousª	inconsistency	indirectness	imprecision					(4.83 to 1.79 lower)	LOW	
Function	(RMDQ 0-24	l) > 4 mor	nths (Better indic	ated by lower	values)							
	randomised trials	,		no serious indirectness	Serious ^b	none	0	-	-	MD 1.3 lower (2.9 lower to 0.3 higher)	⊕000 VERY LOW	CRITICAL
Quality o	of life (SF-36	- Physica	I function 0-100)	<4 months (Be	etter indicated	by lower values)						
	randomised trials	very serious ^a		no serious indirectness	Serious ^b	none	0	1	1	MD 4.3 higher (1.2 lower to 9.8 higher)	⊕OOO VERY LOW	CRITICAL
Healthca	Healthcare utilisation - Number of healthcare visits <4 months (Better indicated by lower values)											
	randomised trials	Serious ^a		no serious indirectness	no serious imprecision	none	169	169	-	MD 1.5 higher (1.22 to 1.78 higher)	⊕⊕⊕O MODERATE	IMPORTANT
Healthca	re utilisation	- Numbe	er of healthcare v	risits > 4 month	ns (Better indic	ated by lower va	lues)					
	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	165	165	-	MD 2.4 higher (1.63 to 3.17 higher)	⊕⊕OO LOW	IMPORTANT
Adverse	events <4 m	onths										
		very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	10/96 (10.4%)	4/49 (8.2%)	RR 1.28 (0.42 to 3.86)	23 more per 1000 (from 47 fewer to 233 more)	⊕000 VERY LOW	IMPORTANT
								0%		-		

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 157: Manipulation/mobilisation versus usual care in low back pain with sciatica

Quality assessment	No of patients	Effect	Quality	Importance
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No of	Design	Risk of	Inconsistency	Indirectness	Imprecision	Other	Manipulation/mobilisation	Usual	Relative	Absolute		
studies	Design	bias	meonsistency	munectness	Imprecision	considerations	mampalation/mobilisation	care	(95% CI)	Absolute		
Pain (0-1	0) <4 months	(Better in	ndicated by lower	· values)								
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	96	96	-	MD 0.9 lower (2.57 lower to 0.77 higher)	⊕OOO VERY LOW	CRITICAL
Pain (0-1	0) > 4 months	s (Better i	ndicated by lowe	r values)								
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	96	96	1	MD 0.4 lower (2.15 lower to 1.35 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	f life (SF-36 -	Physical	health composite	e, 0-100) <4 mon	iths (Better in	ndicated by lower	values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	96	96	-	MD 3.4 higher (3.23 lower to 10.03 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	f life (SF-36-	Mental he	alth composite, (0-100) <4 month	s (Better indi	cated by lower va	alues)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	96	96	-	MD 0 higher (4.76 lower to 4.76 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	f life (SF-36 -	Physical	health composite	e, 0-100) > 4 mo	nths (Better i	ndicated by lowe	r values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	96	96	1	MD 1.5 higher (4.85 lower to 7.85 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	f life (SF-36 -	Mental he	ealth composite)	> 4 months (Bet	tter indicated	by lower values)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	96	96	-	MD 0.7 higher (4.88 lower to 6.28 higher)	⊕OOO VERY LOW	CRITICAL
Function	(RMDQ 0-24)	<4 montl	hs (Better indicat	ed by lower val	ues)							
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	96	96	-	MD 2.5 lower (6.27 lower to 1.27 higher)	⊕000 VERY	CRITICAL

Low back pain and sciatica in over 16s Quality assessment

											LOW		
Function (RMDQ 0-24) > 4 months (Better indicated by lower values)													
1		- ,	no serious inconsistency	no serious indirectness	Serious ^b	none	96	96	-	MD 1.3 lower (5.07 lower to 2.47 higher)		CRITICAL	
Adverse	Adverse events <4 months												
1		- ,	no serious inconsistency	no serious indirectness	Serious ^b	none	29/96 (30.2%)	40/49 (81.6%)	RR 0.72 (0.49 to 1.07)	229 fewer per 1000 (from 416 fewer to 57 more)	⊕OOO VERY LOW	IMPORTANT	

Table 158: Manipulation/mobilisation versus usual care in low back pain without sciatica

			Quality asse	essment			No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation/mobilisation	Usual care	Relative (95% CI)	Absolute		
Pain (NRS	S 0-10) <4 mc	onths (Bet	ter indicated by l	ower values)								
	randomised trials			no serious indirectness	Serious⁵	none	37	35	-	MD 1.2 lower (2.26 to 0.14 lower)	⊕⊕OO LOW	CRITICAL
Pain (NRS	S 0-10) >4 mo	onths (Bet	ter indicated by l	ower values)								
	randomised trials			no serious indirectness	Serious ^b	none	37	35	-	MD 0.9 lower (1.98 lower to 0.18 higher)	⊕⊕OO LOW	CRITICAL
Function	(ODI 0-100) <	4 months	(Better indicated	by lower value	s)							
		- ,		no serious indirectness	Serious ^b	none	105	92	-	MD 6.43 lower (10.93 to 1.93 lower)	⊕OOO VERY LOW	CRITICAL
Function	(ODI 0-100) >	4 months	(Better indicated	d by lower value	s)							

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

1	randomised trials		no serious inconsistency	no serious indirectness	very serious ^b	none	37	35	-	MD 2.3 lower (9.14 lower to 4.54 higher)	⊕OOO VERY LOW	CRITICAL			
Respond	Responder criteria (>30% reduction pain) <4 months														
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	35/37 (94.6%)	20/35 (57.1%)	RR 1.66 (1.23 to 2.23)	377 more per 1000 (from 131 more to 703 more)	⊕⊕OO LOW	IMPORTANT			
Respond	der criteria (>	50% reduc	tion pain) <4 mo	nths											
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	28/37 (75.7%)	14/35 (40%)	RR 1.89 (1.21 to 2.95)	356 more per 1000 (from 84 more to 780 more)	⊕⊕OO LOW	IMPORTANT			
Respond	der criteria (>3	30% reduc	tion ODI) <4 mor	nths											
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	28/37 (75.7%)	17/35 (48.6%)	RR 1.56 (1.06 to 2.29)	272 more per 1000 (from 29 more to 627 more)	⊕⊕OO LOW	IMPORTANT			
Respond	der criteria (>	50% reduc	tion ODI) <4 mor	nths											
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	19/37 (51.4%)	14/35 (40%)	RR 1.28 (0.77 to 2.14)	112 more per 1000 (from 92 fewer to 456 more)	⊕⊕OO LOW	IMPORTANT			

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 159: Manipulation/mobilisation versus soft tissue techniques (massage) in low back pain without sciatica

			Quality ass	sessment			No of patients			Effect	Overlite v	I	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation/mobilisation versus massage	Control	Relative (95% CI)	Absolute	Quality	Importance	
Pain (VAS	Pain (VAS 0-10) <4 months (range of scores: 0-10; Better indicated by lower values)												

2	randomised trials	- ,	no serious inconsistency	no serious indirectness	no serious imprecision	none	110	81	-	MD 0.36 lower (0.98 lower to 0.26 higher)	⊕⊕OO LOW	CRITICAL		
Pain (VA	S 0-10) > 4 m	onths (rar	nge of scores: 0-	10; Better indica	ited by lower va	alues)								
1	randomised trials	-)	no serious inconsistency	no serious indirectness	Serious ^b	none	40	47	-	MD 0.59 lower (1.58 lower to 0.4 higher)	⊕OOO VERY LOW	CRITICAL		
Function	Function (RMDQ 0-24) <4 months (range of scores: 0-24; Better indicated by lower values)													
1	randomised trials	,	no serious inconsistency	no serious indirectness	Serious ^b	none	45	49	-	MD 1.38 lower (3.41 lower to 0.65 higher)	⊕OOO VERY LOW	CRITICAL		
Function	(RMDQ 0-24)	> 4 mont	ths (range of sco	res: 0-24; Better	indicated by lo	ower values)								
1	randomised trials	,	no serious inconsistency	no serious indirectness	Serious ^b	none	41	47	-	MD 1.77 lower (3.76 lower to 0.22 higher)	⊕OOO VERY LOW	CRITICAL		

^a Downgraded by one increment if the majority of the evidence was at high risk of bias, and downgraded by two increments if the majority of the evidence was at very high risk of bias ^b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

Table 160: Manipulation/mobilisation versus belts/corsets in low back pain without sciatica

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			Quality asse	essment			No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation/mobilisation versus belts/corsets	Control	Relative (95% CI)	Absolute	Quality	Importance
Pain (VAS	S 0-10) <4 mo	nths (rang	ge of scores: 0-10); Better indicate	ed by lower v	values)						
1		- 3		no serious indirectness	Serious ^b	none	65	25	-	MD 0.82 lower (2.07 lower to 0.43 higher)	⊕OOO VERY LOW	CRITICAL

a Downgraded by one increment if the majority of the evidence was at high risk of bias, and downgraded by two increments if the majority of the evidence was at very high risk of bias b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

Table 161: Manipulation/mobilisation versus exercise in low back pain with or without sciatica (mixed population)

			Quality asse	essment			No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation/mobilisation		Relative (95% CI)	Absolute	Quanty	importance
Pain severity (NRS, 0-10) < 4 months (range of scores: 0-10; Better indicated by lower values)												
		,		no serious indirectness	Serious ^b	none	13	11	-	MD 1.08 lower (2.76 lower to 0.6 higher)	⊕OOO VERY LOW	CRITICAL
Function ((RMDQ, 0-24)	< 4 month	ns (range of score	s: 0-24; Better in	dicated by lo	ower values)						
		,		no serious indirectness	Serious ^b	none	13	11	-	MD 3.21 lower (7.38 lower to 0.96 higher)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 162: Manipulation/mobilisation versus interferential therapy in low back pain with or without sciatica (mixed population)

			Quality as	sessment			No of patients			Effect	Our life	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation/mobilisation versus interferential therapy	Control	Relative (95% CI)	Absolute	Quality	Importanc
Quality of	f life (EQ-5D,	0-1) <4 n	nonths (range of	scores: 0-1; Be	tter indicated b	y higher values)						
		,			no serious imprecision	none	63	65	-	MD 0 higher (0.22 lower to 0.22 higher)	⊕⊕OO LOW	CRITICAL

				•		_						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	52	55	-	MD 0.05 lower (0.23 lower to 0.13 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life (SF-36-	General I	health 0-100) <4 r	nonths (range o	of scores: 0-10	0; Better indicated	d by higher values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	63	65	-	MD 0.38 lower (6.05 lower to 5.29 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life (SF-36	- Physical	I function 0-100)	<4 months (ran	ge of scores: 0	-100; Better indica	ated by higher values)					
1	randomised trials	,	no serious inconsistency	no serious indirectness	no serious imprecision	none	63	65	-	MD 4.64 higher (20.63 lower to 29.91 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life (SF-36	- Physical	I role limitation 0	-100) <4 months	s (range of sco	res: 0-100; Better	indicated by higher values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	63	65	-	MD 2.79 lower (16.97 lower to 11.39 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life (SF-36	- Bodily p	ain 0-100) <4 mo	nths (range of s	scores: 0-100; I	Better indicated b	y higher values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	63	65	-	MD 0.21 higher (7.61 lower to 8.03 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life (SF-36	- Vitality 0)-100) <4 months	(range of score	es: 0-100; Bette	er indicated by hig	her values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	63	65	-	MD 1.85 higher (4.73 lower to 8.43 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life (SF-36	- Social fu	unction 0-100) <4	months (range	of scores: 0-10	00; Better indicate	ed by higher values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	63	65	-	MD 3.05 higher (5.74 lower to 11.84 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life (SF-36	- Mental h	nealth 0-100) <4 m	nonths (range o	f scores: 0-100	; Better indicated	by higher values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	63	65	-	MD 2.35 higher (3.01 lower to	⊕⊕OO LOW	CRITICAL

7.71 higher)

Quality o	of life (SF-36 -	Mental h	ealth 0-100) > 4 r	months (range o	of scores: 0-10); Better indicated	l by higher values)					
		- ,		no serious indirectness	Serious ^b	none	52	55	-	MD 3.88 higher (2.86 lower to 10.62 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	of life (SF-36 -	Emotion	al role limitation	0-100) > 4 mont	ths (range of s	cores: 0-100; Bette	er indicated by higher values)					
		very serious ^a		no serious indirectness	no serious imprecision	none	52	55	-	MD 2.6 higher (11.98 lower to 17.18 higher)	⊕⊕OO LOW	CRITICAL
Pain (VA	S 0-10) < 4 m	onths (ra	nge of scores: 0-	10; Better indic	ated by lower	values)						
		very serious ^a		no serious indirectness	no serious imprecision	none	63	65	-	MD 0.15 higher (0.71 lower to 1.01 higher)	⊕⊕OO LOW	CRITICAL
Pain (VA	S 0-10) > 4 m	onths (ra	nge of scores: 0-	10; Better indic	ated by lower	values)						
		- ,		no serious indirectness	Serious ^b	none	52	55	-	MD 0.83 higher (0.19 lower to 1.85 higher)	⊕OOO VERY LOW	CRITICAL
Function	(RMDQ 0-24)) <4 mont	ths (range of sco	res: 0-24; Bette	r indicated by I	ower values)						
		,		no serious indirectness	Serious ^b	none	63	65	-	MD 0.97 lower (2.64 lower to 0.7 higher)	⊕OOO VERY LOW	CRITICAL
Function	(RMDQ 0-24)) > 4 mon	ths (range of sco	ores: 0-24; Bette	er indicated by	lower values)						
		very serious ^a		no serious indirectness	no serious imprecision	none	63	65	-	MD 0.19 higher (1.68 lower to 2.06 higher)	⊕⊕OO LOW	CRITICAL

^a Downgraded by one increment if the majority of the evidence was at high risk of bias, and downgraded by two increments if the majority of the evidence was at very high risk of bias ^b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs.

Table 163: Manipulation/mobilisation versus ultrasound therapy in low back pain without sciatica

	•	•			•	•						
			Quality ass	essment			No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation/mobilisation versus ultrasound therapy	Control	Relative (95% CI)	Absolute	Quanty	importance
Pain (VA	S 0-10) <4 mo	onths (ran	ge of scores: 0-1	0; Better indica	ted by lower	values)						
1		- ,	no serious inconsistency	no serious indirectness	Serious ^b	none	56	56	-	MD 1.65 higher (0.63 to 2.67 higher)	⊕OOO VERY LOW	CRITICAL
Pain (VA	S 0-10) > 4 m	onths (rar	nge of scores: 0-	10; Better indica	ited by lower	values)						
1		- ,	no serious inconsistency	no serious indirectness	Serious ^b	none	40	33	,	MD 1.51 higher (0.1 to 2.92 higher)	⊕000 VERY LOW	CRITICAL
Function	(ODI 0-100) <	4 months	(range of score	s: 0-100; Better	indicated by	lower values)						
1		- ,	no serious inconsistency	no serious indirectness	Serious ^b	none	56	56	-	MD 7.8 higher (2.41 to 13.19 higher)	⊕OOO VERY LOW	CRITICAL
Function	(ODI 0-100) >	4 month	s (range of score	es: 0-100; Better	indicated by	lower values)						
1			no serious inconsistency	no serious indirectness	Serious ^b	none	40	33	-	MD 5.2 higher (2.65 lower to 13.05 higher)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by one increment if the majority of the evidence was at high risk of bias, and downgraded by two increments if the majority of the evidence was at very high risk of bias ^b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

Table 164: Manipulation/mobilisation versus self-management in low back pain with or without sciatica (mixed population)

Quality assessment	No of patients	Effect	Quality	Importance
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No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation/mobilisation versus self- management	Control	Relative (95% CI)	Absolute			
Pain (VA	S 0-10) <4 m	onths (rang	ge of scores: 0-1	0; Better indica	ited by lower v	alues)							
2	trials				no serious imprecision	none	0	-	-	MD 0.18 lower (0.92 lower to 0.56 higher)	⊕⊕⊕⊕ HIGH	CRITICAL	
Function	Function (ODI 0-100) <4 months (range of scores: 0-100; Better indicated by lower values)												
1	trials			no serious indirectness	Serious ^a	none	39	38	-	MD 5.4 lower (10.32 to 0.48 lower)	⊕⊕⊕O MODERATE	CRITICAL	

^a Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

Table 165: Manipulation/mobilisation versus non-steroidal anti-inflammatories (NSAIDs) in low back pain without sciatica

			Quality as:	sessment			No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation/mobilisation		Relative (95% CI)	Absolute	Quanty	Importance
Pain seve	erity (VAS, 0-1	l0) < 4 mo	onths (range of sc	ores: 0-10; Bette	er indicated by	lower values)						
	randomised trials				no serious imprecision	none	58	57	-	MD 0.2 lower (0.89 lower to 0.49 higher)	⊕⊕⊕O MODERATE	CRITICAL
Function	(RMDQ, 0-24)) < 4 mont	ths (range of scor	es: 0-24; Better	indicated by lo	wer values)						
1	randomised trials				no serious imprecision	none	58	57	-	MD 0.4 lower (2.06 lower to 1.26 higher)	⊕⊕⊕O MODERATE	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 166: Manipulation/mobilisation versus non-steroidal anti-inflammatories (NSAIDs) in low back pain with or without sciatica (mixed population)

			Quality as:	sessment			No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation/mobilisation versus NSAIDs	Control	Relative (95% CI)	Absolute	quanty	Importance
Pain (VA	S 0-10) <4 mc	onths (ran	ge of scores: 0-1	0; Better indica	ted by lower v	alues)						
	randomised trials		no serious inconsistency		no serious imprecision	none	56	40	-	MD 0.80 lower (1.66 lower to 0.06 higher)	⊕⊕⊕O MODERATE	CRITICAL
Function	(RMDQ 0-24)	<4 mont	hs (range of scor	res: 0-10; Better	indicated by l	ower values)						
	randomised trials				no serious imprecision	none	94	77	-	MD 1.96 lower (3.29 to 0.62 lower)	⊕⊕⊕O MODERATE	CRITICAL

^a Downgraded by one increment if the majority of the evidence was at high risk of bias, and downgraded by two increments if the majority of the evidence was at very high risk of bias

Table 167: Manipulation/mobilisation versus combination of inteventions (exercise + education) in low back pain with or without sciatica (mixed population)

	populat	,											
			Quality asse	essment			No of patient	s		Effect			
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation/mobilisation		Relative (95% CI)	Absolute	Quality	Importance	
Pain seve	Pain severity (NRS, 0-10) < 4 months (range of scores: 0-10; Better indicated by lower values)												
		- ,		no serious indirectness	Serious ^b	none	13	10	-	MD 1.78 lower (3.22 to 0.34 lower)	⊕000 VERY LOW	CRITICAL	
Function	Function (RMDQ, 0-24) < 4 months (range of scores: 0-24; Better indicated by lower values)												

		,	no serious inconsistency	no serious indirectness	Serious ^b	none	13	10	1	MD 4.85 lower (8.88 to 0.82 lower)	⊕OOO VERY LOW	CRITICAL
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a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

J.8.4 Mixed modality manual therapy

Table 168: Mixed modality manual therapy versus usual care in low back pain without sciatica

			Quality asses	ssment		No of patients			Effect	Quality	Importance			
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Mixed modality manual therapy	UC	Relative (95% CI)	Absolute	Quality	Importance		
Pain sever	Pain severity (Melzak pain score, 0-5) < 4 months (range of scores: 0-5; Better indicated by lower values)													
1	randomised trials	very serious ^a		no serious indirectness	Serious ^b	none	8	10	-	MD 0.9 lower (1.4 to 0.39 lower)	⊕OOO VERY LOW	CRITICAL		

a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 169: Mixed modality manual therapy versus sham in low back pain without sciatica

			Quality assess	ment		No of patients		Effect	t	Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Mixed modality manual therapy	Sham	Relative (95% CI)	Absolute	,	
Responder criteria <4 months												
1		no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^a	none	-	-	RR 1.38 (1.16 to 1.64)		⊕⊕⊕O MODERATE	CRITICAL

a Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

Table 170: Mixed modality manual therapy versus sham in low back pain with or without sciatica (mixed population)

		,					itiioat seiatica (iiii			,		
			Quality asses	sment			No of patients	•		Effect	6 111	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Mixed modality manual therapy versus sham	Control	Relative (95% CI)	Absolute	Quality	Importance
Pain (NRS	6 0-10) <4 mo	nths (range o	f scores: 0-10; Be	tter indicated by	lower value	s)						
1	randomised trials		no serious inconsistency	no serious indirectness	Seriousª	none	15	14	ı	MD 0.28 higher (0.46 lower to 1.02 higher)		CRITICAL
Pain (NRS	6 0-10) > 4 mo	nths (range o	of scores: 0-10; Bo	etter indicated b	y lower value	es)						
1	randomised trials		no serious inconsistency		very serious ^a	none	15	14	-	MD 0.32 lower (1.24 lower to 0.6 higher)	⊕⊕OO LOW	CRITICAL
Function	(ODI change	score 0-100) •	<4 months (Better	indicated by lov	wer values)							
1	randomised trials		no serious inconsistency	no serious indirectness	Seriousª	none	15	14	ı	MD 2.03 lower (8.54 lower to 4.48 higher)	⊕⊕OO LOW	CRITICAL
Function	(ODI change	score 0-100)	>4 months (Better	indicated by lov	ver values)							
1	randomised trials	Serious ^b	no serious inconsistency	no serious indirectness	Seriousª	none	15	14	-	MD 1.26 lower (8.44 lower to 5.92 higher)	⊕⊕OO LOW	CRITICAL

Table 171: Mixed modality manual therapy versus manipulation/mobilisation in low back pain without sciatica

Quality assessment	No of patients	Effect	Quality	Importance

^a Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs ^b Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Mixed modality manual therapy versus manipulation/mobilisation	Control	Relative (95% CI)	Absolute		
Pain (VA	S 0-10) <4 m	onths (rai	nge of scores: 0-	10; Better indic	ated by lower	values)						
1		,	no serious inconsistency	no serious indirectness	Serious ^b	none	48	45	-	MD 0.54 lower (1.89 lower to 0.81 higher)	⊕OOO VERY LOW	CRITICAL
Pain (VA	S 0-10) > 4 m	onths (ra	inge of scores: 0	-10; Better indic	cated by lower	values)						
1		- ,	no serious inconsistency		no serious imprecision	none	49	40	-	MD 0.16 lower (1.1 lower to 0.78 higher)	⊕⊕OO LOW	CRITICAL
Function	(RMDQ 0-24) <4 mon	ths (range of sco	res: 0-24; Bette	er indicated by	lower values)						
1		,	no serious inconsistency	no serious indirectness	Serious ^b	none	48	45	-	MD 0.69 lower (2.48 lower to 1.1 higher)	⊕OOO VERY LOW	CRITICAL
Function (RMDQ 0-24) > 4 months (range of scores: 0-24; Better indicated by lower values)												
1		,	no serious inconsistency		no serious imprecision	none	48	41	-	MD 0.27 higher (1.48 lower to 2.02 higher)	⊕⊕OO LOW	CRITICAL

^a Downgraded by one increment if the majority of the evidence was at high risk of bias, and downgraded by two increments if the majority of the evidence was at very high risk of bias ^b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

Table 172: Mixed modality manual therapy versus soft tissue techniques (massage) in low back pain without sciatica

		-	Quality asse	ssment			No of patients			Effect	Ovelite		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Mixed modality manual therapy versus massage	Control	Relative (95% CI)	Absolute	Quality	Importance	
Pain (VAS	Pain (VAS 0-10) <4 months (range of scores: 0-10; Better indicated by lower values)												

1	randomised trials	,	no serious inconsistency	no serious indirectness	Serious ^b	none	48	49	-	MD 0.74 lower (1.38 to 0.1 lower)	⊕OOO VERY LOW	CRITICAL			
Pain (VAS	Pain (VAS 0-10) > 4 months (range of scores: 0-10; Better indicated by lower values)														
1	randomised trials	,	no serious inconsistency	no serious indirectness	Serious ^b	none	49	47	1	MD 0.75 lower (1.61 lower to 0.11 higher)	⊕OOO VERY LOW	CRITICAL			
Function	Function (RMDQ 0-24) > 4 months (range of scores: 0-24; Better indicated by lower values)														
1	randomised trials	- ,	no serious inconsistency	no serious indirectness	Serious ^b	none	48	49	-	MD 1.5 lower (3.18 lower to 0.18 higher)	⊕OOO VERY LOW	CRITICAL			
Function	(RMDQ 0-24)	<4 months	(range of scores:	0-24; Better indi	cated by low	er values)									
1	randomised trials	- ,	no serious inconsistency	no serious indirectness	Serious ^b	none	48	49	-	MD 2.07 lower (3.86 to 0.28 lower)	⊕OOO VERY LOW	CRITICAL			

^a Downgraded by one increment if the majority of the evidence was at high risk of bias, and downgraded by two increments if the majority of the evidence was at very high risk of bias ^b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

Table 173: Mixed modality manual therapy versus traction in low back pain without sciatica

			Quality asse	ssment			No of patients			Effect	Qualita	I
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Mixed modality manual therapy versus traction	Control	Relative (95% CI)	Absolute	Quality	Importance
Pain (VAS	0-10) <4 mon	ths (range	of scores: 0-10; Be	etter indicated by	lower value	s)						
1	randomised trials	- ,		no serious indirectness	Serious ^b	none	30	30	-	MD 1 lower (1.66 to 0.34 lower)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by one increment if the majority of the evidence was at high risk of bias, and downgraded by two increments if the majority of the evidence was at very high risk of bias

Table 174: Mixed modality manual therapy versus exercise (biomechanical) in low back pain without sciatica

			Quality asse	essment			No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Mixed modality manual therapy versus biomechanical exercise		Relative (95% CI)	Absolute	Quanty	importance
Pain (Mel	zak pain scale	e 0-5) <4 r	months (range of	scores: 0-5; Bett	er indicated	by lower values)						
1		- 3		no serious indirectness	Serious ^b	none	8	10	-	MD 0.5 lower (1.03 lower to 0.03 higher)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by one increment if the majority of the evidence was at high risk of bias, and downgraded by two increments if the majority of the evidence was at very high risk of bias ^b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

Combination interventions – manual therapy adjunct J.8.4.1

Low back pain with sciatica J.8.4.2

Table 175: Manual therapy (manipulation) plus self-management (education) plus exercise (aerobic) compared with self-management (education) plus exercise (aerobic plus McKenzie)

			Quality asse	essment			No of pa	itients		Effect	O lite.	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation + education + exercise (aerobic)		Relative (95% CI)	Absolute	Quality	Importance
Pain (VAS	change sco	re) - <4 m	onths (measured	I with: VΔS: ran	ne of scores	· 0-10· Retter ind	icated by lower values					

^b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

		,		no serious indirectness	Serious ^b	none	10	15	-	MD 0.9 lower (2.49 lower to 0.69 higher)	⊕OOO VERY LOW	CRITICAL
Function	(ODI, 0-100)	<4 month	s (range of score	s: 0-100; Better	indicated by	/ lower values)						
		- 3		no serious indirectness	Serious ^b	none	10	15	-	MD 2.86 higher (4.44 lower to 10.16 higher)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 176: Manual therapy (soft tissue techniques – muscle energy technique) plus biomechanical exercise (McKenzie) plus self management (unsupervised exercise) versus biomechanical exercise (McKenzie) plus self management (unsupervised exercise

			Quality asse	ssment			No of pat	ients		Effect	Ovelite	
No of studies	Design Inconsistency Ingirectnes				Imprecision	Other considerations	Manual + ex + self manag	Ex + self manag	Relative (95% CI)	Absolute	Quanty	Importance
Pain sever	rity (VAS, 0-10)) < 4 montl	hs (range of scores	: 0-10; Better ind	icated by low	ver values)						
	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	20	20	-	MD 0.1 lower (0.72 lower to 0.52 higher)	⊕OOO VERY LOW	CRITICAL
Function (ODI, 0-100) < 4	months (range of scores: 0-	100; Better indica	ited by lower	values)						
	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	20	20	-	MD 0.86 lower (4.12 lower to 2.4 higher)	⊕⊕OO LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

b Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

NICE.

Table 177: Manual therapy (soft tissue techniques – muscle energy technique) plus biomechanical exercise (McKenzie) plus self management (unsupervised exercise) versus standard treatment (massage + laser + TENS) plus self management

	(anoupe		xe. 6.56, 16.543	standard tre	u e	assage : lase.	· 12110/ piac	s sen management				
			Quality asse	essment			N	lo of patients		Effect		
No of studies	I DESIGN I INCONSISTANCY I INGIFECTIONS IMPRECISIONI					Other considerations	Manual + ex + self manag	Std treatment (massage + TENS + laser) + self manag	Relative (95% CI)	Absolute	Quality	Importance
Pain seve	erity (VAS, 0-1	0) < 4 mo	nths (range of sco	ores: 0-10; Bette	r indicated b	y lower values)						
1	randomised trials			no serious indirectness	very serious ^b	none	20	20	-	MD 3.29 lower (4.03 to 2.55 lower)	⊕OOO VERY LOW	CRITICAL
Function (ODI, 0-100) < 4 months (range of scores: 0-100; Better indicated by lower values)												
1	randomised trials			no serious indirectness	very serious ^b	none	20	20	1	MD 19.07 lower (24.26 to 13.88 lower)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Low back pain without sciatica J.8.4.3

Table 178: Manual therapy (soft tissue techniques - massage) plus self-management (exercise prescription) versus Postural therapy (Alexander technique -6 lessons)

	teemin	•	Quality ass	sessment			No of patients			Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Massage + self-management (exercise prescription) versus Alexander technique (6 lessons)	Control	Relative (95% CI)	Absolute	Quality	Importance
Qualty of	life (SF-36 p	hysical c	omponent summ	nary) >4months	(follow-up 1 y	ears; range of sc	ores: 0-100; Better indicated by	higher	values)			

-												
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	56	58	-	MD 1.59 higher (7.27 lower to 10.45 higher)	⊕⊕⊕O MODERATE	CRITICAL
Qualty o	of life (SF-36 n	nental co	mponent summa	ry) >4 months	(follow-up 1 ye	ears; range of sco	res: 0-100; Better indicated by h	nigher va	alues)			
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	56	58	-	MD 1.37 lower (9.31 lower to 6.57 higher)	⊕⊕⊕O MODERATE	CRITICAL
Pain (Vo	on Korff pain s	scale) >4r	months (follow-u	p 1 years; rang	e of scores: 0-	10; Better indicat	ed by lower values)					
1	randomised trials	Seriousª	no serious inconsistency		no serious imprecision	none	56	58	-	MD 0.22 lower (1.19 lower to 0.75 higher)	⊕⊕⊕O MODERATE	CRITICAL
Functio	n (RMDQ, 0-24	4) >4 mor	nths (follow-up 1	years; range of	scores: 0-24;	Better indicated b	oy lower values)					
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	58	56	-	MD 0.93 lower (2.84 lower to 0.98 higher)	⊕⊕OO LOW	CRITICAL
Healthc	are utilisation	(primary	care contacts) >	4months (follo	w-up 1 years;	Better indicated b	y lower values)					
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	56	58	-	MD 0.16 lower (0.47 lower to 0.15 higher)	⊕⊕OO LOW	IMPORTANT
Healthc	are utilisation	(prescrip	otions) >4months	(follow-up 1 ye	ears; Better in	dicated by lower v	values)					
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	56	58	-	MD 0.04 lower (0.55 lower to 0.47 higher)	⊕⊕⊕O MODERATE	IMPORTANT

^a Downgraded by one increment if the majority of the evidence was at high risk of bias ^b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs.

Table 179: Manual therapy (soft tissue techniques - massage) plus self-management (exercise prescription) versus Postural therapy (Alexander technique -(24 lessons)

		10.0 (163301137									
			Quality as:	sessment			No of patients			Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Massage + self-management (exercise prescription) versus Alexander technique (24 lessons)	Control	Relative (95% CI)	Absolute	Quality	Importance
Qualty o	f life (SF-36 p	hysical c	omponent summ	nary, 0-100) >4 :	months (follow	· ∕-up 1 years; ranç	ge of scores: 0-100; Better indic	ated by	higher va	ılues)		
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	56	61	-	MD 8.47 lower (17.15 lower to 0.21 higher)	⊕⊕OO LOW	CRITICAL
Qualty of life (SF-36 mental component summary, 0-100) >4 months (follow-up 1 years; range of scores: 0-100; Better indicated by higher values)												
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	56	61	-	MD 1.01 lower (9.32 lower to 7.3 higher)	⊕⊕⊕O MODERATE	CRITICAL
Pain (Vo	n Korff pain s	scale) >4	months (follow-ເ	ıp 1 years; ranç	ge of scores: 0	-10; Better indica	ted by lower values)	•				
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	serious ²	none	57	61	-	MD 0.68 higher (0.28 lower to 1.64 higher)	⊕⊕OO LOW	CRITICAL
Function	(RMDQ, 0-24	4) >4 mon	ths (follow-up 1	years; range of	scores: 0-24;	Better indicated	by lower values)					
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	56	61	-	MD 1.77 higher (0.11 lower to 3.65 higher)	⊕⊕OO LOW	CRITICAL
Healthca	re utilisation	(primary	care contacts) >	4 months (follo	ow-up 1 years;	Better indicated	by lower values)					
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	56	61	-	MD 0.12 lower (0.42 lower to 0.18 higher)	⊕⊕⊕O MODERATE	IMPORTANT
Healthca	re utilisation	(prescrip	otions) >4 months	s (follow-up 1 y	ears; Better in	dicated by lower	values)					

	randomised trials		no serious indirectness	Serious⁵	none	87	6	-	MD 0.49 lower (1.14 lower to 0.16 higher)	⊕⊕OO LOW	IMPORTANT
a Downgra	aded by one i	if the majority of the	ne evidence was	at high risk of	bias.						

^b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs.

Table 180: Manual therapy (manipulation) plus exercise (McKenzie) compared with exercise (biomechanical - core stability)

			Quality asse	ssment			No of patients	S		Effect			
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation + exercise (McKenzie)	core stability	Relative (95% CI)		Quality	Importance	
Function ((ODI, 0-100) <4	months (follow-up 4 weeks	; measured with:	ODI; range	of scores: 0-100; B	etter indicated by lowe	r values)					
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	40	46	-	MD 4 lower (11.34 lower to 3.34 higher)	⊕⊕OO LOW	CRITICAL	
Function (inction (ODI, 0-100) >4 months (follow-up 12 months; measured with: ODI; range of scores: 0-100; Better indicated by lower values)												
1	randomised trials	Serious ^a		no serious indirectness	Serious ^b	none	40	46	-	MD 3.7 lower (11.46 lower to 4.06 higher)	⊕⊕OO LOW	CRITICAL	

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

Table 181: Manual therapy (manipulation) plus exercise (McKenzie) compared with exercise (biomechanical – stretching)

			Quality asse	ssment			No of patients	s		Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation + exercise (McKenzie) +	etrotching	Relative (95% CI)	Absolute	Quality	Importance
Function ((ODI, 0-100) <4	4 months (follow-up 4 weeks	; measured with	: ODI; range	of scores: 0-100; E	Better indicated by lowe	r values)				

1	randomised trials			no serious indirectness	Serious ^b	none	40	37	-	MD 2.7 lower (10.29 lower to 4.89 higher)	⊕⊕OO LOW	CRITICAL		
Function	Function (ODI, 0-100) >4 months (follow-up 12 months; measured with: ODI; range of scores: 0-100; Better indicated by lower values)													
1	randomised trials			no serious indirectness	Serious ^b	none	40	37	-	MD 2 higher (5.46 lower to 9.46 higher)	⊕⊕OO LOW	CRITICAL		

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 182: Manual therapy (manipulation) + exercise (aerobic) compared to exercise (aerobic)

			Quality asse	essment			No of patie	nts		Effect	Ovalita		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation + exercise (aerobic)	exercise (aerobic)	Relative (95% CI)	Absolute	Quality	Importance	
Pain (VAS	ain (VAS, 0-10) <4 months (follow-up 6 weeks; measured with: VAS; range of scores: 0-10; Better indicated by lower values)												
1		- 3	no serious inconsistency	no serious indirectness	Serious ^b	none	15	18	-	MD 0.9 lower (2.68 lower to 0.88 higher)	⊕OOO VERY LOW	CRITICAL	
Function	Function (Quebec back pain disability scale) - <4 months (follow-up 6 weeks; range of scores: 20-100; Better indicated by lower values)												
1		,	no serious inconsistency	no serious indirectness	Serious ^b	none	15	18	-	MD 10.7 lower (23.45 lower to 2.05 higher)	⊕OOO VERY LOW	CRITICAL	

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

Table 183: Manual therapy (manipulation) plus exercise (aerobic) compared with exercise (biomechanical)

|--|

b Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation + exercise (aerob)	exercise (biomech)	Relative (95% CI)	Absolute		
Pain (VAS	6 0-10) - <4 mc	onths (follo	ow-up 6 weeks; m	easured with: V	AS; range of	scores: 0-10; Bette	er indicated by lower	values)				
1		- ,	no serious inconsistency	no serious indirectness	very serious ^b	none	15	18	,	MD 0.07 lower (1.64 lower to 1.5 higher)	⊕OOO VERY LOW	CRITICAL
Function	(Quebec back	pain disa	bility scale 0-100)	- <4 months (fol	low-up 6 wee	eks; range of score	es: 20-100; Better ind	icated by low	er values)		
1		- ,	no serious inconsistency	no serious indirectness	very serious ^b	none	15	18	-	MD 1.48 lower (14.26 lower to 11.3 higher)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

Table 184: Manual therapy (manipulation) plus exercise (biomechanical) compared with exercise (aerobic)

			Quality asse	essment			No of patier	nts		Effect	Ovality	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation + exercise (biomech)	exercise (aerobic)	Relative (95% CI)	Absolute	Quanty	Importance
Pain (VAS	6 0-10) - <4 mc	onths (follo	ow-up 6 weeks; m	easured with: VA	AS; range of	scores: 0-10; Bett	er indicated by lower	values)				
1	randomised trials	,		no serious indirectness	Serious ^b	none	21	18	,	MD 1.89 lower (3.4 to 0.38 lower)	⊕OOO VERY LOW	CRITICAL
Function (Quebec back pain disability scale 0-100) - <4 months (follow-up 6 weeks; range of scores: 20-100; Better indicated by lower values)												
1	randomised trials	,		no serious indirectness	Serious ^b	none	21	18	-	MD 11.45 lower (23.54 lower to 0.64 higher)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

b Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

Table 185: Manual therapy (manipulation) plus exercise (biomechanical) compared with exercise (biomechanical)

	Quality assessment No of patients Effect Qu											Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation + exercise (biomech)	exercise (biomech)	Relative (95% CI)	Absolute	Quality	Importance	
Pain (VAS	Pain (VAS 0-10) - <4 months (follow-up 6 weeks; measured with: VAS; range of scores: 0-10; Better indicated by lower values)												
	randomised trials	,		no serious indirectness	Serious ^b	none	21	18	-	MD 1.06 lower (2.32 lower to 0.2 higher)	⊕OOO VERY LOW	CRITICAL	
Function (Quebec back pain disability scale 0-100) - <4 months (follow-up 6 weeks; range of scores: 20-100; Better indicated by lower values)													
1	randomised trials	,			very serious ^b	none	21	18	-	MD 2.23 lower (14.36 lower to 9.9 higher)	⊕OOO VERY LOW	CRITICAL	

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 186: Manual therapy (manipulation) plus exercise (biomechanical) compared with Manual therapy (manipulation) plus exercise (aerobic)

			Quality asse	essment			No of p	atients		Effect	Ovalite		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation + exercise (biomech)	manipulation +	Relative (95% CI)	Absolute	Quality li	mportance	
Pain (VAS	Pain (VAS 0-10) - <4 months (follow-up 6 weeks; measured with: VAS; range of scores: 0-10; Better indicated by lower values)												
1		- ,		no serious indirectness	Serious ^b	none	21	15	-	MD 0.99 lower (2.52 lower to 0.54	⊕000 VERY	CRITICAL	

b Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

										higher)	LOW	
Function	(Quebec bac	k pain dis	ability scale 0-10	0) - <4 months (follow-up 6 v	veeks; range of so	cores: 20-100; Better	indicated by lower	values)			
1	randomised trials	,		no serious indirectness	very serious ^b	none	21	15	-	MD 0.75 lower (12.99 lower to 11.49 higher)	⊕000 VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

Table 187: Manual therapy (mixed modality - manipulation plus soft tissue techniques - massage) compared with sham

			Quality asse	essment			No of patient	s		Effect	Quality		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation + massage	sham	Relative (95% CI)	Absolute	Quanty	Importance	
Pain (Pain	ain (Pain disability index) - <4 months (follow-up 3 weeks; range of scores: 0-70; Better indicated by lower values)												
		no serious risk of bias		no serious indirectness	no serious imprecision	none	54	52	1	MD 0.6 lower (4.26 lower to 3.06 higher)	⊕⊕⊕⊕ HIGH	CRITICAL	
Function	nction (RMDQ, 0-24) <4 months (follow-up 3 weeks; measured with: RMDQ; range of scores: 0-24; Better indicated by lower values)												
	randomised trials	no serious risk of bias		no serious indirectness	Serious ^a	none	54	52	-	MD 0.5 higher (0.74 lower to 1.74 higher)	⊕⊕⊕O MODERATE	CRITICAL	

^a Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

J.8.4.4 Overall: low back pain with/without sciatica

Table 188: Manual therapy plus self-management (home exercise) compared with self-management (home exercise) plus exercise

			Quality as	sessment			No of patients Effect				Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manual therapy + home exercise	home exercise + exercise	Relative (95% CI)	Absolute	Quality	importance
Pain (0-10	0 VAS conve	rted to 0-1	10) <4 months (Be	tter indicated by	lower values)							
1	randomised trials		no serious inconsistency		no serious imprecision	none	21	27	-	MD 1.7 higher (0.55 to 2.85 higher)	⊕⊕⊕O MODERATE	CRITICAL
Pain (0-10	0 VAS conve	rted to 0-1	I0) >4 months (Be	tter indicated by	lower values)							
1	randomised trials		no serious inconsistency		no serious imprecision	none	22	27	-	MD 1.4 higher (0.26 to 2.54 higher)	⊕⊕⊕O MODERATE	CRITICAL
Function	(ODI, 0-100) <	4 months	(Better indicated	by lower values								
1	randomised trials		no serious inconsistency		no serious imprecision	none	21	27	-	MD 12 higher (4.5 to 19.5 higher)	⊕⊕⊕O MODERATE	CRITICAL
Function	(ODI, 0-100) >	4 months	(Better indicated	by lower values)								
1	randomised trials		no serious inconsistency		no serious imprecision	none	21	27	-	MD 9 higher (1.19 to 16.81 higher)	⊕⊕⊕O MODERATE	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 189: Manual therapy (traction) plus infra-red plus exercise (biomechanical – stretch) compared with infra-red plus exercise (biomechanical – stretch)

			Quality as	sessment			No of pat	tients		Effect	Ovolity	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Traction + infra-red + stretch	infra-red + stretch	Relative (95% CI)	Absolute	Quanty	importance
Pain (NR	S 0-10) - <4 m	onths (Be	tter indicated by I	lower values)								
1	randomised trials	very seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	34	37	-	MD 0.3 lower (0.91 lower to 0.31 higher)	⊕OOO VERY LOW	CRITICAL
Pain (NR	S 0-10) - >4 m	onths (Be	tter indicated by I	lower values)								
1	randomised trials	very seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	32	35	-	MD 0.9 lower (1.45 to 0.35 lower)	⊕⊕OO LOW	CRITICAL
Function	(ODI, 0-100) <	<4 months	(Better indicated	by lower values	s)							
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	34	37	-	MD 1.6 lower (3.11 to 0.09 lower)	⊕⊕OO LOW	CRITICAL
Function	(ODI, 0-100) >	4 months	(Better indicated	by lower values	s)							
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	32	35	-	MD 3.3 lower (4.66 to 1.94 lower)	⊕⊕OO LOW	CRITICAL
Healthca	re utilisation (medicatio	on use) <4 months	S								
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	8/34 (23.5%)	11/37 (29.7%)	RR 0.79 (0.36 to 1.73)	62 fewer per 1000 (from 190 fewer to 217 more)	⊕OOO VERY LOW	IMPORTANT
Healthcar	re utilisation (medicatio	on use) >4 months	3								
1	randomised trials	very seriousª	no serious inconsistency	no serious indirectness	very serious ^b	none	5/33 (15.2%)	8/35 (22.9%)	RR 0.66 (0.24 to 1.82)	78 fewer per 1000 (from 174 fewer to 187 more)	⊕OOO VERY LOW	IMPORTANT

a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 190: Manual therapy (manipulation) plus electrotherapy (interferential) compared with electrotherapy (interferential)

			Quality as	sessment			No of pati	ents		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation + interferential	interferential	Relative (95% CI)	Absolute	Quanty	importance
Quality of	life (EQ-5D)	- <4 mont	hs (Better indicate	ed by lower valu	es)							
	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	66	65	1	MD 0.01 lower (0.15 lower to 0.13 higher)		CRITICAL
Quality of	life (EQ-5D)	- >4 mont	hs (Better indicate	ed by lower valu	es)							
	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	51	55	-	MD 0.05 higher (0.06 lower to 0.16 higher)		CRITICAL
Quality of	life (SF-36 P	hysical fu	nctioning, 0-100)	<4 months (Bett	er indicated by	lower values)						
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	very serious ^b	none	66	65	-	MD 3.69 higher (3.56 lower to 10.94 higher)	⊕000 VERY LOW	CRITICAL
Quality of	life (SF-36 P	hysical fu	nctioning, 0-100)	>4 months (Bett	er indicated by	lower values)						
	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	51	55	-	MD 9.69 higher (0.32 to 19.06 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality of life (SF-36 Role physical, 0-100) <4 months (Better indicated by lower values)												
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	very serious ^b	none	66	65	-	MD 1.36 lower (15.64 lower to 12.92 higher)	⊕000 VERY LOW	CRITICAL
Quality of	life (SF-36 R	ole physic	cal, 0-100) >4 mor	nths (Better indic	cated by lower v	values)						

		1	1		1	1						1
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	51	55	ı	MD 11.4 higher (6.1 lower to 28.9 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	f life (SF-36 B	odily pair	n, 0-100) <4 mont	hs (Better indica	ited by lower va	lues)						
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	66	65	ı	MD 0.48 lower (8.33 lower to 7.37 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	life (SF-36 B	odily pair	n, 0-100) >4 mont	hs (Better indica	ited by lower va	lues)						
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	51	55	-	MD 6 higher (3.8 lower to 15.8 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	life (SF-36 G	eneral he	alth, 0-100) <4 m	onths (Better inc	dicated by lower	r values)						
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	very serious ^b	none	66	65	-	MD 1.89 higher (3.87 lower to 7.65 higher)	⊕000 VERY LOW	CRITICAL
Quality of	Flife (SF-36 G	eneral he	alth, 0-100) >4 m	onths (Better inc	dicated by lower	r values)						
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	51	55	-	MD 3.43 higher (4.21 lower to 11.07 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	f life (SF-36 V	itality, 0-1	00) <4 months (E	Better indicated	by lower values)				, , , , , , , , , , , , , , , , , , ,		
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	66	65	-	MD 0.89 higher (5.72 lower to 7.5 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	life (SF-36 V	itality, 0-1	00) >4 months (E	Better indicated	by lower values)						
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	51	55	-	MD 7 higher (0.89 lower to 14.89 higher)	⊕⊕OO LOW	CRITICAL
Quality of	life (SF-36 S	ocial fund	ctioning, 0-100) <	4 months (Bette	r indicated by lo	ower values)						
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	66	65	-	MD 2.88 higher (5.96 lower to 11.72 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	f life (SF-36 S	ocial fund	ctioning, 0-100) >	4 months (Bette	r indicated by lo	ower values)						

	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	51	55	-	MD 8.1 higher (5.44 lower to 21.64 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	f life (SF-36 R	ole emoti	onal, 0-100) <4 m	onths (Better inc	dicated by lowe	r values)						
I	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	66	65	1	MD 4.02 higher (10.94 lower to 18.98 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	life (SF-36 R	ole emoti	onal, 0-100) >4 m	onths (Better inc	dicated by lowe	r values)						
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	51	55	-	MD 10.8 higher (4.34 lower to 25.94 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	life (SF-36 M	lental hea	lth, 0-100) <4 mor	nths (Better indi	cated by lower	values)						
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	66	65	-	MD 4.81 higher (0.78 lower to 10.4 higher)	⊕⊕OO LOW	CRITICAL
Quality of	f life (SF-36 M	lental hea	lth, 0-100) >4 mor	nths (Better indi	cated by lower	values)						
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	51	55	-	MD 9.46 higher (2.53 to 16.39 higher)	⊕⊕⊕O MODERATE	CRITICAL
Pain (0-10	00 VAS conve	erted to 0-	10) <4 months (B	etter indicated b	y lower values)							
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	66	65	-	MD 0.33 lower (1.2 lower to 0.54 higher)	⊕⊕⊕O MODERATE	CRITICAL
Pain (0-10	00 VAS conve	erted to 0-	10) >4 months (B	etter indicated b	y lower values)							
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	51	55	1	MD 0.08 higher (0.97 lower to 1.13 higher)		CRITICAL
Pain seve	erity (McGill P	ain Rating	g Index, range no	t stated) - <4 mo	nths (Better inc	dicated by lower v	alues)					
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	66	65	-	MD 0.77 lower (4.41 lower to 2.87 higher)	⊕⊕⊕O MODERATE	CRITICAL
Pain seve	erity (McGill P	ain Rating	g Index, range no	t stated) - >4 mo	enths (Better inc	dicated by lower v	alues)					

1	randomised trials		no serious inconsistency		no serious imprecision	none	51	55	-	MD 0.9 lower (5.21 lower to 3.41 higher)	⊕⊕⊕O MODERATE	CRITICAL	
Function	Function (RMDQ, 0-24) <4 months (Better indicated by lower values)												
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	66	65	-	MD 1.09 lower (2.75 lower to 0.57 higher)		CRITICAL	
Function	Function (RMDQ, 0-24) >4 months (Better indicated by lower values)												
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	51	55	-	MD 1.6 lower (3.51 lower to 0.31 higher)	⊕⊕OO LOW	CRITICAL	

Table 191: Manual therapy (manipulation) plus exercise (biomechanical – core stability) compared with exercise (biomechanical – core stability)

			Quality as:	sessment			No of patie	ents		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation + exercise (strength)	exercise (strength)	Relative (95% CI)	Absolute		
Medicatio	on use - >4 m	onths										
		- ,		no serious indirectness	Serious ^b	none	19/52 (36.5%)	60%	RR 0.61 (0.39 to 0.94)	234 fewer per 1000 (from 36 fewer to 366 fewer)	⊕OOO VERY LOW	IMPORTANT
Function	(ODI 0-100) >	4 months	(Better indicated	d by lower value	s)							
		very seriousª			no serious imprecision	none	52	40	-	MD 10.3 higher (4.3 to 16.3 higher)	⊕⊕OO LOW	CRITICAL

^a Downgraded by 1 increment if the majority of evidence was at high risk of bias, and downgraded by 2 increments if the majority of evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID, and downgraded by 2 increments if the confidence interval crossed both MIDs

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 192: Manual therapy (manipulation) plus exercise (biomechanical - strength) compared with pharmacological (NSAID) plus exercise (biomechanical - strength)

			Quality asse	essment			No of pat	ients		Effect	Quality	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation + exercise (strength)	NSAID + exercise (strength)	Relative (95% CI)	Absolute	Quanty	Importance
Pain (11-b	oox scale 0-10)) - <4 mo	nths (Better indica	ated by lower val	ues)							
1		,		no serious indirectness	Serious ^b	none	56	40	-	MD 0.8 lower (1.66 lower to 0.06 higher)	⊕OOO VERY LOW	CRITICAL
Function	(RMDQ, 0-24)	<4 month	ns (range of score	s: 0-24; Better in	dicated by lo	ower values)						
1		,		no serious indirectness	Serious ^b	none	56	40		MD 5.8 lower (12.77 lower to 1.17 higher)		CRITICAL

a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 193: Manual therapy (manipulation) plus exercise (biomechanical - stretch) compared with pharmacological (NSAID) plus exercise (biomechanical - strength)

			Quality asse	essment			No of pat	ients		Effect	Ouglitu	lm no uto u o o
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation + exercise (stretch)	NSAID + exercise (strength)	Relative (95% Absolute CI)		Quanty	Importance
Pain (11-l	oox scale 0-10)) - <4 moi	nths (Better indica	ated by lower val	lues)							
1	randomised trials	very serious ^a		no serious indirectness	Serious ^b	none	36	40	-	MD 0.2 lower (1.21 lower to 0.81 higher)	⊕OOO VERY LOW	CRITICAL

Downgraded by 1 increment if the confidence interval crossed one wild of downgraded by 2 increments if the confidence interval crossed both wilds

Function	(RMDQ, 0-24)	<4 month	ns (Better indicate	d by lower value	s)					
1		- ,		no serious indirectness	Serious ^b	none	36	40	MD 2.5 lower (10.18 lower to 5.18 higher)	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 194: Mixed modality manual therapy plus self-management compared with self-management

			Quality as	sessment	ı.		No of pati	ents		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	MIXED MODALITY+ self-management	self- management	Relative (95% CI)	Absolute		
Quality of	f life (SF-36 F	Physical c	omponent summ	ary score 0-100)) <4 months (E	Setter indicated by	/ lower values)					
	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	259	227	-	MD 2.52 higher (1.23 to 3.81 higher)	⊕⊕OO LOW	CRITICAL
Quality o	f life (SF-36 F	Physical c	omponent summ	ary score 0-100)) >4 months (B	Setter indicated by	/ lower values)					
	randomised trials	- 3	no serious inconsistency		no serious imprecision	none	252	221	-	MD 1.68 higher (0.08 to 3.28 higher)	⊕⊕OO LOW	CRITICAL
Quality of	f life (SF-36 N	Mental co	mponent summa	ry score 0-100)	<4 months (Be	tter indicated by I	ower values)					
	randomised trials	very serious ^a	no serious inconsistency		no serious imprecision	none	259	227	-	MD 2.87 higher (1.26 to 4.48 higher)	⊕⊕OO LOW	CRITICAL
Quality o	f life (SF-36 N	Mental co	mponent summa	ry score 0-100)	>4 months (Ber	tter indicated by I	ower values)					
	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	252	221	-	MD 1.68 higher (0.32 lower to 3.68 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	f life (EQ-5D,	0-10) <4	months (Better in	dicated by high	ner values)							
1	randomised	very	no serious	no serious	no serious	none	342	346	-	MD 0.05 higher	⊕⊕00	CRITICAL

	trials	serious	inconsistency	indirectness	imprecision					(0.01 to 0.09 higher)	LOW	
		3003	<u> </u>	1						(()		
Quality o	f life (EQ-5D,	0-10) >4	months (Better in	dicated by high	ner values)					<u> </u>		
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	342	346	-	MD 0.04 higher (0.01 lower to 0.08 higher)	⊕⊕OO LOW	CRITICAL
Pain (Mo	dified Von Ko	orff scale	0-100 converted	to 0-10) - <4 mo	nths (Better in	dicated by lower v	/alues)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	275	239	-	MD 0.87 lower (1.3 to 0.44 lower)	⊕⊕OO LOW	CRITICAL
Pain (Mo	dified Von Ko	orff scale	0-100 converted	to 0-10) - >4 mo	nths (Better in	dicated by lower v	/alues)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	264	235	-	MD 0.59 lower (1.04 to 0.13 lower)	⊕⊕OO LOW	CRITICAL
Function	(RMDQ, 0-24	l) - <4 mo	nths (Better indic	ated by lower v	alues)							
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	287	256	-	MD 1.57 lower (2.37 to 0.77 lower)	⊕⊕OO LOW	CRITICAL
Function	(RMDQ, 0-24	l) - >4 mo	nths (Better indic	ated by lower v	alues)							
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	273	248	-	MD 1.01 lower (1.84 to 0.18 lower)	⊕⊕OO LOW	CRITICAL
Function	(Modified Vo	n Korff s	cale 0-100 conve	rted to 0-10) - <	4 months (Bette	er indicated by lov	wer values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	275	239	-	MD 0.4 lower (0.83 lower to 0.03 higher)	⊕⊕OO LOW	CRITICAL
Function	(Modified Vo	on Korff s	cale 0-100 conve	rted to 0-10) - >	4 months (Bette	er indicated by lov	wer values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	262	235	-	MD 0.57 lower (0.99 to 0.14 lower)	⊕⊕OO LOW	CRITICAL
Respond	er criteria (>3	30% impre	ovement in RMD0	Q) - <4 months								
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	193/268 (72%)	125/255 (49%)	RR 1.47 (1.27 to 1.7)	221 more per 1000 (from 123 more to 333 more)	⊕⊕OO LOW	IMPORTANT

Respond	er criteria (>3	0% impro	ovement in RMD0	ર) - >4 months								
1		- 3		no serious indirectness	Serious ^b	none	187/275 (68%)	0%	RR 1.21 (1.06 to 1.39)	118 more per 1000 (from 34 more to 219 more)	⊕OOO VERY LOW	IMPORTANT

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 195: Mixed modality manual therapy plus self-management compared with self-management

			Quality as	sessment			No of patient	ts.		Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Mixed modality manual therapy + exercise (biomech) + self- management	self- management	Relative (95% CI)	Absolute	Quality	Importance
Quality o	of life (SF-36	Physical	component sum	mary score, 0-1	100) <4 months	(Better indicated	d by lower values)					
	randomised trials	- ,			no serious imprecision	none	231	227	-	MD 2.55 higher (1.22 to 3.88 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life (SF-36	Physical	component sum	mary score, 0-1	100) >4 months	s (Better indicated	d by lower values)					
	randomised trials	,		no serious indirectness	no serious imprecision	none	221	221	-	MD 2.53 higher (0.78 to 4.28 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life (SF-36	Mental co	omponent summ	ary score, 0-10	0) <4 months (Better indicated	by lower values)					
	randomised trials	- ,			no serious imprecision	none	231	227	-	MD 2.3 higher (0.68 to 3.92 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life (SF-36	Mental co	omponent summ	ary score, 0-10	0) >4 months (Better indicated	by lower values)					
1	randomised trials	- ,		no serious indirectness	Serious ^b	none	221	221	-	MD 1.3 higher (0.75 lower to	⊕OOO VERY	CRITICAL

										3.35 higher)	LOW	
Quality o	of life (EQ-5D	, 0-10) <4	months (Better	indicated by hi	gher values)							
1		very seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	322	326	-	MD 0.03 higher (0 to 0.07 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life (EQ-5D	, 0-10) >4	months (Better	indicated by hi	gher values)							
1	randomised trials	very seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	322	326	-	MD 0.05 higher (0 to 0.1 higher)	⊕⊕OO LOW	CRITICAL
Pain (mo	dified Von K	orff 0-10	0 converted to 0	-10 scale) - <4 n	nonths (Better	indicated by low	er values)					
1	randomised trials	very seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	246	239	-	MD 0.82 lower (1.26 to 0.38 lower)	⊕⊕OO LOW	CRITICAL
Pain (mo	dified Von K	orff 0-100	0 converted to 0	-10 scale) - >4 n	nonths (Better	indicated by low	er values)					
1	randomised trials	very seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	245	235	-	MD 0.67 lower (1.13 to 0.21 lower)	⊕⊕OO LOW	CRITICAL
Function	(RMDQ, 0-24	4) <4 moı	nths (Better indi	cated by lower	values)							
1	randomised trials	very seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	258	256	-	MD 1.87 lower (2.65 to 1.09 lower)	⊕⊕OO LOW	CRITICAL
Function	(RMDQ, 0-24	4) >4 moı	nths (Better indi	cated by lower	values)							
1	randomised trials	very seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	257	248	-	MD 1.3 lower (2.12 to 0.48 lower)	⊕⊕OO LOW	CRITICAL
Function	(modified V	on Korff	0-100 converted	to 0-10 scale) -	<4 months (B	etter indicated by	lower values)					
1	randomised trials	very seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	246	239	-	MD 0.55 lower (0.97 to 0.14 lower)	⊕⊕OO LOW	CRITICAL
Function	(modified V	on Korff	0-100 converted	to 0-10 scale) -	>4 months (B	etter indicated by	lower values)					

1		,			no serious imprecision	none	246	235	-	MD 0.67 lower (1.11 to 0.23 lower)	⊕⊕OO LOW	CRITICAL
Responder criteria (>30% improvement in RMDQ) <4 months												
1		- ,	no serious inconsistency		no serious imprecision	none	185/260 (71.2%)	0%	RR 1.45 (1.25 to 1.68)	221 more per 1000 (from 123 more to 333 more)	⊕⊕OO LOW	IMPORTANT
Responder criteria (>30% improvement in RMDQ) >4 months												
1		,	no serious inconsistency	no serious indirectness	Serious ^b	none	180/246 (73.2%)	0%	RR 1.31 (1.14 to 1.49)	-	⊕OOO VERY LOW	IMPORTANT

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 196: Mixed modality manual therapy + exercise (biomechanical) compared to exercise (biomechanical) + self-management for low back pain with or without sciatica (mixed population)

Quality assessment							No of patients		Effect		∩ualitv	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Exercise + self management	self management + manual therapy	Relative (95% CI)		Quanty	ps. tanoo
Function (RMDQ 0-24) - < 4 months (Better indicated by lower values)												
					no serious imprecision	none	287	225	-	MD 0.38 lower (1.17 lower to 0.41 higher)	⊕⊕OO LOW	CRITICAL
Function (RMDQ 0-24) - > 4 months (Better indicated by lower values)												
					no serious imprecision	none	273	216	-	MD 0.59 lower (1.42 lower to 0.24 higher)	⊕⊕OO LOW	CRITICAL

Pain (Vo	n Korff 0-10) -	< 4 mont	hs (Better indica	ted by lower val	ues)							
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	275	204	-	MD 0.38 lower (0.83 lower to 0.06 higher)	⊕⊕OO LOW	CRITICAL
Pain (Vo	n Korff 0-10) -	> 4 mont	hs (Better indica	ted by lower val	ues)				•			
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	264	200	-	MD 0.01 higher (0.46 lower to 0.49 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life (SF36 0-	100) - < 4	months: Physica	al component (B	Setter indicated	by lower values)						
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	259	191	-	MD 0.21 higher (1.08 lower to 1.5 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life (SF36 0-	100) - > 4	months: Physica	al component (B	Setter indicated	by lower values)						
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	252	194	-	MD 0.21 lower (1.85 lower to 1.43 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life (SF36 0-	100) - < 4	months: Mental	component (Bet	tter indicated by	/ lower values)			<u>.</u>			
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	259	191	-	MD 2.4 higher (0.69 to 4.11 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life (SF36 0-	100) - > 4	months: Mental	component (Bet	tter indicated by	y lower values)						
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	252	194	-	MD 1.32 higher (0.77 lower to 3.41 higher)	⊕⊕OO LOW	CRITICAL
Function	(Von Korff 0-	·10) - < 4 r	months (Better in	dicated by lowe	r values)							
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	275	205	-	MD 0.14 higher (0.29 lower to 0.57 higher)	⊕⊕OO LOW	CRITICAL
unction	(Von Korff 0-	·10) - > 4 r	months (Better in	dicated by lowe	r values)							

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 197: Manual therapy (manipulation) plus exercise (biomechanical) plus self-management compared with self-management

			Quality as	sessment			No of patien	ts		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation + exercise (biomech) + self- management	self- management	Relative (95% CI)	Absolute	Quanty	Importance
Quality o	f life (15D 0 to	o 1) - >4 n	nonths (Better in	dicated by lowe	r values)							
		- ,	no serious inconsistency	no serious indirectness	no serious imprecision	none	63	67	-	MD 0.01 lower (0.03 lower to 0.01 higher)	⊕⊕OO LOW	CRITICAL
Pain (0-10	00 VAS conve	erted to 0	-10) - >4 months	(Better indicate	d by lower valu	ies)						
	randomised trials	,	no serious inconsistency	no serious indirectness	Serious ^b	none	96	100	-	MD 0.65 lower (1.3 lower to 0 higher)	⊕000 VERY LOW	CRITICAL
Function	(ODI, 0-100)	>4 month	s (Better indicate	ed by lower valu	ies)							
		- ,	no serious inconsistency	no serious indirectness	Serious ^b	none	96	100	-	MD 2.8 lower (6.05 lower to 0.45 higher)	⊕000 VERY LOW	CRITICAL
Healthca	re utilisation	(visits to	physicians) >4 m	onths (Better in	ndicated by low	er values)						
		very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	96	100	-	MD 0.3 lower (1.13 lower to 0.53 higher)	⊕⊕OO LOW	IMPORTANT
Healthcar	re utilisation	(visits to	physiotherapy or	other therapies	s) >4 months (E	Better indicated b	y lower values)					

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		,		no serious indirectness	Serious ^b	none	96	100	-	MD 1.6 higher (0.5 lower to 3.7 higher)	⊕OOO VERY LOW	IMPORTANT
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^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 198: Manual therapy (mixed modality: manipulation plus soft tissue techniques - massage) plus exercise (biomech) plus self-management compared with exercise (McKenzie) plus self-management

			Quality as:	sessment			No of pat	iients	I	Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation + massage + exercise (biomech) + self- management	exercise (McKenzie) + self-management	Relative (95% CI)	Absolute	Quality	Importance
Pain (ba	ck and leg pa	ain 0-60)	- <4 months (Be	tter indicated b	y lower value	s)						
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	161	168	-	MD 1.4 lower (4.14 lower to 1.34 higher)	⊕⊕⊕O MODERATE	CRITICAL
Pain (ba	ck and leg pa	ain 0-60)	- >4 months (Be	tter indicated b	y lower value	s)						
1	randomised trials		no serious inconsistency		no serious imprecision	none	163	161	-	MD 2.8 lower (5.77 lower to 0.17 higher)	⊕⊕⊕O MODERATE	CRITICAL
Function	n (RMDQ, 0-2	4) <4 mo	nths (Better indi	cated by lower	values)							
1	randomised trials		no serious inconsistency		no serious imprecision	none	161	168	-	MD 1.5 lower (2.76 to 0.24 lower)	⊕⊕⊕O MODERATE	CRITICAL
Function	n (RMDQ, 0-2	4) >4 mo	nths (Better indi	cated by lower	values)							
1	randomised trials	Seriousª	no serious inconsistency		no serious imprecision	none	163	161	-	MD 1.5 lower (2.87 to 0.13 lower)	⊕⊕⊕O MODERATE	CRITICAL

Healthca	are utilisatior	ı (contac	t with healthcare	in previous 2	months) <4 m	onths						
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	70/160 (43.8%)	35.3%	RR 1.24 (0.95 to 1.62)	85 more per 1000 (from 18 fewer to 219 more)	⊕⊕OO LOW	IMPORTANT
Healthca	are utilisatior	ı (contac	t with healthcare	in previous 2	months) >4 m	onths						
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	89/163 (54.6%)	87/162 (53.7%)	RR 1.02 (0.83 to 1.24)	11 more per 1000 (from 91 fewer to 129 more)	⊕⊕⊕O MODERATE	IMPORTANT
Respond	der criteria ("	Success	" - decrease 5 po	oints or absolu	te score belov	w 5 points on RM	DQ) <4 months					
1	randomised trials		no serious inconsistency		no serious imprecision	none	95/161 (59%)	120/168 (71.4%)	RR 0.83 (0.7 to 0.97)	121 fewer per 1000 (from 21 fewer to 214 fewer)	⊕⊕⊕O MODERATE	CRITICAL
Respond	der criteria ("	Success	" - decrease 5 po	oints or absolu	te score belov	w 5 points on RM	DQ) >4 months					
1	randomised trials		no serious inconsistency		no serious imprecision	none	101/163 (62%)	113/161 (70.2%)	RR 0.88 (0.75 to 1.03)	84 fewer per 1000 (from 175 fewer to 21 more)	⊕⊕⊕O MODERATE	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 199: Manual therapy (manipulation) + exercise +self-management (education + advice to stay active) compared with exercise + self-management (education + advice to stay active)

			Quality asse	essment			No of pat	ients		Effect	O like	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation + education + exercise + self-management		Relative (95% CI)	Absolute	Quanty	Importance

Table 199: Manual therapy (manipulation) + exercise +self-management (education + advice to stay active) compared with exercise + self-man

Pain (0-1	00 VAS conv	erted to 0)-10) - <4 months	(Better indicate	ed by lower v	/alues)						
1		- 3		no serious indirectness	Serious ^b	none	31	33	-	MD 0.58 lower (1.49 lower to 0.33 higher)	⊕OOO VERY LOW	CRITICAL
Function	(ODI, 0-100)	<4 month	ns (Better indicat	ed by lower val	ues)							
1		- 3			very serious ^b	none	31	33	-	MD 0 higher (7.25 lower to 7.25 higher)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 200: Manual therapy (manipulation) + self-management (advice) + pharmacological therapy (NSAIDs) compared with usual care

			Quality asses	ssment			No of patients			Effect	Qualita	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Manipulation + self management + NSAIDS	Usuai	Relative (95% CI)	Absolute	Quality	Importance
Function	(RMDQ, 0-24	change sco	re) < 4 months (fo	llow-up 16 week	s; range of s	cores: 0-24; Bette	r indicated by lower valu	ıes)				
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^a	none	37	35	-	MD 2.54 lower (4.37 to 0.71 lower)	⊕⊕⊕O MODERATE	CRITICAL
Function	(RMDQ, 0-24	change sco	re) > 4 months (fo	llow-up 24 week	s; range of s	cores: 0-24; Bette	r indicated by lower valu	ıes)				
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^a	none	36	35	-	MD 2.58 lower (4.41 to 0.75 lower)	⊕⊕⊕O MODERATE	CRITICAL
Quality of	f life (SF-36 b	odily pain, 0	-100 change score	e) < 4 months (fo	ollow-up 16 w	eeks; range of sc	ores: 0-100; Better indic	ated by	higher v	alues)		
1	randomised trials		no serious inconsistency	no serious indirectness	very serious ^a	none	37	35	-	MD 1.83 higher (3.54 lower to 7.2 higher)	⊕⊕OO LOW	CRITICAL
Quality of	f life (SF-36 p	hysical func	tion, 0-100 change	e score) < 4 mon	ths (follow-u	p 16 weeks; rang	e of scores: 0-100; Bette	r indica	ted by hi	gher values)		

1			no serious inconsistency	no serious indirectness	Serious ^a	none	37	35	-	MD 4.77 higher (1.96 lower to 11.5 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality o	f life (SF-36 b	odily pain, 0	-100 change scor	e) > 4 months (fo	ollow-up 24 v	veeks; range of sc	ores: 0-100; Better indic	ated by	higher v	alues)		
1			no serious inconsistency	no serious indirectness	Serious ^a	none	36	35	-	MD 3.38 higher (1.99 lower to 8.75 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality o	f life (SF-36 p	hysical func	tion, 0-100 chang	e score) > 4 mon	ths (follow-u	up 24 weeks; rang	e of scores: 0-100; Bette	r indica	ted by hi	gher values)		
1			no serious inconsistency	no serious indirectness	very serious ^a	none	36	35	-	MD 3 lower (9.73 lower to 3.73 higher)	⊕⊕OO LOW	CRITICAL

^a Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

Acupuncture

Acupuncture versus placebo/sham J.9.1

Table 201: Acupuncture versus placebo/sham in low back pain without sciatica

			Qua	lity assessme	nt		No of p	patients		Effect	Over life.	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Acupuncture	Placebo/sham	Relative (95% CI)	Absolute	Quality	Importance
Quality o	of life (SF-36	Physical	component sum	nmary score 0	–100) ≤4 mont	ths (range of scores: 0–100; B	etter indicated	d by higher val	ues)	,		
		no serious risk of		no serious indirectness	Serious ^b	none	510	442	-	MD 2.44 higher (0.65 lower to 5.54 higher)	⊕⊕OO LOW	CRITICAL

	1		T	T	1	1						
2	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	no serious imprecision	none	510	442	-	MD 0.13 lower (1.25 lower to 1.51 higher)	⊕⊕⊕⊕ HIGH	CRITICAL
Quality	of life (SF-36	6 Physica	l component sur	mmary score 0	–100) > 4 mor	ths (range of scores: 0–100; B	etter indicate	d by higher va	lues)			
2	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^b	none	510	440	-	MD 2.24 higher (0.92 to 3.56 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality	of life (SF-36	6 Mental o	component sum	mary score 0–1	100) > 4 month	s (Better indicated by lower va	alues)			•		
2	randomised trials	no serious risk of bias	Serious ^c	no serious indirectness	no serious imprecision	none	510	440	-	MD 1.23 higher (2.14 lower to 4.6 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality	of life (SF-36	6 General	health 0–100) ≤4	1 months (Bett	er indicated b	y lower values)						
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^b	none	40	40	-	MD 5.6 higher (4.37 lower to 15.57 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality	of life (SF-36	6 Physica	I function 0–100) ≤4 months (B	etter indicate	d by lower values)						
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^b	none	40	40	-	MD 13.1 higher (3.81 to 22.39 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality	of life (SF-36	6 Physica	l role limitation (0–100) ≤4 mon	ths (Better inc	icated by lower values)						
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^b	none	40	40	-	MD 23 higher (7.57 to 38.43 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality	of life (SF-36	Bodily p	oain 0–100) ≤4 m	onths (Better i	ndicated by lo	wer values)					•	
2	randomised trials	no serious	no serious inconsistency	no serious indirectness	Serious ^b	none	180	110	-	MD 8.85 higher (3.58 to 14.12	⊕⊕⊕O MODERATE	CRITICAL

												<u>.</u>
		risk of								higher)		
		bias										
Quality	of life (SF-36	Vitality (0–100) ≤4 months	s (Better indica	ated by lower	values)						
1		no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^b	none	40	40	-	MD 10.8 higher (0.46 to 21.14 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality	of life (SF-36	Social fu	unction 0–100)≤4	months (Bette	er indicated by	y lower values)						
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	40	40	-	MD 7.2 higher (2.47 lower to 16.87 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality	of life (SF-36	Mental h	nealth 0–100) ≤4 i	months (Bette	r indicated by	lower values)						
1			no serious inconsistency	no serious indirectness	no serious imprecision	none	40	40	-	MD 1.2 higher (8.73 lower to 11.13 higher)	⊕⊕⊕⊕ HIGH	CRITICAL
Quality	of life (SF-36	Emotion	al role limitation	0–100) ≤4 mo	nths (Better ir	ndicated by lower values)						
1		no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^b	none	40	40	-	MD 5 higher (9.64 lower to 19.64 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality	of life (SF-36	Bodily p	ain 0–100) > 4 m	onths (Better i	indicated by lo	ower values)						
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	137	68	-	MD 8.4 higher (1.71 to 15.09 higher)	⊕⊕⊕O MODERATE	CRITICAL
Pain se	verity (VAS ()–10) ≤4 n	nonths (range of	scores: 0-10;	Better indicat	ed by lower values)						
8	randomised trials	no serious risk of bias	Serious ^c	no serious indirectness	Serious ^b	none	864	806	-	MD 0.80 lower (1.29 to 0.32 lower)	⊕⊕OO LOW	CRITICAL

Low back pain and sciatica in over 16s Quality assessment

Pain sev	verity (VAS ()–10) > 4 ı	months (range o	f scores: 0-10	; Better indica	ted by lower values)						
	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	no serious imprecision	none	758	700	-	MD 0.26 lower (0.51 lower to 0.01 higher)	⊕⊕⊕ HIGH	CRITICAL
Functio	n (RMDQ, 0-	·24) >4 mo	onths (Better ind	icated by lowe	er values)							
	randomised trials	no serious risk of bias	very serious	no serious indirectness	very serious ^b	none	147	152	-	MD 0.20 lower (1.52 lower to 1.12 higher)	⊕OOO VERY LOW	CRITICAL
Functio	n (RMDQ, 0-	·24) ≤4 mo	onths (Better ind	icated by lowe	er values)							
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	no serious imprecision	none	192	199	1	MD 1.38 lower (6.08 lower to 3.31 higher)	⊕⊕⊕⊕ HIGH	CRITICAL
Functio	n (ODI) ≤4 m	onths [ch	nange score] (Be	tter indicated	by lower value	es)						
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^b	none	57	58	-	MD 0.15 lower (0.30 lower to 0.00 higher)	⊕⊕⊕O MODERATE	CRITICAL
Functio	n (ODI) > 4 n		hange score] (Be	etter indicated	by lower valu	es)						
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	57	59	-	MD 0.2 lower (0.5 lower to 0.1 higher)	⊕⊕⊕O MODERATE	CRITICAL
Functio	n (FFbH-R/	HFAQ) <	4 months (Bett	er indicated l	ov higher val	ues)				'		
	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision ^b	none	513	446	-	MD 4.05 higher (1.22 to 6.88 higher)	⊕⊕⊕⊕ HIGH	CRITICAL
Functio	n (FFbH-R/	HFAQ) >	4 months (Bett	er indicated I	oy higher val	ues)						

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2	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	no serious imprecision ^b	none	514	444	1	MD 4.22 higher (1.32 to 7.13 higher)	⊕⊕⊕⊕ HIGH	CRITICAL
Function	n (PDI) ≤4 m	onths (Be	etter indicated by	lower values)								
2	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	180	115	ı	MD 3.17 lower (6.3 to 0.05 lower)	⊕⊕⊕⊕ HIGH	CRITICAL
Function	n (PDI) >4 m	onths (Be	etter indicated by	lower values)								
2	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	no serious imprecision	none	177	133	1	MD 2.58 lower (5.82 lower to 0.67 higher)	⊕⊕⊕⊕ HIGH	CRITICAL
Function	n (HFAQ) ≤4	months (Better indicated	by lower value	es)							
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	no serious imprecision	none	373	376	-	MD 4.10 lower (7.37 to 0.83 lower)	⊕⊕⊕⊕ HIGH	CRITICAL
Function	n (HFAQ) >4	months (Better indicated	by lower value	es)							
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	377	376	-	MD 4.60 lower (1.31 to 7.89 lower)	⊕⊕⊕⊕ HIGH	CRITICAL
Psychol	ogical distre	ess (BDI)	≤4 months (Bette	er indicated by	lower values							
1		no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^b	none	57	58	-	MD 0.18 lower (0.38 to 0.02 lower)	⊕⊕⊕O MODERATE	CRITICAL
Psychol	ogical distre	ess (BDI)	> 4 months (Bett	er indicated b	y lower values	s)						
1	randomised trials	no serious	no serious inconsistency	no serious indirectness	no serious imprecision	none	57	59	-	MD 0.08 lower (0.31 lower to 0.15	⊕⊕⊕⊕ HIGH	CRITICAL

		risk of bias								higher)		
Devehol	!	<u> </u>	l S) ≤4 months (Be	attor indicated	by lower valu	06)						
1	randomised trials	no	no serious inconsistency	no serious	no serious imprecision	none	40	45	-	MD 2.60 lower (4.86 to 0.34 lower)	⊕⊕⊕⊕ HIGH	CRITICAL
Psychol	ogical distre	ess (HAD	S) > 4 months (B	etter indicated	by lower valu	ies)						
1			no serious inconsistency	no serious indirectness	no serious imprecision	none	40	45	-	MD 1.5 lower (3.63 lower to 0.63 higher)	⊕⊕⊕⊕ HIGH	CRITICAL
Psychol	logical distre	ess (CES-	D) ≤4 months (B	etter indicated	l by lower valu	ies)						
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	140	70	-	MD 0.5 lower (3.14 to 2.14 higher)	⊕⊕⊕⊕ HIGH	CRITICAL
Psychol	logical distre	ess (CES-	D) > 4 months (E	Better indicated	d by lower val	ues)						
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	137	68	-	MD 2.5 lower (5.26 lower to 0.26 higher)	⊕⊕⊕O MODERATE	CRITICAL
Serious	adverse eve	ents (not	treatment related	I)	•				,			
2	randomised trials		no serious inconsistency	no serious indirectness	very serious ^b	none	25/527 (4.7%)	5.7%		11 more per 1000 (from 21 fewer to 71 more)	⊕⊕OO LOW	IMPORTANT
Adverse	e effects (po	ssibly rela	ated to treatment	t)								
2	randomised trials		no serious inconsistency	no serious indirectness	very serious ^b	none	21/298 (7%)	8.6%	RR 2.19 (0.09 to 53.93)	102 more per 1000 (from 78 fewer to 1000 more)	⊕⊕OO LOW	IMPORTANT

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Days wi	Days with analgesics <4 months (Better indicated by lower values)													
1		no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^b	none	140	70	1	MD 2.9 lower (5 to 0.8 lower)	⊕⊕⊕O MODERATE	IMPORTANT		
Respon	Responder criteria (50%)													
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	36/47 (76.6%)	29.3%	RR 2.62 (1.59 to 4.32)	475 more per 1000 (from 173 more to 973 more)	MODERATE	IMPORTANT		

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 202: Acupuncture vs placebo/sham in low back pain with/without sciatica (overall population)

			Quality asses	ssment			No of	oatients		Effect	Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Acupuncture	Placebo/sham	Relative (95% CI)	Absolute			
Pain seve	ain severity (VAS 0-10) <4 months (Better indicated by lower values)												
2				no serious indirectness	Serious ^a	none	47	43	-	MD 0.52 lower (1.27 lower to 0.24 higher)		CRITICAL	
Function	(RMDQ, 0-24)	<4 months	(Better indicated	by lower values	s)								
2				no serious indirectness	Serious ^a	none	47	43	-	MD 0.83 lower (2.97 lower to 1.31 higher)		CRITICAL	
Overall -	Responder cr	riteria (impr	ovement in function	on >35%) <4 mc	nths								
1				no serious indirectness	Serious ^a	none	50/68 (73.5%)	96/137 (70.1%)	OR 1.19 (0.62 to 2.28)	35 more per 1000 (from 109 fewer to 142 more)	⊕⊕⊕O MODERATE	IMPORTANT	

^{°12 &}gt; 75%; unexplained hetrogeneity. RE analysis used.

Overall (Overall (mixed) Adverse effects possibly related to treatment													
2	randomised trials			no serious indirectness	Serious ^a	none	4/93 (4.3%)	7/163 (4.3%)	RR 0.95 (0.29 to 3.08)	2 fewer per 1000 (from 30 fewer to 89 more)	0000	IMPORTANT		

^a Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

J.9.2 Acupuncture versus usual care

Table 203: Acupuncture versus usual care in low back pain without sciatica

			Quality asse	essment			No of pati	ents		Effect	Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Acupuncture	Usual care	Relative (95% CI)	Absolute			
Quality of	uality of life (SF-36 Physical component score 0-100) <4 months (range of scores: 0-100; Better indicated by higher values)												
2	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	no serious imprecision	none	510	435	-	MD 5.11 higher (2.83 to 7.39 higher)	⊕⊕⊕⊕ HIGH	CRITICAL	
Quality of	ality of life (SF-36 Mental component score 0-100) <4 months (range of scores: 0-100; Better indicated by higher values)												
2	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	serious ¹	none	510	435	-	MD 1.74 higher (0.29 to 3.19 higher)	⊕⊕⊕O MODERATE	CRITICAL	
Quality of	f life (SF-12 P	hysical comp	oonent score 0-10	0) 4 months - 1 y	ear (range of se	cores: 0-100; Bette	er indicated by	y higher	values)				
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	373	364	-	MD 5.8 higher (4.36 to 7.24 higher)	⊕⊕OO LOW	CRITICAL	
Quality of	f life (SF-12 M	lental compo	nent score 0-100)	4 months - 1 yea	ar (range of sco	res: 0-100; Better	indicated by h	nigher va	alues)				
1	randomised trials	no serious risk of bias	no serious inconsistency		no serious imprecision	none	373	364	-	MD 1.5 higher (0.15 lower to 3.15 higher)	⊕⊕⊕⊕ HIGH	CRITICAL	
Quality of	f life (SF-36 B	odily pain 0-	100)<4 months (ra	inge of scores: 0	-100; Better ind	licated by higher	/alues)						

Quality assessment

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Function	(PDI) 4 month	ns-1 vear (Be	etter indicated by	lower values)								
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	serious²	none	40	46	-	MD 6.7 lower (11.53 to 1.87 lower)	⊕⊕⊕O MODERATE	CRITICAL
Function	(HFAQ) <4 m	onths (Bette	r indicated by low	er values)								
3	randomised trials	no serious risk of bias	very serious ⁴	no serious indirectness	serious ²	none	1844	1771	-	MD 11.68 lower (23.2 to 0.17 lower)	⊕OOO VERY LOW	CRITICAL
Function	(HFAQ) > 4 m	onths (Bette	er indicated by lov	ver values)								
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	serious ²	none	337	364	-	MD 11.10 lower (14.49 to 7.71 lower)	⊕⊕⊕O MODERATE	CRITICAL
Psycholo	gical distress	(CES-D 0-10	00) < 4 months (B	etter indicated b	y lower values)			,				
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	140	74	-	MD 0.8 lower (3.6 lower to 2 higher)	⊕⊕⊕O MODERATE	CRITICAL
Psycholo	gical distress	(HADS 0-42) < 4 months (Bet	ter indicated by	lower values)							
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	40	46	-	MD 2.8 lower (4.91 to 0.69 lower)	⊕⊕OO LOW	CRITICAL
Psycholo	gical distress	(HADS 0-42) 4 months - 1 yea	ar (Better indicat	ed by lower val	ues)						
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	40	46	-	MD 2.3 lower (4.48 to 0.12 lower)	⊕⊕OO LOW	CRITICAL
Serious a	dverse event	s (not treatm	ent related)									
2	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	very serious ²	none	25/527 (4.7%)	6.8%	RR 0.93 (0.52 to 1.67)	5 fewer per 1000 (from 33 fewer to 46 more)	⊕⊕OO LOW	IMPORTANT
Respond	er criteria (50°	%)										
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	31/47 (66%)	13.9%	RR 4.75 (2.05 to 10.99)	521 more per 1000 (from 146 to 1000 more)	⊕⊕⊕O MODERATE	IMPORTANT
Days with	n analgesics (Better indica	ted by lower valu	es)	1		()	ļ.	1 .0.00,			

	randomised trials			no serious indirectness	serious ²	none	140	74	ı	MD 4.30 lower (6.44 to 2.16 lower)	⊕⊕OO LOW	IMPORTANT
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^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 204: Acupuncture versus usual care in low back pain with/without sciatica (overall population)

			Quality ass	essment			No of pati	ients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Acupuncture	Usual care	Relative (95% CI)	Absolute		
Quality of	life (EQ5D 0-	–1) ≤4 month	s (Better indicate	d by lower value	es)							
	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	96	42	-	MD 0.1 higher (0.01 to 0.19 higher)	⊕⊕OO LOW	CRITICAL
Quality of	life (EQ5D 0-	–1) > 4 month	ns (Better indicate	ed by lower valu	es)							
	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	145	68	-	MD 0.01 higher (0.05 lower to 0.08 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality of	f life (SF-36 G	eneral health	n 0–100) ≤4 montl	ns (Better indica	ted by lower va	lues)						
		very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	74	69	-	MD 7.4 higher (1.35 to 13.45 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	life (SF-36 P	hysical role l	limitation 0–100) :	≤4 months (Bette	er indicated by	lower values)						
		very seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	74	69	-	MD 14.9 higher (1.58 to 28.22 higher)	⊕⊕OO LOW	CRITICAL
Quality of	f life (SF-36 b	odily pain 0–	100) ≤4 months (l	Better indicated	by lower values	s)						
2	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	218	139	-	MD 5.12 higher (0.22 to 10.03 higher)	⊕000 VERY LOW	CRITICAL

c Heterogeneity, I²=89%, unexplained by subgroup analysis.
d I² >50% and ≤75%; unexplained hetrogeneity. RE analysis used.
e I² >75%; unexplained heterogeneity. RE analysis used.

Quality o	f life (SF-36 P	hysical fund	ction 0–100) ≤4 m	onths (Better in	dicated by lowe	r values)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	74	69	-	MD 8.2 higher (1.54 to 14.86 higher)	⊕000 VERY LOW	CRITICAL
Quality o	f life (SF-36 V	itality 0–100) ≤4 months (Bet	ter indicated by	lower values)							
1	randomised trials	very seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	74	69	-	MD 10.1 higher (3.19 to 17.01 higher)	⊕⊕OO LOW	CRITICAL
Quality o	f life (SF-36 S	ocial function	oning 0–100) ≤4 n	nonths (Better ir	dicated by low	er values)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	74	69	-	MD 7.2 higher (0.77 lower to 15.17 higher)	⊕000 VERY LOW	CRITICAL
Quality o	f life (SF-36 N	lental health	n 0–100) ≤4 month	ns (Better indica	ted by lower va	lues)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	74	69	-	MD 4.6 higher (2.39 lower to 11.59 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	f life (SF-36 E	motional ro	le limitation 0–10	0) ≤4 months (Be	etter indicated I	by lower values)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	74	69	-	MD 13.4 higher (0.11 lower to 26.91 higher)	⊕000 VERY LOW	CRITICAL
Quality o	f life (SF-36 B	odily pain 0	–100) > 4 months	(Better indicate	d by lower valu	ies)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	145	67	-	MD 6.1 higher (0.6 lower to 12.8 higher)	⊕OOO VERY LOW	CRITICAL
Pain seve	erity (VAS 0-1	l0) ≤4 month	ns (Better indicate	ed by lower valu	es)							
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	25	20	-	MD 1.28 lower (2.09 to 0.47 lower)	⊕OOO VERY LOW	CRITICAL
Pain seve	erity (VAS 0-1	10) > 4 mont	hs (Better indicat	ed by lower valu	ies)	•						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	135	57	-	MD 0.1 lower (0.4 lower to 0.2 higher)	⊕⊕OO LOW	CRITICAL
Function	(RMDQ 0-24)) ≤4 months	(Better indicated	by lower values	·)							

Low back pain and sciatica in over 16s Quality assessment

2	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	56	44	-	MD 2.24 lower (3.43 to 1.06 lower)	⊕OOO VERY LOW	CRITICAL
Function	(ODI) >4 mon	ıths (Better i	ndicated by lowe	r values)								
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	134	57	-	MD 1.0 higher (4.16 lower to 6.16 higher)	⊕⊕⊕O MODERATE	CRITICAL
Overall -	Responder cr	riteria (impro	vement in function	on >35%) <4 mon	ths	•						
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^b	none	50/68 (73.5%)	31/70 (44.3%)	OR 3.49 (1.71 to 7.15)	292 more per 1000 (from 133 more to 408 more)	0000	IMPORTANT

a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 205: Acupuncture versus list control in low back pain with/without sciatica

			Quality as	sessment			No of p	oatients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Acu	Waiting list	Relative (95% CI)	Absolute		
Overall SF36 (change scores, <4 months) - Physical (range of scores: 0-100; Better indicated by lower values)												
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	1350	1244	-	MD 4.7 higher (4 to 5.4 higher)	⊕⊕OO LOW	CRITICAL
Overall S	F36 (change s	cores, <4	months) - Mental (range of scores:	0-100; Better in	dicated by lower v	alues)					
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	1350	1244	-	MD 2.1 higher (1.4 to 2.8 higher)	⊕⊕⊕O MODERATE	CRITICAL
Overall Si	F36 (change s	cores, >4	months) - Physica	l (range of score	s: 0-100; Better	indicated by lower	values)					
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	1309	1183	-	MD 0.6 higher (0.2 lower to 1.4 higher)	⊕⊕⊕O MODERATE	CRITICAL
Overall S	F36 (change s	scores, >4	months) - Mental (range of scores:	0-100; Better in	dicated by lower v	alues)					

1	randomised trials	Seriousª			no serious imprecision	none	1309	1183	-	MD 0.2 higher (0.6 lower to 1 higher)	⊕⊕⊕O MODERATE	CRITICAL		
Prescripti	rescription of analgesics													
1	randomised trials	Seriousª			no serious imprecision	none	285/1350 (21.1%)	22.7%		16 fewer per 1000 (from 43 fewer to 18 more)		IMPORTANT		

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Acupuncture versus electrotherapy (TENS) J.9.3

Table 206: Acupuncture versus electrotherapy (TENS) in low back pain without sciatica

			Quality asse	ssment			No of patie	nts		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Acupuncture	TENS	Relative (95% CI)	Absolute		
Pain (VAS	0–10) ≤4 mon	ths (Better	indicated by lower	values)								
2	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	16	16	-	MD 1.54 lower (3.43 lower to 0.36 higher)	⊕⊕OO LOW	CRITICAL
Function (inction (RMDQ 0–24) ≤4 months (Better indicated by lower values)											
1	randomised trials	very seriousª	no serious inconsistency	no serious indirectness	very serious ^b	none	7	6	-	MD 0.8 lower (5.38 lower to 3.78 higher)	⊕OOO VERY LOW	CRITICAL
Function (JOA score 0-1	7) ≤4 mon	ths (Better indicate	d by higher value	s)							
1	randomised trials		no serious inconsistency	no serious indirectness	serious ^d	none	10	10	-	MD 1.42 lower (3.09 lower to 0.25 higher)	⊕⊕OO LOW	CRITICAL
Adverse e	vents											
1	randomised	Serious ^a	no serious	no serious	very	none	3/10	3/10	RR 1 (0.26 to	0 fewer per 1000 (from 222	⊕000	IMPORTANT

trials	inconsistency	indirectness se	erious ^b	(30%)	(30%)	3.81)	fewer to 843 more)	VERY	
					30%		0 fewer per 1000 (from 222 fewer to 843 more)	LOW	

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Acupuncture versus NSAIDs J.9.4

Table 207: Acupuncture versus NSAIDs in low back pain with/without sciatica (overall population)

			Quality asse	ssment			No of pati	ents		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Acupuncture	NSAIDs	Relative (95% CI)	Absolute		
Pain (VAS	0–10) intramı	ıscular dic	clofenac ≤4 months	(Better indicate	d by lower va	alues)						
	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	29	29	-	MD 1.5 higher (0.11 to 2.89 higher)	⊕⊕OO LOW	CRITICAL
Pain (VAS	0–10) oral did	: lofenac ≤₄	4 months (Better in	dicated by lower	values)							
	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	24	20	-	MD 0.37 lower (0 to 0.47 higher)	⊕OOO VERY LOW	CRITICAL
Pain (VAS	0–10) > 4 moi	nths (Bette	er indicated by low	er values)	'							
	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	29	29	-	MD 0.2 lower (1.33 lower to 0.93 higher)	⊕⊕OO LOW	CRITICAL
Function (ODI/RMDQ) ≤	4 months (Better indicated b	y lower values)								
	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	53	49	-	SMD 0.39 higher (0.01 lower to 0.78 higher)	⊕⊕OO LOW	CRITICAL
Function ((ODI 0-100) >	4 months	(Better indicated b	y lower values)								

1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	29	29	-	MD 7.6 lower (16.47 lower to 1.27 higher)	⊕⊕OO LOW	CRITICAL
Healthcar	e utilisation (li	npatient ca	are) > 4 months									
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	19/29 (65.5%)	27/29 (93.1%) 93.1%		279 fewer per 1000 (from 65 fewer to 438 fewer) 279 fewer per 1000 (from 65 fewer to 438 fewer)	⊕⊕OO LOW	IMPORTANT
Healthcare	e utilisation (d	luration of	hospital stay) > 4	months (Better in	dicated by le	ower values)						
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	29	29	-	MD 5.38 lower (10.73 to 0.03 lower)	⊕⊕OO LOW	IMPORTANT

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Combined interventions – acupuncture adjunct J.9.5

Table 208: Acupuncture plus electrotherapy (TENS) compared with usual care in low back pain without sciatica

	or / toup arrec	u. c p.u.s	cicciotherapy	(12:10) compan			ack pain min							
			Quality asse	ssment			No of patier	nts		Effect				
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Acupuncture + TENS	usual care	Relative (95% CI)	Absolute	Quality	Importance		
Pain (0–10	ain (0–100 VAS converted to 0–10) - ≤4 months (follow-up 10 weeks; measured with: VAS; range of scores: 0–10; Better indicated by lower values)													
1	randomised trials	- ,	no serious inconsistency	no serious indirectness	very serious ^b	none	6	7	-	MD 0.89 lower (3.18 lower to 1.4 higher)	⊕OOO VERY LOW	CRITICAL		
Disability (RMDQ 0-24) -	≤4 months	s (follow-up 10 wee	ks; measured with	n: RMDQ; ran	ige of scores: 0-24	; Better indicated	by lowe	r values)					
1		,	no serious inconsistency	no serious indirectness	very serious ^b	none	6	7	-	MD 1.2 lower (4.84 lower to 2.44 higher)	⊕OOO VERY LOW	CRITICAL		

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 209: Acupuncture plus electrotherapy (TENS) compared with electrotherapy (TENS) in low back pain without sciatica

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			Quality asses	ssment			No of patient	s		Effect	Qualife.	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Acupuncture + TENS	TENS	Relative (95% CI)	Absolute	Quality	Importance
Pain (0–10	0 VAS convert	ed to 0–10) - ≤4 months (follow	v-up 10 weeks; me	easured with	: VAS; range of sco	ores: 0-10; Better	indica	ited by lo	ower values)		
		- 3			very serious ^b	none	6	6	-	MD 0.88 lower (2.95 lower to 1.19 higher)	⊕OOO VERY	CRITICAL

											LOW				
Disability	isability (RMDQ 0–24) - ≤4 months (follow-up 10 weeks; measured with: RMDQ; range of scores: 0–24; Better indicated by lower values)														
1	randomised trials	- ,		no serious indirectness	very serious ^b	none	6	6	-	MD 1 lower (4.15 lower to 2.15 higher)	⊕OOO VERY LOW	CRITICAL			

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 210: Acupuncture plus manual therapy (massage) compared with usual care in low back pain without sciatica

			Quality as:	sessment		No of patien	ts		Effect	Ouglitus			
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Acupuncture + massage	usual care	Relative (95% CI)	Absolute	Quality	Importance	
Pain (prop	Pain (proportion of baseline value) - ≤4 months (follow-up 4 weeks; measured with: VAS; range of scores: 0–10; Better indicated by lower values)												
	randomised trials	-)			no serious imprecision	none	27	24	-	MD 0.38 lower (0.55 to 0.21 lower)	⊕⊕OO LOW	CRITICAL	

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 211: Acupuncture plus exercise (biomech plus aerobic) plus self-management compared with exercise (biomechanical plus aerobic) plus selfmanagement in low back pain without sciatica

			Quality asse	essment			No of patie	nts		Effect	Ovality						
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Acupuncture + exercise (biomech + aerobic)	exercise (biomech + aerobic)	Relative (95% CI)	Absolute	Quality	Importance					
Quality of	life (EQ-5D)	- ≤4 month	ns (follow-up 3 mo	uality of life (EQ-5D) - ≤4 months (follow-up 3 months; measured with: EQ-5D; range of scores: 0–1; Better indicated by higher values)													

1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	24	27	-	MD 0.06 lower (0.23 lower to 0.11 higher)	⊕⊕OO LOW	CRITICAL		
Quality o	uality of life (EQ-5D) - >4 months (follow-up 6 months; measured with: EQ5D; range of scores: 0–1; Better indicated by higher values)													
1	randomised trials			no serious indirectness	serious ²	none	24	27	-	MD 0.11 higher (0 to 0.22 higher)	⊕⊕OO LOW	CRITICAL		
Pain (VA	S 0–10) - ≤4 m	onths (fo	llow-up 3 months	measured with	: VAS; range	of scores: 0-10;	Better indicated by lowe	r values)						
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	24	27	-	MD 1.19 higher (0.34 lower to 2.72 higher)	⊕⊕OO LOW	CRITICAL		
Pain (VA	ain (VAS 0–10) - >4 months (follow-up 6 months; measured with: VAS; range of scores: 0–10; Better indicated by lower values)													
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	24	27	-	MD 0.29 lower (1.87 lower to 1.29 higher)	⊕⊕OO LOW	CRITICAL		
Disability	/ (ODI) - ≤4 mo	onths (foll	ow-up 3 months;	measured with:	ODI; range o	of scores: 0–100; I	Better indicated by lowe	r values)						
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	24	27	-	MD 1.36 higher (4.45 lower to 7.17 higher)	⊕⊕OO LOW	CRITICAL		
Disability	/ (ODI) - >4 mo	onths (foll	ow-up 6 months;	measured with:	ODI; range o	of scores: 0-100; I	Better indicated by lowe	r values)						
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	24	27	-	MD 4 lower (12.41 lower to 4.41 higher)	⊕⊕OO LOW	CRITICAL		

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Electrotherapies J.10

Table 212: TENS versus sham for low back pain in low back pain without sciatica

Quality assessment No	No of patients Eff	ect Quality	Importance
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No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	TENS versus sham	Control	Relative (95% CI)	Absolute		
SF-36; str	atum = withou	t sciatica -	· Physical function	; outcome ≤4 mor	nths (range of sc	ores: 0-100; Better	indicated by	higher	values)			
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	15	12	-	MD 19.41 higher (5.79 to 33.03 higher)	⊕⊕OO LOW	CRITICAL
SF-36; str	atum = withou	t sciatica -	Social function; o	utcome ≤4 month	s (range of score	es: 0-100; Better in	dicated by hi	gher val	ues)			
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	15	12	-	MD 17.70 higher (5.97 to 29.43 higher)	⊕⊕OO LOW	CRITICAL
SF-36; str	atum = withou	t sciatica -	Physical role limit	tation; outcome ≤	4 months (range	of scores: 0-100; E	Better indicat	ed by hi	gher valu	ues)		
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	15	12	-	MD 52.76 higher (23.03 to 9 higher)	⊕⊕OO LOW	CRITICAL
SF-36; str	atum = withou	t sciatica -	Emotional role lin	nitation; outcome	≤4 months (rang	je of scores: 0-100;	Better indica	ated by	higher va	alues)		
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	15	12	-	MD 33.36 higher (11.14 to 55.58 higher)	⊕⊕OO LOW	CRITICAL
SF-36; str	atum = withou	t sciatica -	- Mental health; ou	tcome ≤4 months	(range of scores	: 0-100; Better indi	cated by higl	her valu	es)			
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	15	12	-	MD 7.39 higher (0.32 to 14.46 higher)	⊕OOO VERY LOW	CRITICAL
SF-36; str	atum = withou	t sciatica -	· Vitality; outcome	≤4 months (range	of scores: 0-100); Better indicated I	y higher val	ues)	•			
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	15	12	-	MD 4.25 higher (2.61 lower to 11.11 higher)	⊕⊕OO LOW	CRITICAL
SF-36; str	atum = withou	t sciatica -	Bodily pain; outco	ome ≤4 months (r	ange of scores: (0-100; Better indica	ted by highe	r values)			
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	15	12	-	MD 14.98 higher (7.56 to 22.4 higher)	⊕⊕OO LOW	CRITICAL
SF-36; str	atum = withou	t sciatica -	General health pe	rception; outcom	e ≤4 months (ran	nge of scores: 0-10); Better indi	cated by	/ higher v	values)		

Low back pain and sciatica in over 16s Quality assessment

1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	15	12	1	MD 10.51 higher (3.51 to 17.51 higher)	⊕⊕OO LOW	CRITICAL
Back pain	% of baseline	; stratum	= without sciatica;	outcome ≤4 mont	ths (Better indica	ted by lower value	s)					
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	15	15	-	MD 33.62 lower (53.27 to 13.97 lower)	⊕⊕⊕O MODERATE	CRITICAL
Back pain	; stratum = wi	thout scia	tica; outcome ≤4 m	onths (range of s	cores: 0-10; Bett	er indicated by low	er values)					
2	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	52	50	-	MD 0.5 lower (0.53 to 0.47 lower)	⊕⊕⊕O MODERATE	CRITICAL
Function,	RMDQ; stratu	m = witho	ut sciatica; outcom	e ≤4 months (ran	ge of scores: 0-2	4; Better indicated	by lower valu	ıes)				
3	randomised trials	seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	241	249	-	MD 0.36 lower (1.4 lower to 0.68 higher)	⊕⊕⊕O MODERATE	CRITICAL
Function,	ODI 0-100; str	atum = wit	thout sciatica; outo	ome ≤4 months (range of scores:	0-100; Better indic	ated by lower	· values)				
1 (a) Down	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	23	21	-	MD 4.40 lower (5.07 to 3.73 lower)	⊕⊕⊕O MODERATE	CRITICAL

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias (b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 213: TENS versus sham for low back pain in low back pain with or without sciatica

	able 213. TENS Versus shall for low back pain in low back pain with or without sciatica													
			Quality as	sessment			No of pat	tients		Effect	Quality	luono atomo		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	TENS versus sham	Control	Relative (95% CI)	Absolute	Quality	Importance		
SF-36 Con	SF-36 Composite scores; stratum +/- sciatica - Physical composite; outcome ≤4 months (range of scores: 0-100; Better indicated by higher values)													
	randomised trials		no serious inconsistency		no serious imprecision	none	91	83	-	MD 1 higher (1.25 lower to 3.25 higher)	⊕⊕⊕O MODERATE	CRITICAL		
SF-36 Con	F-36 Composite scores; stratum +/- sciatica - Mental composite; outcome ≤4 months (range of scores: 0-100; Better indicated by higher values)													

1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	91	83	-	MD 0.2 higher (3.29 lower to 3.69 higher)	⊕⊕⊕O MODERATE	CRITICAL
Back pain	n (VAS cm); st	ratum +/-	sciatica; outcome	≤4 months (rang	e of scores: 0-1	0; Better indicated	by lower val	lues)				
1	randomised trials	,	no serious inconsistency	no serious indirectness	very serious ^b	none	15	26	-	MD 0.01 lower (1.75 lower to 1.73 higher)	⊕000 VERY LOW	CRITICAL
Back pain	ı VAS: improv	ement of	≥50% from baselin	ne; stratum = +/- s	sciatica; outcom	ne ≤4 months						
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	26/104 (25%)	7/104 (6.7%)	RR 3.71 (1.69 to 8.18)		⊕⊕⊕O MODERATE	CRITICAL
Function;	stratum +/- s	ciatica; ou	tcome ≤4 months	(range of scores	: 0-24; Better in	dicated by lower v	alues)					
1	randomised trials	,	no serious inconsistency	no serious indirectness	no serious imprecision	none	15	26	-	MD 1 lower (4.53 lower to 2.53 higher)	⊕⊕OO LOW	CRITICAL
Roland-M	orris Disabilit	y Questio	nnaire: improvem	ent of 4 points (n	nedian 15 at bas	eline); stratum = +	/- sciatica; o	utcome	≤4 months			
1	randomised trials		no serious inconsistency	no serious indirectness	very serious ^b	none	29/110 (26.4%)	(25%)	to 1.65)	12 more per 1000 (from 82 fewer to 162 more)	VERY LOW	CRITICAL

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias (b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 214: TENS versus usual care for low back pain in low back pain without sciatica

			I care for low be	en pann m ie ii	back pain to	out stratitu							
			Quality ass	sessment		No of patie	ents		Effect	Qualify			
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	TENS versus usual care	Control	Relative II (95% Absolute CI)		Quality	Importance	
Pain VAS;	stratum = wit	hout sciati	ca; outcome ≤4 mo	onths (range of so	cores: 0-10; Bett	er indicated by low	ver values)						
		- ,			no serious imprecision	none	33	37	-	MD 0.45 higher (0.37 to 0.53 higher)	⊕⊕OO LOW	CRITICAL	
Function F	function RMDQ final values; stratum = without sciatica;, outcome ≤4 months (range of scores: 0-24; Better indicated by lower values)												

2	randomised trials	very serious ^{a,}	no serious inconsistency	no serious indirectness	very serious ^b	none	12	14	-	MD 0.20 lower (3.08 lower to 2.68 higher)	⊕OOO VERY LOW	CRITICAL			
Function	Function ODI 0-100 change scores,; stratum = without sciatica; outcome ≤4 months (range of scores: 0-100; Better indicated by lower values)														
1	randomised trials	serious ^a	no serious		no serious	none	21	23	-	MD 6.80 higher (5.17 to 8.43 higher)	⊕⊕⊕O MODERATE	CRITICAL			

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias (b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 215: TENS versus usual care for low back pain in low back pain with or without sciatica

		Quality ass		No of patie	ents		Effect	O alifa.						
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	TENS versus usual care	Control	Relative (95% CI)	Absolute	Quality	Importance		
Pain VAS;	ain VAS; stratum +/- sciatica; outcome ≤4 months (range of scores: 0-10; Better indicated by lower values)													
	randomised trials	very seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	53	49	-	MD 0.25 lower (1.06 lower to 0.56 higher)	⊕⊕OO LOW	CRITICAL		
Quebec Ba	ack Pain Disal	oility Scale;	stratum +/- sciatica	a; outcome ≤4 mo	nths (range of so	cores: 0-100; Better	indicated by lo	wer valu	ies)					
		serious ^{a,b}	inconsistency	no serious indirectness	no serious imprecision	none	53	49	-	MD 0.85 higher (5.21 lower to 6.91 higher)	⊕⊕OO LOW	CRITICAL		

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias (b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 216: TENS versus corset for low back pain without sciatica

	Quality assessment									Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	TENS versus corset	Control	Relative (95%	Absolute	·	·

									CI)					
Pain; stratu	Pain; stratum = without sciatica; outcome ≤4 months (range of scores: 0-10; Better indicated by lower values)													
	randomised trials	- ,		no serious indirectness	serious ^b	none	20	24	-	MD 0.63 higher (1.07 lower to 2.33 higher)	⊕OOO VERY LOW	CRITICAL		

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias (b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 217: TENS versus manipulation for low back pain without sciatica

			Quality asse	ssment			No of patient	s		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	TENS versus manipulation	Control	Relative (95% CI)	Absolute	Quanty	importance
Pain; stratum = without sciatica; outcome ≤4 months (range of scores: 0-10; Better indicated by lower values)												
1	randomised trials	,		no serious indirectness	serious ^b	none	20	43	-	MD 1.45 higher (0.09 lower to 2.99 higher)	⊕000 VERY LOW	CRITICAL

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias (b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 218: TENS versus massage for low back pain without sciatica

			Quality asse	ssment			No of patie	nts		Effect	- Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	TENS versus massage	Contro	Relative (95% CI)	Absolute	Quanty	importance	
Pain; stratum = without sciatica; outcome ≤4 months (range of scores: 0-100; Better indicated by lower values)													
1	randomised	very	no serious	no serious	serious ^b	none	20	20	-	MD 0.76 higher (0.95	⊕OOO	CRITICAL	

	trials	serious ^a	inconsistency	indirectness						lower to 2.47 higher)	VERY LOW	
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- (a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias
- (b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 219: TENS versus massage for low back pain with or without sciatica

			Quality as:	sessment			No of patie	ents		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	TENS versus massage	Control	Relative (95% CI)	Absolute		
Pain rating	n rating index change (%); stratum +/- sciatica; outcome ≤4 months (Better indicated by lower values)											
	randomised trials	very serious ^a		no serious indirectness	no serious imprecision	none	20	21	-	MD 32.3 lower (36.58 to 28.02 lower)	⊕⊕OO LOW	CRITICAL
Responder: >50% decrease in pain; outcome ≤4 months												
	randomised trials	very serious ^a			imprecision	none	17/20 (85%)	8/21 (38.1%)		469 more per 1000 (from 95 more to 1000 more)		IMPORTANT

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 220: PENS versus sham for low back pain without sciatica

			Quality ass	sessment		No of pation	ents		Effect	Ovalite				
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	PENS versus sham		Relative (95% Absolute CI)		Quality	Importance		
SF-36 Com	SF-36 Composite scores; stratum = without sciatica - Mental composite; chronic low back pain; outcome >4 months (Better indicated by higher values)													
	randomised trials			no serious indirectness	serious ^b	none	92	92	-	MD 2.38 lower (6.34 lower to 1.57 higher)	⊕⊕OO LOW	CRITICAL		
SF-36 Com	posite scores	; stratum =	without sciatica - F	Physical composit	te; chronic low ba	ack pain; outcome	>4 months (Bette	er indicat	ed by higher values)				

Low back pain and sciatica in over 16s Quality assessment

4							00	00		MD 4 22 laws (0.20 laws	0000	CRITICAL
	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	92	92		MD 1.23 lower (8.28 lower to 5.82 higher)	⊕⊕⊕O MODERATE	CRITICAL
SF-36 Don	nain scores; st	ratum = w	ithout sciatica - Phy	sical function; ch	ronic low back pa	ain; outcome ≤4 mo	nths (range	of so	cores: 0-	100; Better indicated by hi	gher values))
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	13	12	-	MD 27.98 higher (15.18 to 40.78 higher)	⊕⊕OO LOW	CRITICAL
SF-36 Don	nain scores; st	ratum = w	ithout sciatica - Soc	ial function; chro	nic low back pain	; outcome ≤4 mont	hs (range o	f scoi	res: 0-10	0; Better indicated by high	er values)	
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	13	12	-	MD 26.87 higher (15.32 to 38.42 higher)	⊕⊕OO LOW	CRITICAL
SF-36 Don	nain scores; st	ratum = w	ithout sciatica - Phy	sical role limitation	on; chronic low b	ack pain; outcome :	≤4 months (range	e of scor	es: 0-100; Better indicated	by higher va	alues)
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	13	12	-	MD 55.76 higher (28.34 to 83.18 higher)	⊕⊕OO LOW	CRITICAL
SF-36 Don	nain scores; st	ratum = w	ithout sciatica - Eme	otional role limita	tion; chronic low	back pain; outcome	e ≤4 months	s (ran	ge of sc	ores: 0-100; Better indicate	ed by higher	values)
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	13	12	-	MD 68.42 higher (44.07 to 92.77 higher)	⊕⊕OO LOW	CRITICAL
SF-36 Don	nain scores; st	ratum = w	ithout sciatica - Mer	ntal health; chroni	c low back pain;	outcome ≤4 months	s (range of	score	s: 0-100	Better indicated by highe	r values)	
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	13	12	-	MD 8.48 higher (1.69 to 15.27 higher)	⊕⊕OO LOW	CRITICAL
SF-36 Don	nain scores; st	ratum = w	ithout sciatica - Vita	lity; chronic low I	oack pain; outcon	ne ≤4 months (rang	e of scores	: 0-10	0; Bette	r indicated by higher value	s)	
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	13	12	-	MD 11.89 higher (3.82 to 19.96 higher)	⊕⊕OO LOW	CRITICAL
SF-36 Don	nain scores; st	ratum = w	ithout sciatica - Boo	lily pain; chronic	low back pain; ou	tcome ≤4 months (range of sc	ores:	0-100; E	setter indicated by higher v	alues)	
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	13	12	-	MD 21.05 higher (14.04 to 28.06 higher)	⊕⊕OO LOW	CRITICAL
SF-36 Don	nain scores; st	ratum = w	ithout sciatica - Ger	eral health perce	ption; chronic lov	v back pain; outcon	ne ≤4 montl	ns (ra	nge of s	cores: 0-100; Better indica	ted by highe	r values)
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	13	12	-	MD 24.23 higher (15.63 to 32.83 higher)	⊕⊕OO LOW	CRITICAL

Pain; stra	tum = without	sciatica; o	utcome ≤4 months	(Better indicated	by lower values)	T	1			T			
2	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	30	29	1	SMD 1.33 lower (1.92 to 0.75 lower)	⊕⊕OO LOW	CRITICAL	
Pain; stra	tum = without	sciatica; o	utcome >4 months	(Better indicated	by lower values)								
2	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	92	92	1	SMD 0.05 lower (0.34 lower to 0.24 higher)	⊕⊕⊕O MODERATE	CRITICAL	
Disability (ODI, change score); stratum = without sciatica; outcome ≤4 months (range of scores: 0-24 or 0-50; Better indicated by lower values)													
1	randomised trials	very serious ^a	serious ^c	no serious indirectness	no serious imprecision	none	13	12	-	MD 11.69 lower (14.92 to 8.46 lower)	⊕OOO VERY LOW	CRITICAL	
Function ((RMDQ, final v	alue); strat	tum = without sciati	ca; outcome ≤4 n	nonths (Better ind	icated by lower valu	ues)						
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	17	17	-	MD 2.93 lower (6.11 lower to 0.25 higher)	⊕⊕OO LOW	CRITICAL	
Function ((RMDQ, final v	alue); strat	tum = without sciati	ica; outcome >4 n	nonths (range of s	cores: 0-24 or 0-50	; Better ind	icated	l by lowe	er values)			
2	randomised trials	very serious ^a	serious ^c	no serious indirectness	no serious imprecision	none	92	92		MD 0.81 higher (0.53 lower to 2.15 higher)	VERY LOW	CRITICAL	

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias (b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.
(c) Downgraded by 1 or 2 increments because heterogeneity, 1²=50%, p=0.04, unexplained by subgroup analysis

Table 221: PENS versus usual care for low back pain with or without sciatica

			care for fow ba	pa								
			Quality as:	sessment		No of patie	nts		Effect	Qualifor		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	PENS versus usual care	Control	Relative (95% CI)	Absolute	Quality	Importance
Pain VAS; stratum +/- sciatica; outcome ≤4 months (range of scores: 0-10; Better indicated by lower values)												
	randomised trials	very serious ^a			no serious imprecision	none	53	49	-	MD 0.05 lower (0.95 lower to 0.85 higher)	⊕⊕OO LOW	CRITICAL

Quebec Ba	ack Pain Disat	oility Scale	; stratum +/- sciatio	a; outcome ≤4 mo	onths (range of so	cores: 0-100; Bette	r indicated by lo	wer valu	ies)			
1	randomised trials	,			no serious imprecision	none	53	49	-	MD 1.62 lower (7.75 lower to 4.51 higher)	⊕⊕OO LOW	CRITICAL

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 222: PENS versus TENS for low back pain without sciatica

			Quality as	sessment			No of pati	ents		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	PENS versus TENS	Control	Relative (95% CI)	Absolute	Quanty	importance
SF-36; stra	atum = withou	t sciatica (range of scores: 0-	100; Better indica	ted by higher val	ues)						
	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	208	240	-	not pooled	⊕OOO VERY LOW	CRITICAL
SF-36; stra	atum = withou	t sciatica -	Physical function;	outcome ≤4 mon	ths (range of sco	res: 0-100; Better i	ndicated by h	igher va	lues)			
	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	13	15	-	MD 8.57 higher (6.78 lower to 23.92 higher)	⊕OOO VERY LOW	CRITICAL
SF-36; stra	atum = withou	t sciatica -	Social functionic;	outcome ≤4 mont	hs (range of scor	es: 0-100; Better in	dicated by hi	gher val	ues)			
	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	13	15	-	MD 9.17 higher (0.08 lower to 18.42 higher)	⊕⊕OO LOW	CRITICAL
SF-36; stra	atum = withou	t sciatica -	Physical role limit	ation; outcome ≤4	months (range o	of scores: 0-100; Be	etter indicated	d by high	ner value	s)		
	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	13	15	ı	MD 3.00 higher (25.48 lower to 31.48 higher)	⊕OOO VERY LOW	CRITICAL
SF-36; stra	ntum = withou	t sciatica -	Emotional role lim	itation; outcome :	≤4 months (range	of scores: 0-100; I	Better indicat	ed by hi	gher valu	ies)		
1	randomised	very	no serious	no serious	no serious	none	13	15	-	MD 35.06 higher (15.13 to	⊕⊕00	CRITICAL

	trials	serious ^a	inconsistency	indirectness	imprecision					54.99 higher)	LOW	
SF-36; st	tratum = withou	ıt sciatica -	Mental health; out	tcome ≤4 months	(range of scores:	0-100; Better indic	ated by highe	r values	s)			
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	13	15	-	MD 1.09 higher (3.26 lower to 5.44 higher)	⊕⊕OO LOW	CRITICAL
SF-36; st	tratum = withou	ıt sciatica -	Vitality; outcome	≤4 months (range	of scores: 0-100;	Better indicated b	y higher value	es)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	13	15	-	MD 7.64 higher (0.58 to 14.7 higher)	⊕⊕OO LOW	CRITICAL
SF-36; st	tratum = withou	ıt sciatica -	Bodily pain; outco	ome ≤4 months (ra	inge of scores: 0	-100; Better indicat	ed by higher	values)				•
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	13	15	-	MD 6.07 higher (2.76 lower to 14.9 higher)	⊕000 VERY LOW	CRITICAL
SF-36; st	tratum = withou	ıt sciatica -	General health pe	rception; outcome	e ≤4 months (rang	ge of scores: 0-100	; Better indica	ited by I	nigher va	lues)		
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	13	15	-	MD 13.72 higher (3.74 to 23.7 higher)	⊕⊕OO LOW	CRITICAL
Pain VAS	S; stratum = wit	hout sciati	ca; outcome ≤4 mo	onths (range of sc	ores: 0-10; Bette	r indicated by lowe	r values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	13	15	-	MD 0.81 lower (2.29 lower to 0.67 higher)	⊕000 VERY LOW	CRITICAL
Function	ı; stratum = witl	hout sciation	ca; outcome ≤4 mo	onths (range of sc	ores: 0-50; Better	indicated by lower	r values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	13	15	-	MD 2.93 lower (6.84 lower to 0.98 higher)	⊕OOO VERY LOW	CRITICAL

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias (b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 223: PENS versus TENS for low back pain with or without sciatica

Quality assessment	No of patients	Effect	Quality	Importance
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No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	PENS versus TENS	Control	Relative (95% CI)	Absolute				
Pain VAS;	Pain VAS; stratum +/- sciatica; outcome ≤4 months (range of scores: 0-10; Better indicated by lower values)													
1		- ,		no serious indirectness	serious ^b	none	53	49	-	MD 0.2 higher (0.65 lower to 1.05 higher)	⊕OOO VERY LOW	CRITICAL		
Function;	Function; stratum +/- sciatica; outcome ≤4 months (range of scores: 0-100; Better indicated by lower values)													
1		- ,		no serious indirectness	serious ^b	none	53	49	-	MD 2.47 lower (8.36 lower to 3.42 higher)	⊕OOO VERY LOW	CRITICAL		

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias (b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 224: Interferential therapy versus placebo/sham for low back pain without sciatica

Quality assessment							No of patients	Effect				
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Interferential therapy versus placebo/sham Control		Relative (95% CI)		Quality	Importance
Back pain	n NRS cm; str	atum = witho	ut sciatica (Better	indicated by low	ver values)							
2				no serious indirectness	no serious imprecision	none	59	58	-	MD 0.85 lower (1.14 to 0.56 lower)	⊕⊕⊕⊕ HIGH	CRITICAL

Table 225: Interferential versus traction for low back pain with or without sciatica

	able 225. Interferential versus traction for low back pain with or without sciutica													
Quality assessment							No of patient	ts		Effect	Quality	Importance		
No of	Design	Risk of	Inconsistency	Indirectness	Imprecision	Other	Interferential	Control	Relative	Absolute				

studies	bias considerations versus traction (95% CI)											
Function; outcome ≤4 months (Better indicated by lower values)												
1		very serious ^a			no serious imprecision	none	61	67	-	MD 0.6 lower (5.68 lower to 4.48 higher)	⊕⊕OO LOW	CRITICAL

Low back pain and sciatica in over 16s Quality assessment

Table 226: Laser versus sham for low back pain with sciatica

Quality assessment						No of patients			Effect	Quality	Immortonoo		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Laser versus sham	Control	Relative (95% CI)	Absolute	Quality	Importance	
Back pain; stratum = with sciatica - final score; outcome at ≤4 months (range of scores: 0-10; Better indicated by lower values)													
2	randomised trials	serious ^a	serious ^c		no serious imprecision	none	40	40	-	MD 0.35 higher (0.28 lower to 0.98 higher)	⊕⊕OO LOW	CRITICAL	
Back pair	Back pain; stratum = with sciatica - change score; outcome at ≤4 months (range of scores: 0-10; Better indicated by lower values)												
1	randomised trials				no serious imprecision	none	182	182	-	MD 1.43 lower (1.56 to 1.3 lower)	⊕⊕⊕O MODERATE	CRITICAL	
Function;	; stratum = wi	th sciatica; o	utcome at ≤4 mor	iths (range of sc	ores: 0-24; Bett	er indicated by lov	ver values)	-					
2	randomised trials		no serious inconsistency	no serious indirectness	serious ^b	none	40	40	1	MD 1.14 lower (3.31 lower to 1.04 higher)	⊕⊕OO LOW	CRITICAL	
Responde	er (Function i	mprovement)	; stratum = with s	ciatica; outcome	e at ≤4 months								
1	randomised trials		no serious inconsistency		no serious imprecision	none	151/182 (83%)	98/182 (53.8%)	RR 1.54 (1.33 to 1.79)	291 more per 1000 (from 178 more to 425 more)	⊕⊕⊕⊕ HIGH	IMPORTANT	

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias (b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.
(c) Downgraded by 1 or 2 increments because heterogeneity, I2=50%, p=0.04, unexplained by subgroup analysis

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 227: Laser versus sham for low back pain without sciatica

			Quality as	sessment			No of pat	tients		Effect	Ovality	l
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Laser versus sham	Control	Relative (95% CI)	Absolute	Quality	Importance
Back pain	; stratum = w	ithout scia	ntica; outcome ≤4 ı	nonths (range of	scores: 0-10; B	etter indicated by I	ower values)				
2	randomised trials	serious ^a	serious ^c	no serious indirectness	no serious imprecision	none	29	28	-	SMD 0.80 lower (1.73 lower to 0.12 higher)	⊕⊕OO LOW	CRITICAL
Back pain	(max pain in	last 24hrs); stratum = witho	ut sciatica; outco	me ≤4 months (range of scores: 0	-10; Better in	dicated	by lower value	es)		
1	randomised trials	serious ^a	no serious inconsistency ^c	no serious indirectness	serious ^b	none	30	31	-	MD 1.6 lower (2.8 to 0.37 lower)	⊕⊕OO LOW	CRITICAL
Responde	er (pain impro	vement >6	60%): stratum = wi	hout sciatica - C	hronic low back	pain; outcome ≤4	months					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	27/38 (71.1%)	12/33 (36.4%)		345 more per 1000 (from 69 more to 804 more)	⊕OOO VERY LOW	IMPORTANT
Function	(RMDQ/ODI);	stratum =	without sciatica; o	utcome ≤4 mont	hs (range of sco	res: 0-0-100; Bette	r indicated b	y lower	values)			
2	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	29	28	-	SMD 0.62 lower (2.55 lower to 1.32 higher)	⊕000 VERY LOW	CRITICAL
Function	(ODI)= withou	t sciatica	< 4 months (Better	indicated by low	ver values)							
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	31	30	-	MD 8.2 lower (13.6 to 2.8 lower)	⊕⊕OO LOW	CRITICAL

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias (b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.
(c) Downgraded by 1 or 2 increments because heterogeneity, I2=50%, p=0.04, unexplained by subgroup analysis

Table 228: Laser versus usual care for low back pain with sciatica

				•										
Quality as	ssessment						No of patie	ents		Effect				
											Quality	Importance		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Laser versus usual care	Control	Relative (95% CI)	Absolute				
Back pain	ack pain; stratum = with sciatica; outcome at ≤4 months (range of scores: 0-10; Better indicated by lower values)													
		no serious risk of bias	no serious inconsistency		no serious imprecision	none	182	182	-	MD 0.92 lower (1.05 to 0.78 lower)	⊕⊕⊕⊕ HIGH	CRITICAL		
Function	Function improvement; stratum = with sciatica; outcome at ≤4 months													
		no serious risk of bias	no serious inconsistency		no serious imprecision	none	151/182 (83%)	33/182 (18.1%)	RR 4.58 (3.34 to 6.27)	649 more per 1000 (from 424 more to 956 more)	⊕⊕⊕⊕ HIGH	IMPORTANT		

Table 229: Laser versus usual care for low back pain with or without sciatica

Quality ass	uality assessment									Effect	Quality	Importance		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Laser versus usual care	Control	Relative (95% CI)	Absolute	Quanty	Importance		
Pain VAS;	ain VAS; stratum: +/- sciatica; outcome ≤4 months (follow-up ≤4 months; range of scores: 0-10; Better indicated by lower values)													
		,			no serious imprecision	none	75	75	-	MD 1.26 lower (1.74 to 0.78 lower)	⊕⊕OO LOW	CRITICAL		
Roland Dis	Roland Disability Questionnaire; stratum: +/- sciatica; outcome ≤4 months (range of scores: 0-24; Better indicated by lower values)													
	trials	serious ^a	inconsistency	indirectness	serious ^b	none	25	25	-	MD 0.8 higher (1.06 lower to 2.66 higher)	⊕OOO VERY LOW	CRITICAL		

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias (b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 230: Laser versus exercise for low back pain with or without sciatica

			and for four back	P		-								
			Quality as	sessment			No of patie	ents		Effect				
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Laser versus exercise	Control	Relative (95% CI)		Quality	Importance		
Pain VAS;	ain VAS; stratum: +/- sciatica; outcome ≤4 months (range of scores: 0-10; Better indicated by lower values)													
1	randomised trials	very serious ^a		no serious indirectness	no serious imprecision	none	25	25	-	MD 1 lower (1.75 to 0.25 lower)	⊕⊕OO LOW	CRITICAL		
Roland Dis	sability Questi	onnaire; s	tratum: +/- sciatica;	outcome ≤4 mon	ths (range of sco	res: 0-24; Better in	dicated by lowe	r values))					
1	randomised trials	very serious ^a		no serious indirectness	no serious imprecision	none	25	25	-	MD 1.1 higher (0.59 lower to 2.79 higher)	⊕⊕OO LOW	CRITICAL		

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 231: Laser versus traction for low back pain with sciatica

			Quality asses	ssment			No of patie	ents		Effect	Quality	Importance		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Laser versus traction	Control	Relative (95% CI)	Absolute	Quanty	importance		
Back pain;	ack pain; stratum = with sciatica; outcome ≤4 months (range of scores: 0-10; Better indicated by lower values)													
1		,	no serious inconsistency	no serious indirectness	serious ^b	none	20	20	-	MD 0.13 lower (1.16 lower to 0.9 higher)	⊕OOO VERY LOW	CRITICAL		
Radicular _I	pain; stratum =	with sciat	ica; outcome ≤4 mo	onths (range of sc	ores: 0-10; B	etter indicated by I	lower values)							
1		- ,	no serious inconsistency	no serious indirectness	serious ^b	none	20	20	-	MD 0.59 lower (1.66 lower to 0.48 higher)	⊕OOO VERY LOW	CRITICAL		

Function; stratum = with sciatica; outcome ≤4 months (range of scores: 0-24; Better indicated by lower values)														
1 randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	20	20	-	MD 2.2 lower (4.84 lower to 0.44 higher)	⊕000 VERY LOW	CRITICAL			

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 232: Ultrasound versus placebo/sham for low back pain with sciatica

			Quality asse	essment			No of patients			Effect	Ovelity	Importance		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Ultrasound versus placebo/sham	Control	Relative (95% CI)	Absolute	Quanty	Importance		
Back pain	Back pain (VAS cm); stratum = with sciatica; outcome at ≤4 months (range of scores: 0-10; Better indicated by lower values)													
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	15	15	1	MD 0.06 lower (2.1 lower to 1.98 higher)	⊕000 VERY LOW	CRITICAL		
Function;	stratum = wit	h sciatica;	outcome at ≤4 mo	onths (range of s	cores: 0-100;	Better indicated b	by lower values)							
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	15	15	-	MD 3.86 higher (2.48 lower to 10.2 higher)	⊕⊕OO LOW	CRITICAL		
Paracetan	nol use; stratu	ım = with	sciatica; outcome	at ≤4 months (Be	tter indicate	d by lower values)								
1	trials		no serious inconsistency	no serious indirectness	serious ^b	none	15	15	-	MD 7.67 lower (21.37 lower to 6.03 higher)	LOW	IMPORTANT		

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 233: Ultrasound versus placebo/sham for low back pain without sciatica

Quality assessment	No of patients	Effect	Quality	Importance

⁽b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

⁽b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Ultrasound versus placebo/sham	Control	Relative (95% CI)	Absolute		
Back pair	n (VAS cm); s	stratum = wi	thout sciatica; ou	tcome at ≤4 mo	nths (range of	scores: 0-10; Bet	ter indicated by lowe	er values	s)			
	randomised trials	serious ^a		no serious indirectness	serious ^b	none	21	18	-	MD 0.22 higher (0.55 lower to 0.99 higher)	⊕⊕OO LOW	CRITICAL
Moderate	(>30%) pain	reduction;	stratum = without	sciatica; outco	me ≤4 months							
		no serious risk of bias		no serious indirectness	serious ^b	none	128/233 (54.9%)		RR 1.02 (0.86 to 1.2)	11 more per 1000 (from 76 fewer to 108 more)	⊕⊕⊕O MODERATE	IMPORTANT
Function	; stratum = w	ithout sciati	ica; outcome at ≤	4 months (range	e of scores: 0-1	00; Better indicat	ed by lower values)					
		very serious ^a			no serious imprecision	none	26	23	-	MD 7.46 lower (13.54 to 1.38 lower)	⊕⊕OO LOW	CRITICAL

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias (b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs

Table 234: Ultrasound versus usual care (both groups had exercise) for low back pain without sciatica

				20 til Bi 0 tilpo i		тот то та жазат р	in without sciatica						
			Quality ass	sessment			No of patients			Effect	.		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Ultrasound versus usual care (both groups had exercise)	Control	Relative (95% CI)	Absolute	Quality	Importance	
SF-36; str	F-36; stratum = without sciatica - Physical function domain; outcome ≤4 months (range of scores: 0-100; Better indicated by higher values)												
		- ,	no serious inconsistency	no serious indirectness	serious ^b	none	20	20		MD 2.75 lower (9.72 lower to 4.22 higher)	⊕OOO VERY LOW	CRITICAL	
SF-36; str	F-36; stratum = without sciatica - Mental health domain; outcome ≤4 months (range of scores: 0-100; Better indicated by higher values)												

-											•	
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	20	20	-	MD 0.7 lower (7.64 lower to 6.24 higher)	⊕OOO VERY LOW	CRITICAL
SF-36; st	ratum = witho	out sciatic	a - Pain domain; o	outcome ≤4 mor	nths (range of so	cores: 0-100; Bette	r indicated by higher values	s)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	20	20		MD 0.25 lower (7.67 lower to 7.17 higher)	⊕⊕OO LOW	CRITICAL
SF-36; st	ratum = witho	out sciatic	a - General health	n domain; outcoi	me ≤4 months (ı	range of scores: 0-	-100; Better indicated by hig	gher val	ues)			
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	20	20	-	MD 5.75 lower (15.34 lower to 3.84 higher)	⊕OOO VERY LOW	CRITICAL
SF-36; st	ratum = witho	out sciatic	a - Social function	n domain; outco	me ≤4 months (range of scores: 0	-10; Better indicated by hig	her valu	es)			
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	20	20	-	MD 1.75 lower (9.54 lower to 6.04 higher)		CRITICAL
SF-36; st	ratum = witho	out sciatic	a - Physical role l	imitation domaii	n; outcome ≤4 n	nonths (range of s	cores: 0-100; Better indicate	ed by hi	gher val	ues)		
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	20	20	-	MD 6 higher (1.55 lower to 13.55 higher)	⊕⊕OO LOW	CRITICAL
SF-36; st	ratum = witho	out sciatic	a - Emotional role	limitation doma	ain; outcome ≤4	months (range of	scores: 0-100; Better indica	ated by	nigher va	alues)		
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	20	20	-	MD 7 higher (2.2 lower to 16.2 higher)	⊕OOO VERY LOW	CRITICAL
SF-36; st	ratum = witho	out sciatic	a - Energy domai	n; outcome ≤4 m	nonths (range of	f scores: 0-100; Be	tter indicated by higher val	ues)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	20	20	-	MD 3.5 lower (11.53 lower to 4.53 higher)		CRITICAL
Pain; stra	atum = withou	ıt sciatica	; outcome ≤4 mor	nths (range of so	cores: 0-10; Bett	ter indicated by lov	ver values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	20	20	-	MD 1.7 lower (2.57 to 0.83 lower)	⊕⊕OO LOW	CRITICAL

Function;	unction; stratum = without sciatica; outcome ≤4 months (range of scores: 0-50; Better indicated by lower values)													
1		- ,			no serious imprecision	none	20	20	-	MD 0.6 lower (2.8 lower to 1.6 higher)	⊕⊕OO LOW	CRITICAL		
Depression	Depression; stratum = without sciatica; outcome ≤4 months (range of scores: 0-63; Better indicated by lower values)													
1		- ,			no serious imprecision	none	20	20		MD 0.75 lower (3.01 lower to 1.51 higher)		CRITICAL		

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias (b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 235: Ultrasound versus laser for low back pain with or without sciatica

			Quality asses	sment			No of pati	ents		Effect	Quality	
No of studies						Other considerations	Ultrasound		Relative (95% CI)	Absolute	Quality	Importance
Back pain;	stratum +/- sci	atica (rang	e of scores: 0-10; Be	tter indicated by lo	ower values)							
		,		no serious indirectness	serious ^b	none	27	35	-	MD 0.37 lower (1.53 lower to 0.79 higher)	⊕OOO VERY LOW	CRITICAL

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias (b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 236: Ultrasound versus traction for low back pain with sciatica

			Quality ass	sessment			No of patien	its		Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Ultrasound versus traction	Control	Relative (95% CI)	Absolute	Quality	Importance
Back pain;	stratum = wi	th sciatica	; outcome ≤4 mont	hs (range of sco	res: 0-10; Better	indicated by lower	values)					

1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	20	20	-	MD 0.44 lower (1.42 lower to 0.54 higher)	⊕OOO VERY LOW	CRITICAL
Function	RMDQ SMD; s	tratum = v	vith sciatica; outco	ome ≤4 months (ra	ange of scores: (0-24; Better indicate	ed by lower values)				
1	randomised trials	very serious ^a	no serious inconsistency		no serious imprecision	none	20	20		MD 0.3 lower (3.46 lower to 2.86 higher)	⊕⊕OO LOW	CRITICAL

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias (b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Combinations of interventions – electrotherapy adjunct J.10.1

Low back pain with sciatica J.10.1.1

Table 237: Electrotherapy (ultrasound) plus exercise (biomechanical plus aerobics) compared with waiting list control

			Quality asse	essment			No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency			Exercise (biomechanical + aerobics) + ultrasound	waiting list control	Relative (95% CI)	Absolute	Quanty	importance	
Back Pair	n (VAS 0-10) -	≤4 month	ns (follow-up 3 we	eks; measured	with: VAS 0-	10; range of score	es: 0-10; Better indicated by	lower value	es)			
	randomised trials			no serious indirectness	serious ^b	none	15	15	-	MD 2.6 lower (4.27 to 0.93 lower)	⊕⊕OO LOW	CRITICAL
Leg Pain	(VAS 0-10) - ≤	4 months	s (follow-up 3 wee	ks; measured w	ith: VAS 0-1	0; range of scores	: 0-10; Better indicated by I	ower values	s)			
	randomised trials			no serious indirectness	serious ^b	none	15	15	-	MD 2 lower (3.73 to 0.27 lower)	⊕⊕OO LOW	CRITICAL
Function	Function (ODI 0-100) - ≤4 months (follow-up 3 weeks; measured with: ODI; range of scores: 0-100; Better indicated by lower values)											
	randomised trials	serious ^a			very serious ^b	none	15	15	-	MD 0.34 lower (7.27 lower to 6.59 higher)	⊕OOO VERY LOW	CRITICAL

Medicatio	on use - ≤4 mo	onths (foll	low-up 3 weeks; r	neasured with: I	Paracetamol	intake; Better indi	cated by lower values)					
1	randomised trials			no serious indirectness	serious ^b	none	15	15	1	MD 22.27 lower (38.26 to 6.28 lower)	⊕⊕OO LOW	IMPORTANT

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias (b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 238: Electrotherapy (ultrasound) plus exercise (biomechanical plus aerobics) compared with exercise (biomechanical plus aerobics)

			Quality ass	essment			No of pat	ients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Ultrasound + exercise (biomechanical + aerobics)	exercise (biomechanical + aerobics)	Relative (95% CI)	Absolute	Quanty	Importance
Back Pai	n (VAS 0-10)	- ≤4 mon	ths (follow-up 3	weeks; measur	ed with: VAS	0-10; range of so	cores: 0-10; Better indica	ited by lower values))			
	randomised trials		no serious inconsistency	no serious indirectness	very serious ^b	none	15	15	-	MD 0.26 lower (2.3 lower to 1.78 higher)	⊕OOO VERY LOW	CRITICAL
Leg Pain	(VAS 0-10) -	≤4 montl	hs (follow-up 3 w	eeks; measure	d with: VAS (0-10; range of sco	ores: 0-10; Better indicat	ed by lower values)				
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	15	15	-	MD 1 higher (1.44 lower to 3.44 higher)	⊕OOO VERY LOW	CRITICAL
Function	(ODI 0-100)	- ≤4 mon	ths (follow-up 3	weeks; measur	ed with: Osw	vestry disability ir	ndex 0-100; range of sco	res: 0-100; Better inc	licated b	y lower values)		
	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	15	15	-	MD 3.86 higher (2.48 lower to 10.2 higher)	⊕⊕OO LOW	CRITICAL
Medication	on use - ≤4 m	nonths (fo	ollow-up 3 weeks	; measured wit	h: Use of par	acetamol; Better	indicated by lower value	es)				
	randomised trials		no serious inconsistency	no serious indirectness	serious ^b	none	15	15	-	MD 7.67 lower (21.37 lower to 6.03 higher)	LOW	IMPORTANT

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Low back pain without sciatica

Table 239: Electrotherapy (laser) plus self-management (education) plus exercise (biomechanical) compared with self-management (education) plus exercise (biomechanical)

		(1010111	cenameary									
			Quality ass	essment			No of pa	atients		Effect		
No of studies	Design	Risk of bias	I Inconsistancy I Ind		Imprecision	Other considerations	Laser + education + exercise (biomechanical)	education + exercise (biomechanical)	Relative (95% CI)		Quality	Importance
Pain (0-1	0 VAS) - <4 m	onths (fo	ollow-up 3 weeks	; measured with	: VAS; range	e of scores: 0-10;	Better indicated by lowe	r values)				
	randomised trials			no serious indirectness	serious ^b	none	50	50	-	MD 1.64 lower (2.42 to 0.86 lower)	⊕⊕OO LOW	CRITICAL

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

(b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 240: Electrotherapy (TENS) plus acupuncture compared with acupuncture

			Quality asse	ssment			No of pa	tients		Effect	Ovelite	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	TENS + acupuncture	acupuncture	Relative (95% CI)	Absolute	Quanty	importance
² ain (0-100	VAS conver	ted to 0-10) - <4 months (folio	ow-up 10 weeks;	measured wi	th: VAS; range of	scores: 0-10; Bet	tter indicated	by lower	values)		
		- ,		no serious indirectness	very serious ^b	none	6	7	-	MD 0.59 higher (1.48 lower to 2.66 higher)	⊕000 VERY LOW	CRITICAL

⁽b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

		- ,		and the second s	very serious ^b	none	6	7	ı	MD 0.2 lower (3.98 lower to 3.58 higher)	⊕OOO VERY LOW	CRITICAL
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⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias (b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 241: Electrotherapy (TENS) plus exercise (biomechanical) compared with sham TENS

			Quality as	sessment			No of patients			Effect	Ouglitu	
No of studies					Other considerations	TENS + exercise (biomechanical)	sham TENS	Relative (95% CI)		Quanty	Importance	
Pain (Borç	y verbal pain r	ating scal	e 0-10) - <4 month	s (follow-up 8 we	eks; measured w	vith: VRS; range of	scores: 0-10; Better ind	icated b	y lower v	alues)		
1	randomised trials	- ,	no serious inconsistency		no serious imprecision	none	21	21	1	MD 0.66 lower (0.7 to 0.62 lower)	⊕⊕OO LOW	CRITICAL
Function (Function (ODI 0-100) - <4 months (follow-up 8 weeks; measured with: ODI; range of scores: 0-100; Better indicated by lower values)											
1	randomised trials	- ,	no serious inconsistency		no serious imprecision	none	21	21	-	MD 7.6 lower (8.77 to 6.43 lower)	⊕⊕OO LOW	CRITICAL

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 242: Electrotherapy (TENS) plus exercise (biomechanical) compared with exercise (biomechanical)

			Quality asse	essment			No of pa	tients		Effect	0			
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	TENS + exercise (biomechanical)	exercise (biomechanical)	Relative (95% Absolute CI)		Quality	Importance		
SF-36 (0-	SF-36 (0-100) - <4 months: Mental health (follow-up 6 weeks; measured with: SF-36; range of scores: 0-100; Better indicated by higher values)													
1	randomised	very	no serious	no serious	serious ^b	none	20	20	-	MD 6.95 higher	⊕000	CRITICAL		

	L		l	h	1	1				(0.44)	\ /EB\ /	1
	trials	serious ^a	inconsistency	indirectness						(0.44 lower to 14.34 higher)	VERY LOW	
										14.34 nigner)	LOVV	
F-36 (0-	100) - <4 mor	ths: Gene	eral health (follow	v-up 6 weeks; m	easured with	n: SF-36; range of	scores: 0-100; Better	indicated by higher	values)			T
	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	20	20	-	MD 6.15 higher (5.3 lower to 17.6 higher)	⊕OOO VERY LOW	CRITICAL
SF-36 (0-	100) - <4 mor	nths: Ener	gy (follow-up 6 v	veeks; measured	d with: SF-36	; range of scores:	0-100; Better indicate	d by higher values)			
	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	20	20	-	MD 16.05 higher (7.72 to 24.38 higher)	⊕000 VERY LOW	CRITICAL
Pain (Bo	rg and PDI -ce	onverted t	to 0-10) - <4 mon	ths (range of sc	ores: 0-10; B	etter indicated by	lower values)					
?	randomised trials	very serious ^a	very serious ^c	no serious indirectness	very serious ^b	none	41	43	-	MD 0.15 higher (0.54 lower to 0.85 higher)	⊕OOO VERY LOW	CRITICAL
unction	(ODI 0-100)	- <4 mont	hs (measured wi	th: ODI; range o	f scores: 0-1	00; Better indicate	ed by lower values)					
!	randomised trials	very serious ^a	very serious ^c	no serious indirectness	very serious ^b	none	41	43	-	MD 2.63 higher (5.61 lower to 4.86 higher)	⊕000 VERY LOW	CRITICAL
Psycholo	ogical distres	s: Beck D	epression Invent	ory (0-63) - <4 m	onths (follow	v-up 6 weeks; mea	asured with: BDI; rang	je of scores: 0-63; E	Better inc	dicated by lower va	lues)	
	randomised trials	- ,	no serious inconsistency	no serious indirectness	serious ^b	none	20	20	-	MD 1.5 lower (3.68 lower to 0.68 higher)	⊕000 VERY LOW	CRITICAL

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias.
(b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.
(c) Downgraded by 1 increment for I2 >50% - 74% and 2 increments for I2 >75%.

Table 243: Electrotherapy (PENS) plus exercise (biomechanical plus aerobics) compared with sham PENS plus exercise (biomechanical plus aerobics)

Quality assessment	No of patients	Effect	Quality	Importance
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No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	PENS + exercise (biomechanical + aerobics)	sham PENS + exercise (biomechanical + aerobics)	Relative (95% CI)	Absolute		
SF-36 (0	-100) - <4 mo	nths: Me	ental component	summary scor	e (follow-up 6	weeks; measure	d with: SF-36; range o	of scores: 0-100; Bette	r indicate	d by higher va	ues)	
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	45	44	-	MD 3.1 lower (8.34 lower to 2.14 higher)	⊕⊕OO LOW	CRITICAL
SF-36 (0	-100) - >4 mo	nths: Me	ental component	summary scor	e (follow-up 6	months; measur	ed with: SF-36; range	of scores: 0-100; Bett	er indicat	ted by higher v	alues)	
1	randomised trials	seriousª	no serious inconsistency	no serious indirectness	serious ^b	none	45	44	-	MD 1.7 lower (7.44 lower to 4.04 higher)	⊕⊕OO LOW	CRITICAL
SF-36 (0	-100) - <4 mo	nths: Ph	ysical compone	nt summary sc	ore (follow-up	6 weeks; measu	red with: SF-36; range	e of scores: 0-100; Bet	ter indica	ted by higher v	alues)	
1	randomised trials	seriousª	no serious inconsistency	no serious indirectness	serious ^b	none	45	44	-	MD 3 lower (13.09 lower to 7.09 higher)	⊕⊕OO LOW	CRITICAL
SF-36 (0	-100) - >4 mo	nths: Ph	ysical compone	nt summary sc	ore (follow-up	6 months; meas	ured with: SF-36; rang	ge of scores: 0-100; Be	etter indic	ated by higher	values)	
1	randomised trials	seriousª	no serious inconsistency	no serious indirectness	serious ^b	none	45	44	-	MD 4.1 lower (15.06 lower to 6.86 higher)	⊕⊕OO LOW	CRITICAL
Pain (Mo	:Gill) - <4 mo	nths (foll	ow-up 6 weeks;	measured with	ı: McGill; rang	e of scores: 0-78;	Better indicated by l	ower values)				
1	randomised trials	seriousª		no serious indirectness	serious ^b	none	45	44	-	MD 1 lower (4.34 lower to 2.34 higher)	⊕⊕OO LOW	CRITICAL
Pain (Mo	:Gill) - >4 mo	nths (foll	ow-up 6 months	; measured wit	th: McGill; ran	ge of scores: 0-7	8; Better indicated by	lower values)				
1	randomised trials	seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	45	44	-	MD 0.7 lower (4.04 lower to 2.64 higher)	⊕⊕⊕O MODERATE	CRITICAL
Function	n (RMDQ) - <4	4 months	(follow-up 6 we	eks; measured	with: RMDQ;	range of scores:	0-24; Better indicated	by lower values)				
1	randomised	seriousª	no serious	no serious	no serious	none	45	44	-	MD 0.4 higher	⊕⊕⊕О	CRITICAL

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	trials		inconsistency	indirectness	imprecision					(1.53 lower to 2.33 higher)	MODERATE	
Function	(RMDQ) - >4	months	(follow-up 6 mo	nths; measure	d with: RMDQ	range of scores	: 0-24; Better indicate	d by lower values)				
	randomised trials			no serious indirectness	serious ^b	none	45	44	-	MD 0.7 higher (1.31 lower to 2.71 higher)	⊕⊕OO LOW	CRITICAL

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias (b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 244: Electrotherapy (ultrasound) plus exercise compared with exercise (biomechanical)

	Quality assessment							f patients		Effect	Ouality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Ultrasound + exercise	exercise (biomechanical)	Relative (95% CI)	Absolute	Quanty	importance
SF-36 (0-1	00) - <4 mont	ths: Menta	al health (follow-up	6 weeks; meas	ured with: SI	F-36; range of sco	res: 0-100; Bette	er indicated by high	er values	s)		
1 -		,		no serious indirectness	very serious ^b	none	19	20	-	MD 1.3 higher (6.09 lower to 8.69 higher)	⊕OOO VERY LOW	CRITICAL
SF-36 (0-1	00) - <4 mont	ths: Gene	ral health (follow-u	ıp 6 weeks; mea	sured with: §	SF-36; range of sc	ores: 0-100; Bet	ter indicated by hig	her value	es)		
		- ,		no serious indirectness	very serious ^b	none	19	20	-	MD 1.27 higher (9.07 lower to 11.61 higher)	⊕OOO VERY LOW	CRITICAL
SF-36 (0-1	00) - <4 mont	ths: Energ	y (follow-up 6 wee	eks; measured w	/ith: SF-36; ra	ange of scores: 0-	100; Better indi	cated by higher valu	ies)			
1		,		no serious indirectness	very serious ^b	none	19	20	-	MD 0.93 higher (8.36 lower to 10.22 higher)	⊕OOO VERY LOW	CRITICAL
Pain (pain	n (pain disabiltiy index 0-50) - <4 months (follow-up 6 weeks; range of scores: 0-50; Bo						icated by lower	values)				
1		- ,		no serious indirectness	very serious ^b	none	19	20	-	MD 0.29 lower (3.07 lower to 2.49 higher)	⊕OOO VERY	CRITICAL

											LOW	
Function	(ODI 0-100) -	<4 month	ıs (follow-up 6 wee	eks; measured w	ith: ODI; ran	ge of scores: 0-10	0; Better indicat	ed by lower values)				
1		- ,		no serious indirectness	very serious ^b	none	19	20	-	MD 0.28 higher (2.03 lower to 2.59 higher)	⊕OOO VERY LOW	CRITICAL
Depression (Beck Depression Inventory (0-63)) - <4 months (follow-up 6 weeks; measured with: BDI; range of scores: 0-63; Better indicated by lower values)												
1		- ,		no serious indirectness	serious ^b	none	19	20	-	MD 0.91 lower (3.05 lower to 1.23 higher)	⊕OOO VERY LOW	CRITICAL

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 245: Electrotherapy (ultrasound) plus exercise plus self-management compared with exercise plus self-management

			Quality asse	essment			No of patients Effect				Quality	Immontono
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Ultrasound + exercise + self- management	exercise + self- management	Relative (95% CI)	Absolute	Quality	Importance
Pain (0-10	00 VAS conve	erted to 0-	·10) - <4 months (follow-up 2 mon	ths; measur	ed with: VAS; ran	ge of scores: 0-10; Bet	ter indicated by lo	wer valu	es)		
1		,	no serious inconsistency	no serious indirectness	serious ^b	none	21	18	-	MD 0.22 higher (0.55 lower to 0.99 higher)	⊕OOO VERY LOW	CRITICAL
Function	(Functional F	Rating Ind	ex) - <4 months (follow-up 2 mon	ths; range o	f scores: 0-40; Be	tter indicated by lower	values)				
1	trials	seriousª	,	indirectness		none	21	18	-	MD 7.7 lower (14.13 to 1.27 lower)	⊕OOO VERY LOW	CRITICAL

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

⁽b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

⁽b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

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Table 246: Electroacupuncture plus self-management (mixed modality - education + home exercise) plus exercise compared with self-management (mixed modality - education + home exercise) plus exercise

			Quality as:	sessment			No of patients Effect			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Electroacupuncture + education + exercise + home exercise	education + exercise + home exercise	Relative (95% CI)	Absolute	Quanty	importance
Pain (NR	S 0-10) - <4 ı	months (Better indicated	by lower value	s)							
		very serious ^a	no serious inconsistency		no serious imprecision	none	24	25	-	MD 1.81 lower (3.07 to 0.55 lower)	⊕⊕OO LOW	CRITICAL
Function	(Aberdeen I	ow back	pain scale 0-100	cvonverted to	0-10 scale) - •	<4 months (Better	r indicated by lower values)					
		- ,	no serious inconsistency	no serious indirectness	serious ^b	none	24	25	-	MD 0.6 lower (1.25 lower to 0.06 higher)	⊕OOO VERY LOW	CRITICAL
Analgesi	c consumpti	ion - <4 n	nonths									
	randomised trials	- ,	no serious inconsistency	no serious indirectness	very serious ^b	none	2/26 (7.7%)	4/26 (15.4%)	RR 0.5 (0.1 to 2.5)	77 fewer per 1000 (from 138 fewer to 231 more)	⊕OOO VERY LOW	IMPORTANT
							raded by 2 increments if the n	15.4%		77 fewer per 1000 (from 139 fewer to 231 more)		

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias (b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 247: Electrotherapy (Interferential) plus manual therapy (manipulation) compared with manual therapy (manipulation)

									, (
	Quality assessment						No of patients Effect			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Interferential + manipulation	manipulation	Relative (95% CI)	Absolute	Quality	importance
Quality o	f life (EQ-5D)	- <4 months	s (Better indicated	d by lower value	s)							
1	randomised trials	seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	66	63	-	MD 0.01 lower (0.15 lower to 0.13 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality o	f life (EQ-5D)	- >4 months	(Better indicated	by lower value	s)							
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	51	52	-	MD 0.1 higher (0.01 lower to 0.21 higher)	⊕⊕OO LOW	CRITICAL
SF-36 (0-	100) - <4 mor	nths: Physic	al functioning (Be	etter indicated b	y lower values)							
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	66	63	-	MD 0.95 lower (8.27 lower to 6.37 higher)	⊕000 VERY LOW	CRITICAL
SF-36 (0-	100) - >4 mor	nths: Physic	al functioning (Be	etter indicated b	y lower values)							
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	51	52	-	MD 12.04 higher (2.6 to 21.48 higher)	⊕⊕⊕O MODERATE	CRITICAL
SF-36 (0-	100) - <4 mor	nths: Role pl	hysical (Better inc	dicated by lower	r values)							
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	66	63	-	MD 1.43 higher (12.96 lower to 15.82 higher)	⊕OOO VERY LOW	CRITICAL
SF-36 (0-	100) - >4 mor	nths: Role pl	hysical (Better inc	dicated by lower	r values)							
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	51	52	-	MD 12.2 higher (5.48 lower to 29.88	⊕000 VERY LOW	CRITICAL

										higher)		
										nigher)		
SF-36 (0-	100) - <4 mor	ths: Bodily	pain (Better indic	ated by lower v	alues)							
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	66	63	-	MD 0.69 lower (8.86 lower to 7.48 higher)	⊕OOO VERY LOW	CRITICAL
SF-36 (0-	100) - >4 mor	nths: Bodily	pain (Better indic	ated by lower v	alues)							
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	51	52	-	MD 12.59 higher (2.65 to 22.53 higher)	⊕⊕⊕O MODERATE	CRITICAL
SF-36 (0-	100) - <4 mor	nths: Genera	al health (Better in	ndicated by lowe	er values)							
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	66	63	-	MD 2.27 higher (3.56 lower to 8.1 higher)	⊕OOO VERY LOW	CRITICAL
SF-36 (0-	100) - >4 mor	nths: Genera	al health (Better ir	ndicated by lowe	er values)							
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	51	52	-	MD 3.27 higher (4.58 lower to 11.12 higher)	⊕OOO VERY LOW	CRITICAL
SF-36 (0-	100) - <4 mor	nths: Vitality	(Better indicated	by lower values	s)							
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	66	63	-	MD 0.96 lower (7.64 lower to 5.72 higher)	⊕000 VERY LOW	CRITICAL
SF-36 (0-	100) - >4 mor	nths: Vitality	(Better indicated	by lower values	s)							
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	51	52	-	MD 5.17 higher (2.93 lower to 13.27 higher)	⊕OOO VERY LOW	CRITICAL
SF-36 (0-	100) - <4 mor	nths: Social	functioning (Bette	er indicated by I	ower values)							
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	very serious ^b	none	66	63	-	MD 0.17 lower (9.05 lower to 8.71 higher)	⊕⊕OO LOW	CRITICAL

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SF-36 (0-	100) - >4 mor	ths: Social	functioning (Bett	er indicated by	lower values)							
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	51	52	-	MD 0.2 lower (13.99 lower to 13.59 higher)	⊕OOO VERY LOW	CRITICAL
SF-36 (0-	100) - <4 mor	iths: Role e	motional (Better i	ndicated by low	er values)							
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	66	63	1	MD 11.85 higher (3.38 lower to 27.08 higher)	⊕⊕OO LOW	CRITICAL
SF-36 (0-	100) - >4 mor	iths: Role e	motional (Better i	ndicated by low	er values)							
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	very serious ²	none	51	52	-	MD 8.2 higher (7.21 lower to 23.61 higher)	⊕OOO VERY LOW	CRITICAL
SF-36 (0-	100) - <4 mor	ıths: Mental	health domain (E	Better indicated	by lower values	3)				<u>, </u>		
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	66	63	1	MD 2.46 higher (3.06 lower to 7.98 higher)	⊕OOO VERY LOW	CRITICAL
SF-36 (0-	100) - >4 mor	iths: Mental	health domain (E	Better indicated	by lower values	· s)						
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	51	52	-	MD 5.58 higher (1.53 lower to 12.69 higher)	⊕⊕OO LOW	CRITICAL
Pain (0-10	00 VAS conve	erted to 0-10	0) - <4 months (Be	etter indicated b	y lower values)							
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	66	63	-	MD 0.48 lower (1.35 lower to 0.39 higher)	⊕⊕OO LOW	CRITICAL
Pain (0-10	00 VAS conve	erted to 0-10	0) - >4 months (Be	etter indicated b	y lower values)							
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	51	52	1	MD 0.75 lower (1.81 lower to 0.31 higher)	⊕⊕OO LOW	CRITICAL
Function	(RMDQ) - <4	months (Be	etter indicated by	lower values)								

1		randomised trials	seriousª	no serious inconsistency		no serious imprecision	none	66	63	-	MD 0.12 lower (1.78 lower to 1.54 higher)	⊕⊕⊕O MODERATE	CRITICAL
F	unction	(RMDQ) - >4	months (Be	tter indicated by	lower values)								
1		randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	51	52	1	MD 1.79 lower (3.77 lower to 0.19 higher)	⊕⊕OO LOW	CRITICAL
(8	a) Down	graded by 1 ir	ncrement if th	ne majority of the e	vidence was at h	igh risk of bias, a	and downgraded by	y 2 increments if the	majority of the	e evidenc	e was at very high ris	sk of bias	

(b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 248: Electrotherapy (laser) plus self-management (home exercise) compared with self-management (home exercise)

			Quality as	sessment			No of pat	ients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Laser + home exercise	home exercise	Relative (95% CI)	Absolute	Quanty	Importance
Pain (VAS	0-10) - <4 mon	ths (Better	r indicated by I	ower values)								
	randomised trials	very serious ^a	very serious ^c	no serious indirectness	serious ^b	none	44	43	-	MD 0.99 lower (2.85 lower to 0.87 higher)	⊕OOO VERY LOW	CRITICAL
Function (ODI 0-100) - <	4 months (Better indicate	d by lower values)							
	randomised trials	very serious ^a	very serious ^c	indirectness	very serious ^b	none	44	43	-	MD 4.00 lower (11.23 lower to 3.23 higher)	⊕OOO VERY LOW	CRITICAL

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias.
(b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

⁽c) Downgraded by 1 increment for I2 >50% - 74% and 2 increments for I2 >75%.

Table 249: Electrotherapy (HILT Laser) + self-management (unsupervised exercise) compared to placebo HILT laser + self-management (unsupervised exercise)

			Quality as	sessment			No of patients		ı	Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	HILT laser + self-management (unsupervised exercise) compared to placebo HILT laser + self-management (unsupervised exercise) for low back pain		Relative (95% CI)		Quality	Importance
Pain seve	erity (VAS, 0	-10) ≤ 4 n	nonths (follow-u	p 12 weeks; Be	etter indicated	by lower values)						
		,		no serious indirectness	serious ^b	none	28	24	-	MD 1.07 lower (1.77 to 0.37 lower)	⊕OOO VERY LOW	CRITICAL
Function	(RMDQ, 0-2	4) ≤ 4 mo	nths (follow-up	12 weeks; Bette	er indicated by	/ lower values)						
		-		no serious indirectness	serious ^b	none	28	24	-	MD 1.42 lower (1.95 to 0.89 lower)	⊕OOO VERY LOW	CRITICAL
Function	(MODQ, 0-1	00) ≤ 4 m	onths (follow-up	12 weeks; Bet	tter indicated I	by lower values)						
	randomised trials	,		no serious indirectness	no serious imprecision	none	28	24	-	MD 3.61 lower (5.62 to 1.6 lower)	⊕⊕OO LOW	CRITICAL

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias (b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 250: Electrotherapy (BEMER + TENS) + exercise + manual therapy (massage) compared to placebo BEMER + TENS + exercise + manual therapy (massage)

(IIIassa ₈	5 -1												
			Quality ass	essment			No of patients			Effect			
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Othor	BEMER + TENS+ exercise + manual therapy (massage) vs placebo BEMER + TENS + manual therapy (massage)	Control	Relative (95% CI)	Absolute	Quality	Importance	
Quality o	ality of life (SF-36 Physical functioning, 0-100) ≤ 4 months (follow-up 15 weeks; Better indicated by lower values)												
1		,		no serious indirectness	very serious ^b	none	13	13	-	MD 0.15 lower (3.95 lower to 3.65 higher)	⊕OOO VERY LOW	CRITICAL	
Quality o	Quality of life (SF-36 Role physical, 0-100) ≤ 4 months (follow-up 15 weeks; Better indicated by lower values)												
1		very serious ^a		no serious indirectness	serious ^b	none	14	14	-	MD 5.63 lower (13.72 lower to 2.46 higher)	⊕OOO VERY LOW	CRITICAL	
Quality o	f life (SF-36 I	Bodily pai	in, 0-100) ≤ 4 mor	nths (follow-up	15 weeks; Be	etter indicated by	lower values)						
1		very serious ^a		no serious indirectness	serious ^b	none	15	18	-	MD 4.01 lower (8.86 lower to 0.84 higher)	⊕OOO VERY LOW	CRITICAL	
Quality o	f life (SF-36 (General h	ealth, 0-100) ≤ 4 r	months (follow-	up 15 weeks	; Better indicated	by lower values)						
1	randomised trials	very serious ^a			very serious ^b	none	12	14	-	MD 1.40 lower (5.18 lower to 2.38 higher)	⊕OOO VERY LOW	CRITICAL	
Quality o	f life (SF-36 \	/itality, 0-	-100) ≤ 4 months	(follow-up 15 w	eeks; Better	indicated by low	er values)						
1		very serious ^a		no serious indirectness	serious ^b	none	10	12	-	MD 5.6 lower (11.13 to 0.07 lower)	⊕OOO VERY LOW	CRITICAL	
Quality o	f life (SF-36	Social fun	ctioning, 0-100) :	≤ 4 months (foll	ow-up 15 we	eks; Better indica	ated by lower values)						

								•	1			
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	13	18	-	MD 0.98 lower (8.25 lower to 6.29 higher)	⊕000 VERY LOW	CRITICAL
Quality o	of life (SF-36	Role emo	tional, 0-100) ≤ 4	months (follow	-up 15 week	s; Better indicated	d by lower values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	13	15	-	MD 3.5 lower (16.38 lower to 9.38 higher)	⊕000 VERY LOW	CRITICAL
Quality o	of life (SF-36 I	Mental he	ealth, 0-100) ≤ 4 m	nonths (follow-u	ıp 15 weeks;	Better indicated I	by lower values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	9	15	-	MD 0.52 lower (6.71 lower to 5.67 higher)	⊕000 VERY LOW	CRITICAL
Quality o	of life (SF-36	Physical o	component sumn	nary score, 0-10	00) ≤ 4 montl	ns (follow-up 15 w	veeks; Better indicated by lower value	es)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	6	10	-	MD 0.93 lower (6.38 lower to 4.52 higher)	⊕000 VERY LOW	CRITICAL
Quality o	of life (SF-36 l	Mental co	mponent summa	ry score, 0-100) ≤ 4 months	(follow-up 15 we	eks; Better indicated by lower values)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	6	10	-	MD 8.66 lower (15.29 to 2.03 lower)	⊕000 VERY LOW	CRITICAL
Pain sev	erity (exercis	e VAS, 0-	-10) ≤ 4 months (f	follow-up 15 we	eks; Better i	ndicated by lower	values)					
1	randomised trials		no serious inconsistency	no serious indirectness	very serious ^b	none	18	19	-	MD 0.42 higher (0.99 lower to 1.83 higher)	⊕OOO VERY LOW	CRITICAL
Pain sev	erity (resting	VAS, 0-1	0) ≤ 4 months (fo	llow-up 15 wee	ks; Better in	dicated by lower v	/alues)					
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	18	19	-	MD 0.72 higher (0.6 lower to 2.04 higher)	⊕000 VERY LOW	CRITICAL
Function	(ODI, 0-100)	≤ 4 mont	hs (follow-up 15	weeks; Better i	ndicated by l	ower values)						
1	randomised	seriousª	no serious	no serious	very	none	18	19	-	MD 1.19 higher	⊕000	CRITICAL

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trials	inconsistency	indirectness	serious ^b			(7.02 lower to	VERY	
						9.40 higher)	LOW	

- (a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias
- (b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

J.11 Psychological interventions

Table 251: Cognitive behavioural approaches versus placebo/sham in low back pain with or without sciatica

			Quality as:	sessment			No of patients			Effect	Ouality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	cognitive behavioural approaches versus placebo/sham	Control	Relative (95% CI)		Quanty	importance
Pain seve	rity - >4 mont	ths (Bette	r indicated by low	er values)								
		very serious ^a	no serious inconsistency		no serious imprecision	none	59	59	-	MD 0.90 higher (3.6 lower to 5.41 higher)	⊕⊕OO LOW	CRITICAL
Function	(ODI, 0-100) >	4 months	(range of scores	: 0-100; Better in	dicated by lowe	er values)						
		very seriousª	no serious inconsistency		no serious imprecision	none	59	59	-	MD 0.7 higher (4.81 lower to 6.21 higher)	⊕⊕OO LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 252: Cognitive behavioural approaches versus usual care/waiting list in low back pain with or without sciatica

	cogt.		rounal apploa		suu. cu. c, iii		v back pain with or with	 			
			Quality ass	sessment		No of patients		Effect	Quality	l	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	cognitive behavioural approaches versus usual care/waiting list	Relative (95% CI)	Absolute	Quality	Importance

Pain sev	erity (VAS, 0-	10 final va	alue) <4 months (range of scores	: 0-10; Better in	idicated by lower	values)					
6	randomised trials	very serious ^a	serious ²	no serious indirectness	no serious imprecision	none	231	227	-	MD 0.66 lower (1.01 to 0.31 lower)	⊕000 VERY LOW	CRITICAL
Pain (VA	AS, 0-10) <4 m	onths (rar	nge of scores: 0-1	0; Better indica	ted by lower va	ılues)						
1	randomised trials	very serious ^a	no serious inconsistency ^b	no serious indirectness	no serious imprecision	none	27	27	-	MD 2.59 lower (3.28 to 1.9 lower)	⊕⊕OO LOW	CRITICAL
Function	n (RMDQ, 0-24	l) <4 mont	ths (range of scor	es: 0-24; Better	indicated by lo	wer values)						
2	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	121	119	-	MD 2.95 lower (4.26 to 1.65 lower)	⊕⊕OO LOW	CRITICAL
Function	n (PDI, 0-70) <	4 months	(range of scores	0-70; Better inc	dicated by lowe	er values)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^c	none	53	50	-	MD 1.20 lower (6.44 lower to 4.04 higher)	⊕000 VERY LOW	CRITICAL
Psychol	ogical distres	s (BDI, 0-0	68)<4 months (rai	nge of scores: 0	-68; Better indi	cated by lower va	lues)					
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^c	none	58	51	-	MD 1.65 lower (3.42 lower to 0.12 higher)	⊕⊕OO LOW	CRITICAL
Quality	of life (SF-36 p	perceived	general health, 0	-5) < 4 months (range of scores	s: 0-5; Better indic	ated by higher values)					
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	143	171	-	MD 0 higher (0.18 lower to 0.18 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality of	of life (SF-36 p	perceived	general health, 0	-5) >4 months (r	ange of scores	: 0-5; Better indica	ated by higher values)					
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	143	171	-	MD 0 higher (0.19 lower to 0.19 higher)	⊕⊕⊕O MODERATE	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by one increment because of heterogeneity, I2 >50% ^c Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

Table 253: Cognitive behavioural approaches versus behavioural therapy in low back pain with or without sciatica

						- /			1			
			Quality as	sessment			No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	cognitive behavioural approaches versus behavioural therapy	Control	Relative (95% CI)	Absolute	Quanty	importance
Pain sev	erity (VAS 0-1	00 conver	ted to 0-10) <4 m	onths (range of	scores: 0-10; B	etter indicated by	lower values)					
1		very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	41	36	-	MD 0.4 lower (1.03 lower to 0.96 higher)	⊕⊕OO LOW	CRITICAL
Pain sev	erity (VAS 0-1	00 conver	ted to 0-10) >4 m	onths (range of	scores: 0-10; B	etter indicated by	lower values)					
1		very serious ^a	no serious inconsistency		no serious imprecision	none	38	35	-	MD 0.07 higher (0.95 lower to 1.09 higher)	⊕⊕OO LOW	CRITICAL
Function	(RMDQ, 0-24)) >4 month	ns (range of score	es: 0-24; Better i	ndicated by low	ver values)		•				
1		very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	38	35	1	MD 2.94 lower (12.17 lower to 6.29 higher)	⊕000 VERY LOW	CRITICAL
Function (RMDQ, 0-24) >4 months (range of scores: 0-24; Better indicated by lower values)												
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	38	35	-	MD 2.11 lower (4.71 lower to 0.49 higher)	⊕⊕OO LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

Table 254: Behavioural therapy versus placebo/sham in low back pain with or without sciatica

			Quality asse	ssment		No of patients			Effect	Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Behavioural therapy versus placebo	Control	Relative		Quanty	importance
Pain severity (VAS, 0-10) <4 months (range of scores: 0-10; Better indicated by lower values)												
	randomised trials			no serious indirectness	Serious ^b	none	16	8	-	MD 1.44 lower (2.88 lower to 0 higher)	⊕⊕OO LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

Table 255: Behavioural therapy versus usual care/waiting list in low back pain with or without sciatica

			Quality asse	essment			No of patients			Effect	Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Behavioural therapy versus usual care/waiting list		Relative (95% CI)	Absolute	Quanty	importance	
Pain seve	rity (Back pai	n log) <4	months (Better inc	dicated by lower	values)								
1		- ,	no serious inconsistency		very serious ^b	none	10	10	-	MD 4.80 lower (15.84 lower to 6.24 higher)	⊕OOO VERY LOW	CRITICAL	
Pain seve	rity (McGill Pa	ain questi	onnaire, 0-78) <4 ı	months (range o	f scores: 0-7	8; Better indicated	l by lower values)						
2		,		no serious indirectness	Serious ^b	none	65	57	1	mean 3.42 lower (8.08 lower to 1.24 higher)	⊕OOO VERY LOW	CRITICAL	
Function	nction (Modified activity form score) >4 months (Better indicated by lower values)												

b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

Table 256: Mindfulness versus usual care/waiting list in low back pain with or without sciatica

			Quality asse	ssment			No of patients			Effect	Qualita	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Mindfulness versus UC/waiting list	Control	Relative (95% CI)	Absolute	Quality	Importance
Pain sever	rity (McGill pa	in questio	nnaire, 0-78) <4 mo	onths (range of s	cores: 0-78; E	Better indicated by	lower values)					

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

2	randomised trials	very serious ^a	very serious ^b	no serious indirectness	Serious ^c	none	58	66	-	MD 5.55 lower (11.7 lower to 0.08 higher)	⊕OOO VERY LOW	CRITICAL
Function	(RMDQ, 0-24)	<4 months	(range of scores:	0-24; Better indic	cated by low	er values)						
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^c	none	19	18	-	MD 1.20 lower (4.55 lower to 2.15 higher)	⊕⊕OO LOW	CRITICAL
Quality of	life (SF-36 glo	bal health	composite, 0-100	<4 months (range)	ge of scores:	0-100; Better indic	cated by higher values)					
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^c	none	19	18	-	MD 1.8 higher (4.56 lower to 8.16 higher)	⊕⊕OO LOW	CRITICAL
Quality of	life (SF-36 me	ental healtl	h composite, 0-100) <4 months (ran	ge of scores	: 0-100; Better indi	cated by higher values)					
2	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^c	none	58	66	-	MD 4.74 higher (2.87 to 6.62 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	life (SF-36 pa	in scale, 0	-100) <4 months (r	ange of scores: 0)-100; Better	indicated by highe	r values)					
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	very serious ^c	none	19	18	-	MD 1.1 higher (4.07 lower to 6.27 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	life (SF-36 ph	ysical fund	ction scale, 0-100)	<4 months (range	e of scores:	0-100; Better indica	ated by higher values)					
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	very serious ^c	none	19	18	-	MD 1.2 higher (5.04 lower to 7.44 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	life (SF-36 ph	ysical hea	Ith composite, 0-1	00) <4 months (ra	inge of score	es: 0-100; Better in	dicated by higher value	s)				
2	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^c	none	58	66	-	MD 3.69 higher (2.59 to 4.8 higher)	⊕OOO VERY LOW	CRITICAL

Low back pain and sciatica in over 16s Quality assessment

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 2 increments because of heterogeneity, I2=75%, p=0.05, unexplained by subgroup analysis ^c Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

Table 257: Cognitive therapy versus usual care/waiting list in low back pain without sciatica

			Quality as	sessment			No of	patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Cognitive versus	Usual care/ waiting list	Relative (95% CI)	Absolute	Quanty	importance
Quality of	life (SF-36 ph	ysical fun	ction, 0-100) >4 m	onths (range of s	cores: 0-100; Be	tter indicated by h	igher values)					
1	randomised trials	very seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	34	29	-	MD 6.7 higher (2.01 lower to 15.41 higher)	⊕000 VERY LOW	CRITICAL
Quality of	life (SF-36 ro	le function	n, 0-100) >4 month	s (range of score	s: 0-100; Better i	ndicated by higher	r values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	34	29	-	MD 9.1 higher (57.12 lower to 75.32 higher)	⊕000 VERY LOW	CRITICAL
Quality of	life (SF-36 bo	dily pain,	0-100) >4 months	(range of scores:	0-100; Better in	dicated by higher v	/alues)	•	•			
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	34	29	-	MD 8.9 higher (2.63 lower to 20.43 higher)	⊕000 VERY LOW	CRITICAL
Quality of	life (SF-36 ge	neral heal	th, 0-100) >4 mont	hs (range of scor	res: 0-100; Bette	r indicated by high	er values)	'	,		<u>'</u>	
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	34	29	-	MD 5 higher (1.12 lower to 11.12 higher)	⊕000 VERY LOW	CRITICAL
Quality of	life (SF-36 vit	ality, 0-10	0) >4 months (rang	ge of scores: 0-10	00; Better indicat	ted by higher value	es)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	34	29	-	MD 12.6 higher (2.44 to 22.76 higher)	⊕000 VERY LOW	CRITICAL
Quality of	life (SF-36 so	cial functi	on, 0-100) >4 mon	ths (range of sco	res: 0-100; Bette	r indicated by high	ner values)		•			
1	randomised	very	no serious	no serious	Serious ^b	none	34	29	-	MD 1.9 higher (9.43	⊕OOO	CRITICAL

1	L			h 11 /	1	1	l		ı		\((= 0) \((
	trials	serious ^a	inconsistency	indirectness						lower to 13.23 higher)	VERY				
			ļ	l .							LOW				
Quality of	life (SF-36 ro	le emotion	nal, 0-100) >4 mont	hs (range of scor	es: 0-100; Bette	r indicated by high	er values)								
*	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	34	29	-	MD 14 higher (7.44 lower to 35.44 higher)	⊕OOO VERY LOW	CRITICAL			
Quality of	uality of life (SF-36 mental health, 0-100) >4 months (range of scores: 0-100; Better indicated by higher values)														
	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	34	29	-	MD 6.8 higher (0.7 lower to 14.3 higher)	⊕OOO VERY LOW	CRITICAL			
Quality of	Quality of life (SF-36 health transition, 0-100) >4 months (range of scores: 0-100; Better indicated by higher values)														
	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	34	29	-	MD 5.6 higher (13.43 lower to 24.63 higher)	⊕OOO VERY LOW	CRITICAL			
Pain seve	rity (VAS 0-10	0 converte	ed to 0-10) <4 mon	ths (range of sco	res: 0-10; Better	indicated by lowe	r values)								
	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	34	29	-	MD 1.09 lower (2.202 lower to 0.22 higher)	⊕OOO VERY LOW	CRITICAL			
Function ((RMDQ, 0-24)	>4 months	s (range of scores	0-24; Better indi	cated by lower v	alues)									
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	34	29	-	MD 1.9 lower (3.84 lower to 0.04 higher)	⊕⊕OO LOW	CRITICAL			

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

Table 258: Cognitive therapy versus usual care/waiting list in low back pain with or without sciatica

	Quality assessment							ients		Effect	Quality	Importance
No of studies			Indirectness	Imprecision	Other considerations	Cognitive tp	UC/WL	Relative (95%	Absolute	-		

									01)					
									CI)					
Pain sever	ity (VAS 0-100	converted	to 0-10) <4 months (range of scores: 0	-10; Better ir	ndicated by lower va	lues)							
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	16	18	-	MD -1.12 lower (2.51 lower to 0.28 higher)	⊕OOO VERY LOW	CRITICAL		
Psycholog	Psychological distress (BDI, 0-63) <4 months (range of scores: 0-63; Better indicated by lower values)													
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	16	18	-	MD 1.53 higher (2.63 lower to 5.69 higher)	⊕OOO VERY LOW	CRITICAL		
Function (Sickness impa	ct profile, 0	0-68) <4 months (ran	ge of scores: 0-68;	Better indic	ated by lower value	s)							
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	16	18	-	MD 1.69 lower (7.34 lower to 3.96 higher)	⊕OOO VERY LOW	CRITICAL		

Low back pain and sciatica in over 16s Quality assessment

Table 259: Cognitive therapy versus exercise (biomechanical plus aerobics) in low back pain without sciatica

	Quality assessment							ients		Effect	Over life.	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Cognitive therapy	Exercise	Relative (95% CI)	Absolute	Quality	Importance
Quality of	life (SF-36 phy	sical funct	ion, 0-100) >4 mont	hs (range of score	tter indicated by hi	gher values)						
1	randomised trials	,		no serious indirectness	Serious ^b	none	34	30	-	MD 6.2 higher (2.51 lower to 14.91 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	life (SF-36 role	function,	0-100) >4 months (r	ange of scores: 0	-100; Better i	ndicated by higher	values)					
1	randomised trials	,		no serious indirectness	Serious ^b	none	34	30	-	MD 3.6 lower (26.21 lower to 19.01 higher)	⊕OOO VERY	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

											LOW	
Quality of	life (SF-36 Box	dily nain 0	-100) >4 months (ra	nge of scores: 0-1	00: Better in	ndicated by higher v	(alues)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	34	30	-	MD 6.8 higher (4.4 lower to 18 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	life (SF-36 gen	eral health	n, 0-100) >4 months	(range of scores:	0-100; Bette	r indicated by high	er values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	34	30	-	MD 1.2 higher (5.45 lower to 7.85 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	life (SF-36 vita	lity 0-100)	>4 months (range of	of scores: 0-100; B	etter indicat	ed by higher values	s)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	34	30	-	MD 12.5 higher (4.02 to 20.98 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	life (SF-36 soc	ial function	n, 0-100) >4 months	(range of scores:	0-100; Bette	er indicated by high	er values)					
1	randomised trials	very seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	34	30	-	MD 3.1 higher (8.47 lower to 14.67 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	life (SF-36 role	emotiona	I, 0-100) >4 months	(range of scores:	0-100; Bette	er indicated by high	er values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	34	30	-	MD 6.6 higher (16.58 lower to 29.78 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	life (SF-36 me	ntal health,	, 0-100) >4 months	(range of scores: (0-100; Better	r indicated by highe	r values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	34	30	-	MD 7.7 higher (1.01 to 14.39 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	life (SF-36 hea	lth transiti	on, 0-100) >4 montl	ns (range of score	s: 0-100; Be	tter indicated by hig	gher values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	34	30	-	MD 2.6 higher (17.36 lower to 22.56 higher)	⊕OOO VERY LOW	CRITICAL

Low back pain and sciatica in over 16s Quality assessment

Pain sever	ain severity (VAS 0-100, converted to 0-10) >4 months (range of scores: 0-10; Better indicated by lower values)													
1	randomised trials	- ,		no serious indirectness	Serious ^b	none	34	30	-	MD 0.6 lower (1.76 lower to 0.56 higher)	⊕OOO VERY LOW	CRITICAL		
Function (I	Function (RMDQ, 0-24) >4 months (range of scores: 0-24; Better indicated by lower values)													
1	randomised trials	- ,		no serious indirectness	Serious ^b	none	34	30	-	MD 1.4 lower (3.34 lower to 0.54 higher)	⊕OOO VERY LOW	CRITICAL		

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by one increment if the confidence interval crossed one MID or by two increments if the confidence interval crossed both MIDs

Combinations of interventions – psychological adjunct J.11.1

Table 260: Psychological therapy (behavioural therapy) plus exercise (aerobic) compared with waiting list in low back pain without sciatica

			Quality asse	essment			No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Behavioural therapy + exercise (aerobic)	waiting list	Relative (95% CI)	Absolute	Quanty	importance
Pain (McG	Gill) - <4 month	ns (follow-	-up 8 weeks; meas	ured with: McGil	l; range of so	cores: 0-78; Better	indicated by lower valu	es)				
	randomised trials	- 3		no serious indirectness	Serious ^b	none	18	19	-	MD 6.17 lower (13.29 lower to 0.95 higher)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

Table 261: Psychological therapy (Behavioural therapy) plus exercise (aerobic) compared with exercise (aerobic) in low back pain without sciatica

Quality assessment	No of patients	Effect	Quality	Importance

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Behavioural therapy + exercise (aerobic)	exercise (aerobic)	Relative (95% CI)	Absolute			
Pain (Mc	Pain (McGill) - <4 months (follow-up 8 weeks; measured with: McGill; range of scores: 0-78; Better indicated by lower values)												
1		- 3		no serious indirectness	Serious ^b	none	18	21		MD 2.74 lower (9.59 lower to 4.11 higher)	⊕OOO VERY LOW	CRITICAL	

Low back pain and sciatica in over 16s Quality assessment

Table 262: Psychological intervention (cognitive behavioural approaches) plus exercise (mixed: biomechanical + aerobic) compared with exercise (mixed: biomechanical + aerobic) in low back pain with or without sciatica

	(IIIIACU. I	Jioineen	anicai + aerobi	cj ili low back	Pain With	or without solu						
Quality assessment							No of patients		Effect			
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	cognitive behavioural approaches + exercise	exercise	Relative (95% CI)		Quality	Importance
Pain (0-10	Pain (0-100 NRS converted to 0-10 scale) - <4 months (range of scores: 0-10; Better indicated by lower values)											
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	43	41	-	MD 0.71 lower (1.8 lower to 0.38 higher)	⊕⊕OO LOW	CRITICAL
Pain (0-10	0 NRS conver	ted to 0-1	0 scale) - >4 mont	ns (range of scor	es: 0-10; Bet	tter indicated by lo	wer values)					
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	34	35	-	MD 1.55 lower (2.78 to 0.32 lower)	⊕⊕OO LOW	CRITICAL
Function	Low back out	come sca	le questionnaire 0	-75 converted to	0-10) - <4 me	onths (range of sc	ores: 0-10; Better indicate	d by high	er value:	s)		
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	43	41	-	MD 0.83 higher (0.06 lower to 1.72 higher)	⊕⊕OO LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

Function (Low back outcome scale questionnaire 0-75 converted to 0-10) - >4 months (range of scores: 0-10; Better indicated by higher values)												
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	34	35	-	MD 1.06 higher (0.06 to 2.06 higher)	⊕⊕OO LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 263: Psychological intervention (cognitive behavioural approaches) plus self-management compared with self-management in low back pain with or without sciatica

			Quality as	sessment		No of patients		Effect		Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	cognitive behavioural approaches + self- management	self- management	Relative (95% CI)	Absolute	Quality	Importance
Pain (0-100 von Korff converted to 0-10 scale) - <4 months (range of scores: 0-10; Better indicated by lower values)												
1		,		no serious indirectness	no serious imprecision	none	355	190	-	MD 0.68 lower (1.06 to 0.3 lower)	⊕⊕OO LOW	CRITICAL
Pain (0-1	Pain (0-100 von Korff converted to 0-10 scale) - >4 months (range of scores: 0-10; Better indicated by lower values)											
1	randomised trials			no serious indirectness	no serious imprecision	none	399	199	-	MD 0.7 lower (1.12 to 0.28 lower)	⊕⊕⊕O MODERATE	CRITICAL
Function	Function (RMDQ, 0-24) <4 months (range of scores: 0-24; Better indicated by lower values)											
1		very serious ^a		no serious indirectness	no serious imprecision	none	355	190	-	MD 0.9 lower (1.63 to 0.17 lower)	⊕⊕OO LOW	CRITICAL
Function (RMDQ 0-24) >4 months (range of scores: 0-24; Better indicated by lower values)												
1	randomised trials			no serious indirectness	no serious imprecision	none	399	199	-	MD 1.3 lower (2.12 to 0.48 lower)	⊕⊕⊕O MODERATE	CRITICAL

Function	(0-100 von K	orff scale	converted to 0-	10) - <4 months	(range of scor	es: 0-10; Better in	ndicated by lower values)				
	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	355	190	-	MD 0.43 lower (0.85 to 0.01 lower)	⊕⊕OO LOW	CRITICAL
unction	(0-100 von K	orff scale	converted to 0-	10) - >4 months	(range of scor	es: 0-10; Better in	dicated by lower values)				
	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	399	199	-	MD 0.84 lower (1.26 to 0.42 lower)	⊕⊕⊕O MODERATE	CRITICAL
Quality o	f life (EQ-5D,	0-1) <4 m	nonths (range of	scores: 0-1; Be	tter indicated b	y higher values)						
	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	349	179	-	MD 0.06 higher (0.01 to 0.11 higher)	⊕⊕OO LOW	CRITICAL
Quality o	f life (EQ-5D,	0-1) >4 m	nonths (range of	scores: 0-1; Be	tter indicated b	y higher values)						
	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	327	163	-	MD 0.05 higher (0.02 to 0.09 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality o	f life (SF-12 p	ohysical c	component, 0-100	0) <4 months (ra	ange of scores:	0-100; Better ind	icated by higher values)					
	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	332	176	-	MD 0.6 higher (1.47 lower to 2.67 higher)	⊕⊕OO LOW	CRITICAL
Quality o	f life (SF-12 p	ohysical c	component, 0-100	0) >4 months (ra	ange of scores:	0-100; Better ind	icated by higher values)					
	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	375	187	-	MD 0.6 lower (2.6 lower to 1.4 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality o	f life (SF-12 r	nental co	mponent, 0-100)	<4 months (ran	ge of scores: ()-100; Better indic	ated by higher values)					
		very serious ^a	no serious inconsistency	no serious indirectness	No serious imprecision	none	332	176	-	MD 1.6 higher (0.34 lower to 3.54 higher)	⊕⊕OO LOW	CRITICAL
Quality o	f life (SF-12 r	nental co	mponent, 0-100)	>4 months (ran	ge of scores: (-100; Better indic	ated by higher values)					

		randomised trials				no serious imprecision	none	375	187	1	MD 3.3 higher (1.29 lower to 5.31 higher)	⊕⊕⊕O MODERATE	CRITICAL
,	a Downgra	aded by 1 incr	ement if the	ne maiority of the	evidence was at	high risk of bias	. and downgraded	by 2 increments if the mai	ority of the evid	lence was	s at verv high risk o	of bias	

Pharmacological interventions

Antidepressants versus placebo J.12.1

Table 264: Tricyclic antidepressants versus placebo (low back pain with/without sciatica population)

Quality a	ssessment						No. of patients		Effect			
No. of studies		Risk of bias	Inconsistency	Indirectness	Imprecision	Other	Tricyclic antidepressants versus placebo		Relative (95% CI)	Absolute	Quality	Importance
Pain sev	erity (follow	/-up ≤4 m	onths; measure	ed with: (DSS 0	0-21 and VAS	0-10); Better inc	dicated by lower val	ues)				
2	Randomised trials	Serious ^a	No serious inconsistency	No serious indirectness	No serious imprecision	None	57	59	-	SMD 0.24 higher (0.13 lower to 0.6 higher)	MODERATE	CRITICAL
Psycholo	ogical distres	s (follow	/-up ≤4 months;	measured wit	th: BDI; range	of scores: 0-63;	Better indicated by	lower	values)			
2	Randomised trials	Serious ^a	No serious inconsistency	No serious indirectness	No serious imprecision	None	59	59	-	MD 1.75 higher (0.05 lower to 3.56 higher)	MODERATE	CRITICAL
Psycholo	ogical distres	s (follow	r-up ≤4 months;	measured wit	th: STAI; range	e of scores: 20-8	30; Better indicated	by lowe	er values)	,	•	,
1	Randomised trials	Very serious ^b	No serious inconsistency	No serious indirectness	Serious ^c	None	38	40	-	MD 2.59 higher (1.28 lower to 6.46 higher)	VERY LOW	CRITICAL
Adverse	events (follo	ow-up ≤4	l months)		•	•	•	-				•
1	Randomised	Serious ^a	No serious	No serious	Serious ^c	None	28/41	29/40	RR 1.02	14 more per 1000	LOW	IMPORTANT

b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

	trials		inconsistency	indirectness		(68.3%)	` '		(from 160 fewer to 239 more)		
Health	care utilisatio	n (follow	-up ≤4 months)								
1	Randomised trials			No serious indirectness	No serious imprecision	· ·	,	(0.44 to	206 fewer per 1000 (from 115 fewer to 268 fewer)	MODERATE	IMPORTANT

- (a) Downgraded by one increment if the majority of the evidence was at high risk of bias
- (b) Downgraded by two increments if the majority of the evidence was at very high risk of bias
- (c) Downgraded by one increment if the confidence interval crossed one MID

Table 265: SSRIs versus placebo (low back pain only and low back pain with/without sciatica population)

			Quality as	sessment			No of pat	ients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	SSRIs versus placebo	Control	Relative (95% CI)	Absolute		
Pain seve	rity (low back	pain pop	ulation) (follow-up	<4 months; mea	asured with: DS	S; range of scores	s: 0-63; Better	rindicate	ed by lower va	alues)		
		,	no serious inconsistency	no serious indirectness	Serious ^b	none	31	22	-	MD 0.90 higher (0.63 lower to 2.43 higher)	VERY LOW	CRITICAL
Pain seve	rity (low back	pain with	/without sciatica	population) (follo	ow-up median <	4 months; Better ii	ndicated by lo	ower valu	ues)			
	randomised trials	Serious ^c			no serious imprecision	none	78	84	-	SMD 0.05 higher (0.26 lower to 0.36 higher)	MODERATE	CRITICAL
Disability	(ODI) (follow-	up <4 mo	nths; range of sco	pres: 0-100; Bette	er indicated by l	ower values)						
	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	44	48	-	MD 2.2 lower (8.11 lower to 3.71 higher)	LOW	CRITICAL
Psycholog	gical distress	, MADRS	(follow-up <4 mon	ths; range of sco	ores: 20-80; Bet	ter indicated by lo	wer values)	1		1		
1	randomised	Serious ^c	no serious	no serious	no serious	none	44	48	-	MD 0.1 lower (3.64	MODERATE	IMPORTANT

	trials		inconsistency	indirectness	imprecision					lower to 3.44 higher)		
Adverse e	vents (low ba	ack pain p	opulation) (follow	-up <4 months)								
		,	no serious inconsistency	no serious indirectness	Serious ^b	none	16/43 (37.2%)	3/26 (11.5%)	RR 3.22 (1.04 to 10.01)	256 more per 1000 (from 5 more to 1000 more)	VERY LOW	IMPORTANT
Adverse e	vents (low ba	ack pain w	rith/without sciatti	ca population) (1	follow-up <4 mo	nths)						
	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	20/22 (90.9%)	31/32 (96.9%)	RR 0.94 (0.81 to 1.09)	58 fewer per 1000 (from 184 fewer to 87 more)	MODERATE	IMPORTANT

Table 266: SNRIs versus placebo (low back pain with/without sciatica)

			Quality as	sessment			No of pat	ients		Effect	Quality	Importanc
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	SNRIs versus placebo	Control	Relative (95% CI)	Absolute		P
ain seve	erity (follow-u	p <4 mont	hs; Better indicate	ed by lower value	es)							
3	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	576	428	-	MD 0.7 lower (0.99 to 0.4 lower)	MODERATE	CRITICAL
unction	(mean change	e) - BPI-I (0-10) (follow-up <	4 months; Better	indicated by lo	wer values)						
	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	575	427	-	MD 0.66 lower (0.91 to 0.41 lower)	MODERATE	CRITICAL

⁽a) Downgraded by two increments if the majority of the evidence was at very high risk of bias

⁽b) Downgraded by one increment if the confidence interval crossed one MID

⁽c) Downgraded by one increment if the majority of the evidence was at high risk of bias

2	randomised	Serious ^a	no serious	no serious	Serious ^b	none	172/310	145/320	RR 1.22	100 more per 1000	LOW	IMPORTANT
					Octions	none					LOVV	IIVII OIXIAIVI
	trials		inconsistency	indirectness			(55.5%)	(45.3%)	(1.05 to 1.43)	` .		
										more)		
EQ-5D (fo	llow-up <4 m	onths: rar	nge of scores: 0-1	: Better indicated	by lower value	s)						
(,	.9	,	,	-,						
2	randomised	Serious ^a	no serious	no serious	no serious	none	446	296	-	MD 0.05 higher (0.01 to	MODERATE	CRITICAL
	trials		inconsistency	indirectness	imprecision					0.09 higher)		
	uiuio		in conclotonoy	in an oothood	in prodictor					o.oo mgnor)		
Adverse e	events			L		l L						
3	randomised	Serious ^a	no serious	no serious	serious ²	none	243/600	87/441	RR 1.39	77 more per 1000 (from	LOW	IMPORTANT
	trials		inconsistency	indirectness			(40.5%)	(19.7%)	(1.17 to 1.65)	34 more to 128 more)		
							(1312,12)	(1011 /0/	(
Healthcar	e utilisation (follow-up	<4 months)	<u>'</u>				L				
	·		·									
1	randomised	Serious ^a	no serious	no serious	no serious	none	65/236	58/121	RR 0.57	206 fewer per 1000	MODERATE	IMPORTANT
	trials		inconsistency	indirectness	imprecision		(27.5%)	(47.9%)	(0.44 to 0.76)	(from 115 fewer to 268		
					,		(=:::://	(, . ,	(,	fewer)		
										iewei)		
L												

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias

Table 267: SNRIs versus placebo (low back with/without sciatica population)

			Quality as:	sessment			No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	SNRI (60 mg) versus placebo (low back pain +/- sciatica)		Relative (95% CI)	Absolute		
SF-36 (Du	F-36 (Duloxetine 60 mg) - Mental component (follow-up <4 months; range of scores: 0-100; Better indicated by higher values)											
	randomised trials				no serious imprecision	none	147	153	-	MD 2.25 higher (0.17 to 4.33 higher)	MODERATE	CRITICAL
SF-36 (Du	loxetine 60 m	ng) - Phys	ical component (f	follow-up <4 mo	nths; range of s	cores: 0-100; Bet	er indicated by higher va	alues)			I	

⁽b) Downgraded by 1 increment if the confidence interval crossed one MID

1												
	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	147	153	-	MD 1.24 higher (0.89 lower to 3.37 higher)	MODERATE	CRITICAL
SF-36 (D	uloxetine 60 n	ng) - Bodi	ly pain (follow-up	<4 months; ran	ge of scores: 0	-100; Better indica	ted by higher values)					
2	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	290	298	-	MD 0.66 higher (0.13 to 1.2 higher)	MODERATE	CRITICAL
SF-36 (D	uloxetine 60 n	ng) - Ment	al health (follow-	up <4 months; r	ange of scores	: 0-100; Better indi	cated by higher values)					
2	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	267	274	-	MD 1.02 higher (0.09 to 1.96 higher)	MODERATE	CRITICAL
SF-36 (D	uloxetine 60 n	ng) - Gene	eral health (follow	/-up <4 months;	range of score	s: 0-100; Better ind	icated by higher values)					
2	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	290	298	-	MD 0.69 higher (0.1 lower to 1.49 higher)	MODERATE	CRITICAL
SF-36 (D	uloxetine 60 n	ng) - Phys	ical functioning	follow-up <4 mc	onths; range of	scores: 0-100; Bet	ter indicated by higher va	lues)				
2	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	288	297	-	MD 0.53 higher (0.47 lower to 1.54 higher)	MODERATE	CRITICAL
SF-36 (D	uloxetine 60 n	ng) - Role	emotional (range	e of scores: 0-10	0; Better indica	ted by higher valu	es)	ļ				
2	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	274	287	-	MD 0.12 higher (0.13 lower to 0.37 higher)	MODERATE	CRITICAL
SF-36 (D	uloxetine 60 n	ng) - Role	-physical (follow-	up 2 months; ra	nge of scores:	0-100; Better indic	ated by higher values)					
2	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	274	287	-	MD 0.01 higher (0.4 lower to 0.43 higher)	MODERATE	CRITICAL

SF-36 (Du	loxetine 60 m	ng) - Socia	al functioning (fo	llow-up <4 mont	hs; range of sco	ores: 0-100; Better	indicated by lower value	es)				
2	randomised trials				no serious imprecision	none	290	298	-	MD 0.01 higher (0.42 lower to 0.44 higher)	MODERATE	CRITICAL
SF-36 (Du	loxetine 60 m	ng) - Vitali	ity (follow-up <4 r	nonths; range o	f scores: 0-100;	Better indicated b	y higher values)					
2	randomised trials				no serious imprecision	none	265	273	-	MD 0.75 higher (0.2 lower to 1.7 higher)	MODERATE	CRITICAL

⁽a) Downgraded by one increment if the majority of the evidence was at high risk of bias

Table 268: SNRIs versus placebo (low back pain with/without sciatica)

		-	Quality as:	sessment			No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	SNRIs versus placebo (low back pain +/- sciatica)	Control	Relative (95% CI)	Absolute		
SF-36 (Du	loxetine 20m	g) - Bodily	y pain (follow-up <	<4 months; rang	e of scores: 0-1	00; Better indicate	ed by higher values)					
1	randomised trials	seriousª		no serious indirectness	no serious imprecision	none	54	108	-	MD 0.15 higher (0.5 lower to 0.8 higher)	MODERATE	CRITICAL
SF-36 (Du	loxetine 20m	g) - Genei	ral health (follow-	up <4 months; ra	ange of scores:	0-100; Better indi	cated by higher values)					
1	randomised trials	serious ^a		no serious indirectness	no serious imprecision	none	54	108	-	MD 0.04 higher (0.94 lower to 1.02 higher)	MODERATE	CRITICAL
SF-36 (Du	loxetine 20m	g) - Menta	l health (follow-u	p <4 months; rar	nge of scores: 0	-100; Better indica	ated by higher values)					
1	randomised trials	seriousª		no serious indirectness	no serious imprecision	none	54	108	-	MD 0.17 lower (1.35 lower to 1.01 higher)		CRITICAL

SF-36 (Duloxetine 20m	ıg) - Phys	ical functioning (follow-up <4 mo	nths; range of s	cores: 0-100; Bette	er indicated by higher va	lues)				
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	54	108	-	MD 0.43 lower (1.68 MC lower to 0.82 higher)	ODERATE	CRITICA
SF-36 (Duloxetine 20m	ıg) - Role-	emotional (follow	/-up <4 months;	range of scores	: 0-100; Better indi	cated by higher values)			I		
1	randomised trials	seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	54	108	-	MD 0.02 higher (0.27 lower to 0.31 higher)	ODERATE	CRITICA
SF-36 (Duloxetine 20m	ıg) - Role	physical (follow-	up <4 months; ra	ange of scores:	0-100; Better indic	ated by higher values)					
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	54	108	-	MD 0.01 higher (0.5 MC lower to 0.52 higher)	ODERATE	CRITICA
SF-36 (Duloxetine 20m	ıg) - Socia	al functioning (fol	low-up <4 days;	range of scores	s: 0-100; Better ind	icated by higher values)			,	'	
1	randomised trials	seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	54	108	-	MD 0.25 higher (0.26 lower to 0.76 higher)	ODERATE	CRITICA
SF-36 (Duloxetine 20m	ng) - Vitali	ty (follow-up <4 r	nonths; range of	scores: 0-100;	Better indicated by	higher values)			<u> </u>		
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	54	108	-	MD 0.22 lower (1.42 MC lower to 0.98 higher)	ODERATE	CRITICA
			T									

⁽a) Downgraded by one increment if the majority of the evidence was at high risk of bias

Table 269: SNRIs versus placebo (low back pain with/without sciatica)

		·	Quality as	sessment		·	No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	SNRIs versus placebo (low back pain +/- sciatica)	Control	Relative (95% CI)	Absolute	•	·

OF 00 (D		\ D				0.400. D.44	. 4					
SF-36 (DI	uloxetine 120	mg) - Boo	ally pain (follow-u	p <4 montns; ra	inge of scores:	0-100; Better indica	ated by higher values)					
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	101	108	-	MD 0.75 higher (0.21 to 1.29 higher)	LOW	CRITICAL
SF-36 (Du	uloxetine 120	mg) - Ger	neral health (follo	w-up <4 months	; range of score	es: 0-100; Better in	dicated by higher values	s)			1	
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	101	108	-	MD 0.15 higher (0.67 lower to 0.97 higher)	MODERATE	CRITICAL
SF-36 (Du	uloxetine 120	mg) - Mei	ntal health (follow	-up <4 months;	range of scores	s: 0-100; Better ind	icated by higher values)					
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	101	108	-	MD 0.08 higher (0.9 lower to 1.06 higher)		CRITICAL
SF-36 (Du	uloxetine 120	mg) - Phy	sical functioning	(follow-up <4 m	nonths; range o	f scores: 0-100; Be	tter indicated by higher	values)				
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	102	108	-	MD 0.32 higher (0.72 lower to 1.36 higher)	MODERATE	CRITICAL
SF-36 (Du	uloxetine 120	mg) - Rol	e-emotional (folio	ow-up <4 months	s; range of scor	res: 0-100; Better in	dicated by higher value	s)				
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	101	108	-	MD 0.06 higher (0.19 lower to 0.31 higher)	MODERATE	CRITICAL
SF-36 (Du	uloxetine 120	mg) - Rol	e physical (follow	/-up <4 months;	range of score	s: 0-100; Better ind	icated by higher values)				!	
1	randomised trials	seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	101	108	-	MD 0.05 higher (0.37 lower to 0.47 higher)	MODERATE	CRITICAL
SF-36 (Du	uloxetine 120	mg) - Soc	cial functioning (f	ollow-up <4 mor	nths; range of s	cores: 0-100; Bette	r indicated by higher va	lues)			·	
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	101	108	-	MD 0.12 lower (0.55 lower to 0.31 higher)		CRITICAL

SF-36 ([Ouloxetine 120	mg) - Vita	lity (follow-up <4	months; range o	f scores: 0-100;	Better indicated b	oy higher values)			
1	randomised trials	serious ^a			no serious imprecision	none	101	108	MD 0.47 lower (1.47 lower to 0.53 higher)	CRITICAL

⁽a) Downgraded by one increment if the majority of the evidence was at high risk of bias

J.12.2 Anti-epileptics versus placebo

Table 270: Gabapentinoids versus placebo (low back pain with sciatica population)

			Quality ass	essment			No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Gabapentinoids versus placebo (low back pain with sciatica)	Control	Relative (95% CI)	Absolute		
Back pai	n at rest (follo	ow-up <4	months; measure	ed with: VAS; ra	inge of score	s: 0-10; Better in	dicated by lower values)	ļ			ļ	
1	randomised trials		no serious inconsistency	no serious indirectness	serious ^b	none	31	34	-	MD 0.21 lower (1.22 lower to 0.8 higher)	LOW	CRITICAL
Back pai	n on moveme	nt (follow	-up <4 months; r	neasured with:	VAS; range o	of scores: 0-10; B	etter indicated by lower value	s)			-	
1	randomised trials		no serious inconsistency	no serious indirectness	serious ^b	none	31	34		MD 0.33 lower (1.15 lower to 0.49 higher)		CRITICAL
Adverse	events											
1	randomised trials		no serious inconsistency	no serious indirectness	serious ^b	none	19/31 (61.3%)	13/34 (38.2%)	RR 1.60 (0.96 to 2.67)	229 more per 1000 (from 15 fewer to 639 more)	LOW	IMPORTANT

⁽a) Downgraded by one increment if the majority of the evidence was at high risk of bias

⁽b) Downgraded by one increment if the confidence interval crossed one MID

⁽b) Downgraded by one increment if the confidence interval crossed one MID

Table 271: Other anticonvulsants versus placebo (Low back pain with/without sciatica)

										Quality	Importance
Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Other anticonvulsants versus placebo (low back pain +/- sciatica)	Control	Relative (95% CI)	Absolute	Quality	Importance
follow-up <	4 months	; measured with	: ODI; range of	scores: 0-100	Better indicated	by lower values)					
andomised ials			no serious indirectness	no serious imprecision	none	48	48	-	MD 4.9 lower (7 to 2.8 lower)	MODERATE	CRITICAL
ity (follow-u	up <4 mo	nths; measured	with: McGill pa	in questionnai	re; range of score	es: 0-78; Better indicated by	lower va	alues)			
andomised ials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	48	48	-	MD 11.4 lower (12.16 to 10.64 lower)	MODERATE	CRITICAL
ysical funct	tion (follo	ow-up <4 months	; range of scor	es: 0-100; Bett	er indicated by h	igher values)					
andomised ials			no serious indirectness	no serious imprecision	none	48	48	-	MD 8 higher (5.07 to 10.93 higher)	MODERATE	CRITICAL
le-physical	(follow-u	ıp <4 months; ra	nge of scores:	0-100; Better in	ndicated by highe	er values)					
andomised ials	serious ^a		no serious indirectness	no serious imprecision	none	48	48	-	MD 7.5 higher (4.42 to 10.58 higher)	MODERATE	CRITICAL
dily pain (fo	ollow-up	<4 months; rang	e of scores: 0-	100; Better ind	icated by higher	l values)					
andomised ials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	48	48	-	MD 2.1 higher (0.49 lower to 4.69 higher)	LOW	CRITICAL
it i	ndomised als ty (follow-tondomised als sical functondomised als e-physical als domised als	ndomised serious ^a ty (follow-up <4 mo ndomised serious ^a sical function (follo ndomised serious ^a se-physical (follow-up ndomised serious ^a	ndomised serious no serious inconsistency ty (follow-up <4 months; measured not make als not serious inconsistency sical function (follow-up <4 months inconsistency e-physical (follow-up <4 months; randomised serious inconsistency e-physical (follow-up <4 months; randomised serious inconsistency lily pain (follow-up <4 months; rangondomised serious no serious inconsistency	ndomised als serious no serious inconsistency indirectness als inconsistency indirectness indirectness als no serious inconsistency inconsistency inconsistency inconsistency indirectness inconsistency inconsistency indirectness inconsistency inconsistency inconsistency indirectness inconsistency	ndomised serious no serious inconsistency indirectness imprecision ty (follow-up <4 months; measured with: McGill pain questionnai andomised serious no serious inconsistency indirectness imprecision sical function (follow-up <4 months; range of scores: 0-100; Better indirectness inconsistency indirectness imprecision als no serious no serious indirectness imprecision als no serious indirectness imp	Indomised serious inconsistency indirectness imprecision inconsistency indirectness imprecision inconsistency indirectness imprecision inconsistency indirectness imprecision inconsistency inconsiste	inconsistency indirectness imprecision ty (follow-up <4 months; measured with: McGill pain questionnaire; range of scores: 0-78; Better indicated by moderate als inconsistency indirectness imprecision inconsistency indirectness imprecision sical function (follow-up <4 months; range of scores: 0-100; Better indicated by higher values) andomised serious inconsistency indirectness imprecision inconsistency indirectness imprecision andomised (follow-up <4 months; range of scores: 0-100; Better indicated by higher values) andomised serious inconsistency inconsistency indirectness imprecision inconsistency indirectness imprecision andomised serious inconsistency indirectness imprecision inconsistency indirectness inconsistency inconsistency indirectness imprecision inconsistency indirectness inconsistency inconsist	Indomised serious no serious inconsistency indirectness imprecision none inconsistency inconsistency indirectness imprecision none inconsistency inconsistency indirectness imprecision none inconsistency indirectness imprecision inconsistency indirectness imprecision none inconsistency indirectness imprecision none inconsistency indirectness imprecision none inconsistency indirectness imprecision inconsistency indirectness imprecision none inconsistency indirectness imprecision none inconsistency indirectness imprecision inconsistency indirectness imprecision none inconsistency indirectness imprecision incons	Indomised serious inconsistency indirectness inconsistency indirectness indirectnes indirectness indirectness indirectness indirectness indirectness	Indomised serious inconsistency indirectness imprecision indirectness	Indomised serious inconsistency indirectness imprecision inconsistency inconsistency indirectness imprecision inconsistency indirectness imprecision inconsistency inconsistency indirectness imprecision inconsistency indirectness indirectne

1	randomised trials	seriousª	no serious inconsistency	no serious indirectness	serious ^b	none	48	48	-	MD 3.5 higher (0.88 to 6.12 higher)	LOW	
SF-36 - \	/itality (follow	v-up <4 m	nonths; range of	scores: 0-100;	Better indicate	d by higher value	es)					
1	randomised trials	seriousª	no serious inconsistency	no serious indirectness	serious ^b	none	48	48	-	MD 6.2 higher (2.88 to 9.52 higher)	LOW	
SF-36 - S	Social functio	ning (fol	low-up <4 month	ns; range of sco	ores: 0-100; Be	tter indicated by I	nigher values)					
1	randomised trials	seriousª	no serious inconsistency	no serious indirectness	serious ^b	none	48	48	-	MD 3.2 higher (0.66 to 5.74 higher)	LOW	CRITICAL
SF-36 - I	Role-emotion	al (follow	-up <4 months;	range of scores	s: 0-100; Better	indicated by high	her values)					
1	randomised trials	seriousª	no serious inconsistency	no serious indirectness	serious ^b	none	48	48	-	MD 2.6 higher (0.53 to 4.67 higher)	LOW	CRITICAL
SF-36 - I	Mental health	(follow-u	 ip <4 months; ra	inge of scores:	0-100; Better in	l ndicated by highe	er values)					
1	randomised trials	seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	48	48	-	MD 5.4 higher (3.14 to 7.66 higher)	MODERATE	CRITICAL
Adverse	events											
1	randomised trials	seriousª	no serious inconsistency	no serious indirectness	serious ^b	none	18/48 (37.5%)	10/48 (20.8%)	RR 1.80 (0.93 to 3.49)	167 more per 1000 (from 15 fewer to 519	LOW	IMPORTANT

⁽a) Downgraded by one increment if the majority of the evidence was at high risk of bias
(b) Downgraded one increment if the confidence interval crossed one MID

NICE. 2016 Anticonvulsants versus usual care (cohort study)

Table 272: Gabapentinoids versus usual care (low back pain with sciatica)

			Quality ass	sessment			No of patients	5		Effect	Quality	Importanc
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Anticonvulsants versus usual care	Control	Relative (95% CI)	Absolute		
Pain inte	 nsity (follow-up	12 weeks	l s; range of score	s: 0-10; Better ir	l ndicated by low	er values)						
	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	564	119	-	MD 1.4 lower (1.81 to 0.99 lower)	VERY LOW	CRITICAL
HADS- aı	nxiety (follow-u	p 12 week	s; range of score	es: 0-21; Better i	ndicated by lov	ver values)		1				
	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	564	119	-	MD 1.8 lower (2.42 to 1.18 lower)	VERY LOW	CRITICAL
HADS- de	epression (folio	ow-up 12 v	veeks; range of s	cores: 0-21; Bet	ter indicated by	y lower values)						
	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	564	119	-	MD 1.9 lower (2.58 to 1.22 lower)	VERY LOW	CRITICAL
SF-12 ph	 ysical (follow-ບ	ıp 12 week	s; range of score	es: 0-100; Better	r indicated by h	igher values)]				
	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	564	119	-	MD 3.9 higher (2.21 to 5.59 higher)	VERY LOW	CRITICAL
SF-12 me	ental (follow-up	12 weeks	; range of scores	s: 0-100; Better i	ndicated by hig	her values)						
	observational studies	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	564	119	-	MD 5.3 higher (3.71 to 6.89 higher)	⊕000 VERY LOW	CRITICAL

1	observational studies	,		no serious imprecision	none	347/564 (61.5%)		244 more per 1000 (from 111 more to 414 more)	VERY LOW	IMPORTANT
							37%	244 more per 1000 (from 111 more to 414 more)		

⁽a) Downgraded by two increments if the majority of the evidence was at very high risk of bias

J.12.4 Muscle relaxants versus placebo

Table 273: Muscle relaxants versus placebo (low back pain with/without sciatica population)

			Quality as	sessment			No of patients			Effect	0 111	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Muscle relaxants versus placebo (low back pain with sciatica)	Control	Relative (95% CI)	Absolute	Quality	Importance
Pain at n	ight (follow-ι	ıp <4 mor	l nths; measured w	ith: VAS; range	of scores: 0-1	00; Better indicate	ed by lower values)	<u> </u>				
2	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	97	96	-	MD 0.26 lower (0.99 lower to 0.48 higher)	MODERATE	CRITICAL
Pain at re	est (follow-up	<4 mont	hs; measured wi	th: VAS; range	of scores: 0-10	0; Better indicated	d by lower values)					
2	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	97	96	-	MD 0.11 lower (0.9 lower to 0.69 higher)	MODERATE	CRITICAL
Pain wall	king (follow-ເ	ıp <4 mor	nths; measured w	vith: VAS; range	e of scores: 0-1	00; Better indicat	ed by lower values)	1				
2	randomised	serious ^a	no serious	no serious	no serious	none	97	96	-	MD 0.19 higher (0.56 lower to 0.95	MODERATE	CRITICAL

⁽b) Downgraded by one increment if the confidence interval crossed one MID

	trials		inconsistency	indirectness	imprecision					higher)		
Muscle s	pasms (follow	w-up 13 -	18 days; range o	of scores: 1-5; E	Better indicated	l by lower values)						
1		· ,	no serious inconsistency	no serious indirectness	serious ^c	none	16	19		MD 0.10 higher (0.03 to 0.17 higher)	VERY LOW	CRITICAL
Adverse	events											
3	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	114/208 (54.8%)	57/204 (27.9%)	RR 1.97 (1.53 to 2.54)	271 more per 1000 (from 148 more to 430 more)	MODERATE	IMPORTANT

⁽a) Downgraded by one increment if the majority of the evidence was at high risk of bias

J.12.5 Muscle relaxants versus usual care

Table 274: Muscle relaxants versus usual care (low back pain without sciatica)

			Quality as	sessment			No of patien			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Muscle relaxants versus usal care	Control	Relative (95% CI)	Absolute		
Pain - Pai	n - Pain on movement (follow-up <4 months; range of scores: 0-10; Better indicated by lower values)											
		- ,	no serious inconsistency		no serious imprecision	none	94	91	-	MD 2.11 lower (2.72 to 1.5 lower)	LOW	CRITICAL
Pain - Pai	n at rest (folio	ow-up <4 i	months; range of	scores: 0-10; Bet	tter indicated by	lower values)						
1		,	no serious inconsistency	no serious indirectness	serious ^b	none	94	91	-	MD 1.53 lower (2.16 to 0.9 lower)	VERY LOW	CRITICAL

⁽b) Downgraded by two increments if the majority of the evidence was at very high risk of bias

⁽c) Downgraded by one increment if the confidence interval crossed one MID

Pain - Pain at night (follow-up <4 months; range of scores: 0-10; Better indicated by lower values)														
1	randomised	very	no serious	no serious	serious ^b	none	94	91	-	MD 1.36 lower (1.98	VERY	CRITICAL		
	trials	serious ^a	inconsistency	indirectness						to 0.74 lower)	LOW			
Adverse e	Adverse effects (follow-up <4 months)													
1	randomised	very	no serious	no serious	very serious ^c	none	12/101	12/96	OR 0.94 (0.4	7 fewer per 1000	VERY	IMPORTAN'		
	trials	seriousª	inconsistency	indirectness			(11.9%)	(12.5%)	to 2.22)	(from 71 fewer to 116 more)	LOW			
			if the americanity of		1									

⁽a) Downgraded by 2 increments if the majority of the evidence was at very high risk of bias

J.12.6 Opioids versus placebo

Table 275: Opioids versus placebo (low back pain population)

			Quality ass	sessment			No of patients			Effect			
No of studies	Design Inconsistence		Inconsistency	Indirectness	Imprecision Other considerations		Opioid analgesics versus placebo (LBP population)	Control	Relative (95% CI)	Absolute	Quality	Importance	
Quality o	f life (Physica	al compor	nent Score, PCS,0)-100)< 4 month	s (follow-up <4	I months; Better i	ndicated by lower value	es)					
1	randomised trials			no serious indirectness	Serious ^b	none	193	196	-	MD 3.9 higher (1.95 to 5.85 higher)	⊕⊕OO LOW	CRITICAL	
Quality o	uality of life (Mental component Score, MCS,0-100)< 4 months (follow-up <4 months; Better indicated by lower values)												

⁽b) Downgraded by 1 increment if the confidence interval crossed one MID

⁽c) Downgraded by 2 increment if the confidence interval crossed two MIDs

1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	193	196	-	MD 3.22 lower (5.37 to 1.07 lower)	⊕⊕⊕O MODERATE	CRITICAL		
Function	n(RMDQ, 0-24)	<4 month	ns (follow-up <4 n	nonths; Better i	ndicated by lov	ver values)								
7	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	790	720	-	MD 1.32 lower (1.88 to 0.75 lower)	⊕⊕⊕O MODERATE	CRITICAL		
Pain inte	ensity (<4 mor	nths) (VAS	6 0-10) (follow-up	<4 months; Be	tter indicated b	y lower values)								
12	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	1848	1420	-	MD 0.59 lower (0.61 to 0.56 lower)	⊕⊕⊕O MODERATE	CRITICAL		
Respond	esponder ≥30%in pain intensity on NRS scale (follow-up <4 months)													
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	95/193 (49.2%)	65/196 (33.2%)	RR 1.48 (1.16 to 1.9)	159 more per 1000 (from 53 more to 298 more)	⊕⊕OO LOW	IMPORTANT		
Respond	der ≥50%in pa	in intensi	ty on NRS scale (follow-up <4 m	onths)									
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	74/193 (38.3%)	48/196 (24.5%)	RR 1.57 (1.16 to 2.12)	140 more per 1000 (from 39 more to 274 more)	⊕⊕OO LOW	IMPORTANT		
Adverse	events													
7	randomised trials	Serious ^a	very serious ^c	no serious indirectness	no serious imprecision	none	356/1004 (35.5%)	121/800 (15.1%)		210 more per 1000 (from 70 more to	⊕000 VERY LOW	IMPORTANT		

									3.92)	442 more)			
Quality o	of life (Individ	ual domai	in scores, SF36,	0-100) < 4 montl	hs - Physical fu	nctioning (Better	indicated by lower valu	ies)					
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	150	146	-	MD 0.7 lower (6.92 lower to 5.52 higher)	⊕000 VERY LOW	CRITICAL	
Quality o	of life (Individ	ual domai	in scores, SF36,	0-100) < 4 month	hs - Role - phys	ical (Better indica	ated by lower values)						
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	149	146	-	MD 10.1 higher (0.6 to 19.6 higher)	⊕⊕OO LOW	CRITICAL	
Quality o	of life (Individ	ual domai	in scores, SF36,	0-100) < 4 monti	ns - Bodily pain	(Better indicated	by lower values)						
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	151	146	-	MD 4.4 higher (0.49 lower to 9.29 higher)	⊕⊕OO LOW	CRITICAL	
Quality o	of life (Individ	ual domai	in scores, SF36,	0-100) < 4 month	ns - Vitality (Be	tter indicated by I	ower values)						
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	151	145	-	MD 0.3 higher (4.65 lower to 5.25 higher)	⊕OOO VERY LOW	CRITICAL	
Quality o	Quality of life (Individual domain scores, SF36, 0-100) < 4 months - Social functioning (Better indicated by lower values)												
1	randomised trials	Very serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	151	146	-	MD 2 higher (4.13 lower to 8.13 higher)	⊕000 VERY LOW	CRITICAL	

Quality o	f life (Individ	ual domai	n scores, SF36, 0	9-100) < 4 month	ns - Role - emo	tional (Better indi	cated by higher values)					
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	151	146	-	MD 13.1 higher (3.89 to 22.31 higher)	⊕⊕OO LOW	CRITICAL
Quality o	f life (Individ	ual domai	n scores, SF36, 0)-100) < 4 month	ns - Mental hea	Ith (Better indicat	ed by lower values)					
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	151	145	-	mean 0 higher (0.74 lower to 7.34 higher)	⊕⊕OO LOW	CRITICAL
Quality o	f life (Individ	ual domai	n scores, SF36, 0)-100) < 4 month	ns - General he	alth (Better indica	ited by lower values)					
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	very serious ^b	none	146	144	-	MD 0.4 lower (5.28 lower to 4.48 higher)	⊕000 VERY LOW	CRITICAL

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias (b) Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

Table 276: Opioids versus placebo (low back pain with sciatic population)

	I		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,								
			Quality asse	essment			No	of patients	Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other Opiod Placeho (LBP with Pelative					
Adverse e	events										

⁽c) Downgraded by two increments due to unexplained herterogeneity ($l^2=87\%$)

trials serious inconsistency indirectness (53%) (52.5%) (0.65 to 1.59) (from 107 fewer to 1.59) VERY LOW (a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias.
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Paracetamol versus placebo J.12.7

Table 277: Paracetamol versus placebo (low back pain with/without sciatica)

			Quality as	sessment			No of patients Effect Paracetamol versus				Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Paracetamol versus placebo (low back pain +/- sciatica)	Control	Relative (95% CI)	Absolute		
Pain inte	nsity (follow-	up <4 moi	nths; measured v	with: VAS; range	e of scores: 0-1	0; Better indicated	d by lower values)					
1		very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	506	505	-	MD 0.1 lower (0.38 lower to 0.18 higher)	LOW	CRITICAL
Function	(follow-up <4	1 months;	measured with:	RMDQ; range of	f scores: 0-24;	l Better indicated b	y lower values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	504	503	-	MD 0 higher (0.57 lower to 0.57 higher)	LOW	CRITICAL
SF-12 Ph	ysical score ((follow-up	<4 months; rang	ge of scores: 0-1	 100; Better indi	cated by higher va	alues)					
1		- ,	no serious inconsistency	no serious indirectness	no serious imprecision	none	252	243	-	MD 0.2 higher (1.33 lower to 1.73 higher)	LOW	CRITICAL
SF-12 Me	ental score (fo	ollow-up <	4 months; range	of scores: 0-10	 0; Better indica	ted by higher valu	ues)					

⁽a) Downgraded by 1 increment if the majority of the evidence was at right risk of blas, and downgraded by 2 increments if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

1	randomised trials	,	no serious inconsistency		no serious imprecision	none	252	243	-	MD 0.9 higher (0.05 lower to 1.85 higher)	LOW	CRITICAL			
Adverse	Adverse events														
1	randomised trials	,	no serious inconsistency	no serious indirectness	serious ^b	none	99/534 (18.5%)	98/531 (18.5%)	RR 1.00 (0.78 to 1.29)	0 fewer per 1000 (from 41 fewer to 54 more)	VERY LOW	IMPORTANT			

⁽a) Downgraded by two increments if the majority of the evidence was at very high risk of bias

J.12.8 NSAIDs versus placebo

Table 278: NSAIDs versus placebo (low back pain with/without sciatica)

			Quality as	sessment			No of patients	3		Effect	Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	NSAID versus placebo (low back pain +/- sciatica)	Control	Relative (95% CI)	Absolute	quanty	mportunos	
Pain inte	nsity <4 mont	hs NSAID	20 mg with/with	out sciatica (fol	low-up 14 days	; range of scores:	0-10; Better indicate	d by low	er values)				
1	randomised trials			no serious indirectness	serious ^b	none	33	35	-	MD 0.23 lower (0.76 lower to 0.3 higher)		CRITICAL	
Pain 0-10 lower val	•	ence) < 4	months low back	c pain without w	ith/without scia	itica (NSAID 60mg) (follow-up 12 weeks	; measu	red with: VA	AS; range of scores:	0-10; Better	indicated by	
2	randomised trials				no serious imprecision	none	210	217	-	MD 1.13 lower (1.57 to 0.7 lower)	MODERATE	CRITICAL	
	Pain 0-10 (mean difference) < 4 months low back pain without with/without sciatica (NSAID 90mg) (follow-up 12 weeks; measured with: VAS; range of scores: 0-10; Better indicated by lower values)												
2	randomised trials			no serious indirectness	serious ^b	none	210	212	-	MD 1.02 lower (1.45 to 0.59 lower)	LOW	CRITICAL	

⁽b) Downgraded by one increment if the confidence interval crossed one MID

	n (mean differ r values)	ence) < 4	months low bac	k pain without v	vith/without sci	atica (NSAID 60mg	g) (follow-up 12 week	ks; measur	ed with: RI	MDQ; range of score	s: 0-24; Bette	er indicated
	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	210	217	-	MD 2.64 lower (3.61 to 1.67 lower)	LOW	CRITICA
	n (mean differ r values)	ence) < 4	months low bac	k pain without w	vith/without sci	atica (NSAID 90mg	g) (follow-up 12 weel	ks; measui	ed with: RI	MDQ; range of score	s: 0-24; Bette	er indicate
	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	210	212	-	MD 2.23 lower (3.19 to 1.26 lower)	LOW	CRITICA
	(mean differer ter indicated l			pain without wi	th/without sciat	ica (NSAID 60mg)	(follow-up 12 weeks	; measure	d with: SF-	12 Physical compone	ent; range of	scores: 0
	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	210	217	-	MD 2.31 higher (0.61 to 4.02 higher)	MODERATE	CRITICA
	(mean differer ter indicated l			pain without wi	th/without sciat	ica (NSAID 90mg)	(follow-up 12 weeks	; measure	d with: SF1	2 - Physical compon	ent; range o	f scores:
	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	210	212	-	MD 2.80 higher (1.1 to 4.49 higher)	MODERATE	CRITICA
	(mean differer ter indicated l			pain without wi	th/without sciat	ica (NSAID 60mg)	(follow-up 12 weeks	; measure	d with: SF-	12 Mental componen	t; range of s	cores: 0-
	randomised trials	seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	210	217	-	MD 0.49 higher (1.06 lower to 2.05 higher)	MODERATE	CRITICA
	(mean differer ter indicated l			pain without wi	th/without sciat	ica (NSAID 90mg)	(follow-up 12 weeks	; measure	d with: SF1	2 - Mental componer	nt; range of s	scores: 0-
	randomised trials	seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	210	212	-	MD 0.07 lower (1.62 lower to 1.47 higher)	MODERATE	CRITICA
dverse	events (follow	w-up 1-12	weeks)									
	randomised trials	seriousª	no serious inconsistency	no serious indirectness	serious ^b	none	289/834 (34.7%)	147/510 (28.8%)	RR 1.11 (0.95 to 1.29)	32 more per 1000 (from 14 fewer to 84 more)	LOW	IMPORTA

⁽a) Downgraded by one increment if the majority of the evidence was at high risk of bias

(b) Downgraded by one increment if the confidence interval crossed one MID

Table 279: NSAIDS versus placebo (low back pain only)

			Quality asse	essment			No of patients	5		Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	NSAID versus placebo low back pain only	Control	Relative (95% CI)	Absolute	Quality	Importance
Pain inten	nsity (VAS 0-1	0 change	score) low back p	ain only- lbuprof	en (follow-up	o 7 days; range of	scores: 0-10; Better i	ndicated	by lower val	ues)		
	randomised trials	seriousª		no serious indirectness	serious ^b	none	103	92	-	MD 1.13 lower (1.85 to 0.41 lower)	LOW	CRITICAL
Pain inter	nsity (VAS 0-1	0 change	score) low back p	ain only- Diclofe	nac-K (follow	/-up 7 days; range	of scores: 0-10; Bett	er indica	ted by lower	values)		
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	107	92	-	MD 1.09 lower (1.83 to 0.35 lower)	LOW	CRITICAL
Adverse e	Adverse events (follow-up <4 months)											
4	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	173/624 (27.7%)	96/401 (23.9%)	RR 1.07 (0.87 to 1.31)	17 more per 1000 (from 31 fewer to 74 more)	LOW	IMPORTANT

⁽a) Downgraded by one increment if the majority of the evidence was at high risk of bias

J.12.9 Antibiotics versus placebo

Table 280: Antibiotics versus placebo (low back pain with/without sciatica)

			Quality ass	sessment			No of patie	nts		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Antibiotics versus placebo	Control	Relative (95% CI)	Absolute	•	·

⁽b) Downgraded by one increment if the confidence interval crossed one MID

	. (0.40) .4			5								
Back pai	in (0-10) - <4 m	nonths (fo	llow-up <4 month	ns; Better indica	ted by lower va	lues)					T	
l	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	76	67	-	MD 1.3 lower (3.46 lower to 0.86 higher)	LOW	CRITICAL
Back pai	in (0-10) - 4-12	months (follow-up 4-12 me	onths; Better inc	dicated by lower	r values)						
	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	77	67	-	MD 2.6 lower (5.08 to 0.12 lower)	LOW	CRITICAL
Disabilit	y (RMDQ) - <4	months (follow-up <4 mon	ths; Better indic	cated by lower v	alues)						
l	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	76	67	-	MD 2.5 lower (7.13 lower to 2.13 higher)	MODERATE	CRITICAL
Disabilit	y (RMDQ) - 4-1	12 months	s (follow-up 4-12 r	months; Better i	ndicated by low	er values)						
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	77	67	-	MD 7 lower (12.56 to 1.44 lower)	MODERATE	CRITICAL
ED-5D -	<4 months (fo	llow-up <	4 months; Better	indicated by low	ver values)							
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	76	67	-	MD 5 higher (15.16 lower to 25.16 higher)	MODERATE	CRITICAL
ED-5D	4-12 months (1	follow-up	4-12 months; Bet	ter indicated by	lower values)		•					
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	77	67	-	MD 15 higher (5.17 lower to 35.17 higher)	MODERATE	CRITICAL
Healthca	re utilisation ((dr consu	Itation for back pa	ain) (follow-up <	4 months)							
I	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	18/77 (23.4%)	28/67 (41.8%)	RR 0.56 (0.34 to 0.92)	184 fewer per 1000 (from 33 fewer to 276 fewer)	LOW	IMPORTAN
Adverse	events (GI co	mplaints)	(follow-up <4 mo	nths)				· '				
1	randomised trials	Serious	no serious inconsistency	no serious indirectness	no serious imprecision	none	59/90 (65.6%)	17/72 (23.6%)	RR 2.78 (1.79 to 4.32)	420 more per 1000 (from 187 more to 784 more)	MODERATE	IMPORTAN

⁽a) Downgraded by1 increment if the majority of the evidence was at high risk of bias

⁽b) Downgraded by 1 increment if the confidence interval crossed one MID

Head to head comparisons Table 281: Anti-epileptics vers

Table 281: Anti-epileptics versus antidepressants (TCAs) low back pain with/without sciatica

			Quality asse	essment			No of patients			Effect	Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Antiepileptic versus antidepressant (TCA)	Control	Relative (95% CI)	Absolute			
Adverse (Adverse events (follow-up 6 weeks)												
1	randomised trials			no serious indirectness	Serious ^b	none	29/97 (29.9%)	17.5%	RR 1.71 (1.02 to 2.87)	124 more per 1000 (from 3 more to 327 more)	LOW	IMPORTANT	

⁽a) Downgraded by1 increment if the majority of the evidence was at high risk of bias

Table 282: Antidepressants versus paracetamol – low back pain with/without sciatica

			Quality asses	sment			No of patients	S		Effect	Quality	lance out on o
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Amitriptyline versus paracetamol	Control	Relative (95% CI)	Absolute	Quality	Importance
Pain (VAS	0-15) (follow	-up 5 weeks;	measured with: V	AS ; range of sc	ores: 0-15; B	Setter indicated by	lower values)		_			
1	randomised no serious no serious inconsistency lindirectness					none	20	19	-	MD 1.83 lower (3.66 lower to 0 higher)	MODERATE	CRITICAL
Psycholog	gical distress	(follow-up 5	weeks; measured	with: Beck depr	ession inven	tory; range of sco	ores: 0-63; Better indic	cated by	lower va	alues)	•	
1				no serious indirectness	Serious ^a	none	20	19	-	MD 2.17 lower (7.35 lower to 3.01 higher)		CRITICAL
Psycholog	gical distress	(follow-up 5	weeks; measured	with: STAI-state	; range of so	cores: 20-80; Bette	r indicated by lower v	/alues)				

⁽b) Downgraded by 1 increment if the confidence interval crossed one MID

Table 283: Opioid plus paracetamol versus opioid – low back pain with/without sciatica

			Quality asse	ssment			No of patients	S		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision		Opioid + non-opioid analgesic versus opioid	Control	Relative (95% CI)	Absolute	Quality	importance
Adverse	events (follov	v-up 10 days	s)									
1		no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^a	none	30/59 (50.8%)	38.4%	RR 0.69 (0.52 to 0.93)	119 fewer per 1000 (from 27 fewer to 184 fewer)		IMPORTANT

⁽a) Downgraded by 1 increment if the confidence interval crossed one MID

Table 284: Opioid plus paracetamol versus NSAIDs—low back pain with/without sciatica

			Quality ass	essment			No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Opioids + non-opioid analgesics versus NSAIDs	Control	Relative (95% CI)	Absolute	Quality	importance
Pain inter	nsity (VAS) (f	ollow-up 1 v	veeks; range of s	cores: 0-10; Bet	ter indicated by	y lower values)						
1		no serious risk of bias			no serious imprecision	none	58	55	-	MD 0.05 higher (0.81 lower to 0.91 higher)		CRITICAL

Adverse	events (follow	v-up 1 week	s)							
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	38/59 (64.4%)	21/62 (33.9%)	305 more per 1000 (from 95 more to 620 more)	IMPORTANT

J.12.11 Combined pharmacological treatments versus placebo

Table 285: Opioid and paracetamol versus placebo- low back pain only

			Quality as:	sessment			No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Combination (opioid and non-opioid analgesics) <4 months, low back pain only	Control	Relative (95% CI)	Absolute	quanty	portaneo
Time to d	onset: percep	otible pair	relief (follow-up	3 days)								
	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	108/141 (76.6%)	95/136 (69.9%)	HR 1.22 (0.92 to 1.62)	70 more per 1000 (from 30 fewer to 158 more)	LOW	CRITICAL
								0%		-		
Time to d	nset: meani	ngful pair	relief (follow-up	3 days)	T							
1 -	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious	none	61/141 (43.3%)	45/136 (33.1%)	HR 1.57 (1.05 to 2.35)	137 more per 1000 (from 13 more to 280 more)	LOW	CRITICAL
								0%		-		
Time to r	emedication	(follow-u	p 3 days)									
	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	very serious ^c	none	18/144 (12.5%)	17/136 (12.5%)	HR 0.93 (0.47 to 1.84)	8 fewer per 1000 (from 64 fewer to 93 more)	VERY LOW	CRITICAL
								0%		-		
Adverse	events (follo	w-up 2.5	days)									

	1		ı	1							1	
2	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	106/308 (34.4%)	30/305 (9.8%)	RR 3.48 (2.06 to 5.44)	244 more per 1000 (from 104 more to 437 more)	MODERATE	CRITICAL
SF McGi	II Pain questi	ionnaire (follow-up 91 day	s; range of sco	res: 0-78; Bett	er indicated by lo	wer values)				,	
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	164	161	-	MD 2.2 lower (4.64 lower to 0.24 higher)	MODERATE	CRITICAL
Pain VAS	S (0-10) (follo	w-up 91 c	lays; range of so	cores: 0-10; Bet	ter indicated b	y lower values)						
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	167	169	-	MD 1.55 lower (2.47 lower to 0.63 higher)	LOW	CRITICAL
SF-36 bo	odily pain (fol	low-up 91	I days; range of	scores: 0-100;	Better indicate	d by higher value	s)					
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	164	163	-	MD 6.4 higher (2.09 to 10.71 higher)	LOW	CRITICAL
SF-36 ge	eneral health	(follow-u	o 91 days; range	of scores: 0-10	00; Better indic	ated by higher va	ilues)					
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	164	163	-	MD 3.5 higher (0.94 lower to 7.94 higher)	MODERATE	CRITICAL
SF-36 me	ental health (follow-up	91 days; range	of scores: 0-10	0; Better indica	ated by higher val	ues)					
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	164	163	-	MD 2.6 higher (1.8 lower to 7 higher)	MODERATE	CRITICAL
SF-36 ph	nysical functi	oning (fol	llow-up 91 days;	range of score	s: 0-100; Bette	r indicated by hig	her values)					
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	164	163	-	MD 3.8 higher (1.83 lower to 9.43 higher)	MODERATE	CRITICAL
SF-36 re	ported health	transitio	n (follow-up 91 o	days; range of s	scores: 0-100;	Better indicated b	y higher values)					
1	randomised	Serious ^a	no serious	no serious	no serious	none	164	163	-	MD 2.2 lower	MODERATE	CRITICAL

	trials		inconsistency	indirectness	imprecision					(7.42 lower to 3.02 higher)		
SF-36 ro	le-emotional	(follow-u	p 91 days; range	of scores: 0-10	00; Better indic	cated by higher va	alues)					
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	164	163	-	MD 1.3 higher (8.02 lower to 10.62 higher)	MODERATE	CRITICAL
SF-36 ro	le-physical (f	ollow-up	91 days; range o	of scores: 0-100); Better indica	ted by higher val	ues)					
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	164	163	-	MD 3.8 higher (4.03 lower to 11.63 higher)	MODERATE	CRITICAL
3F-36 sc	ocial function	ing (follo	w-up 91 days; ra	nge of scores:	0-100; Better i	ndicated by highe	er values)					
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	164	163	-	MD 0.7 lower (6.2 lower to 4.8 higher)	MODERATE	CRITICAI
SF36 he	alth survey -	SF-36 vita	ality (follow-up 9	1 days; range o	of scores: 0-10	0; Better indicated	d by higher values)	•				
I	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	164	163	-	MD 1.3 higher (3.16 lower to 5.76 higher)	MODERATE	CRITICAL
Function	n (RMDQ 0-24) (follow-	up 91 days; rang	je of scores: 0-2	24; Better indic	cated by lower val	lues)					
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	164	163	-	MD 0.9 lower (2.16 lower to 0.36 higher)	MODERATE	CRITICAL

⁽a) Downgraded by one increment if the majority of the evidence was at high risk of bias(b) Downgraded by one increment if the confidence interval crossed one MID

Table 286: Opioid and paracetamol versus placebo- low back pain only

			Quality ass	essment			No of patients			Effect	Quality	Importance
No of	Design	Risk of	Inconsistency	Indirectness	Imprecision	Other	Combination (opioid and	Control	Relative	Absolute		

⁽c) Downgraded by two increments if the confidence interval crossed both MIDs

studies		bias				considerations	non-opioid analgesics) <4 months, low back pain with/without sciatica		(95% CI)			
Adverse	events (follo	w-up <4 r	months)					_				
2		no serious risk of bias	no serious inconsistency	no serious indirectness	no serious imprecision	none	116/150 (77.3%)	71/145 (49%)	RR 1.57 (1.31 to 1.89)	279 more per 1000 (from 152 more to 436 more)	HIGH	IMPORTANT
								39.1%		223 more per 1000 (from 121 more to 348 more)		
Respond	ler criteria pa	ain reduct	ion >30% (follow	-up 2 weeks)								
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	serious ¹	none	49/85 (57.6%)	37/90 (41.1%)	RR 1.4 (1.03 to 1.91)	164 more per 1000 (from 12 more to 374 more)	MODERATE	IMPORTANT
								41.1%		164 more per 1000 (from 12 more to 374 more)		
Function	(Korean OD	l 0-100) (f	ollow-up 2 weeks	s; range of sco	res: 0-100; Be	tter indicated by	ower values)					
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	serious ¹	none	83	87	-	MD 4.04 higher (0.16 to 7.91 higher)	MODERATE	CRITICAL
Korean S	Short Form-3	6 Bodily p	pain (follow-up 2	weeks; range of	of scores: 0-10	00; Better indicate	ed by higher values)					
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	no serious imprecision	none	83	87	-	MD 1.6 higher (3.54 lower to 6.74 higher)	HIGH	CRITICAL
Korean S	Short Form-3	6 General	health (follow-u	p 2 weeks; ran	ge of scores: (0-100; Better indi	cated by higher values)	,				
1	randomised trials	no serious risk of	no serious inconsistency	no serious indirectness	Serious ^a	none	83	87	-	MD 4.59 higher (0.52 to 8.66 higher)	MODERATE	CRITICAL

		bias										
Corean	Short Form-3	6 health s	survey (change s	cores) - Mental	l health (follow	/-up 2 weeks: rand	ge of scores: 0-100; Better inc	licated b	ov higher va	alues)		
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	83	87	-	MD 2.09 higher (5.1 lower to 9.28 higher)	HIGH	CRITICAL
Korean	Short Form-3	6 Physica	al functioning (fo	llow-up 2 week	s; range of sc	ores: 0-100; Bette	r indicated by higher values)					
I	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	no serious imprecision	none	83	87	-	MD 3.15 higher (2.03 lower to 8.33 higher)	HIGH	CRITICAI
Korean	Short Form-3	6 Reporte	ed health transiti	on (follow-up 2	weeks; range	of scores: 0-100;	Better indicated by higher va	lues)				
I	trials	no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^a	none	83	87	-	MD 11.17 lower (19.63 to 2.71 lower)	MODERATE	CRITICAL
Korean	Short Form-3	6 Role em	notional (follow-u	ıp 2 weeks; rar	nge of scores:	0-100; Better indi	cated by higher values)					
I		no serious risk of bias	no serious inconsistency	no serious indirectness	no serious imprecision	none	83	87	-	MD 0.66 higher (7.94 lower to 9.26 higher)	HIGH	CRITICAL
Korean	Short Form-3	6 Role ph	ysical (follow-up	2 weeks; rang	e of scores: 0	-100; Better indica	ated by higher values)					
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^a	none	83	87	-	MD 7.35 higher (0.35 to 14.35 higher)	MODERATE	CRITICAL
Korean	Short Form-3	6 Social f	unctioning (follo	w-up 2 weeks;	range of score	es: 0-100; Better i	ndicated by higher values)					
1		no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^a	none	83	87	-	MD 5.14 higher (1.88 lower to 12.16 higher)	MODERATE	CRITICAL
Corean	Short Form-3	6 Vitality	follow-up 2 wee	ks; range of so	ores: 0-100; B	etter indicated by	higher values)				'	

1 randomised no no serious no serious serious inconsistency indirectness risk of bias no serious inconsistency indirectness serious serious inconsistency indirectness serious no serious serious no serious serious no serious no serious no serious no serious serious no serious	87	-	MD 5.32 higher (0.63 lower to 11.27 higher)	MODERATE	CRITICAL
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⁽a) Downgraded by one increment if the confidence interval crossed one MID

Table 287: Opioid and paracetamol versus other treatment (anticonvulsants) placebo- low back pain with/without sciatica

	Quality assessment							patients	1	Effect			
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	CONSIDERATIONS	(onioid and non-	anticonvulsant at <4 months, low back pain only	Relative (95% CI)	Absolute	Quality	Importance	
Numer o	Numer of people discontinued due to adverse events												
		no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^a	none	3/30 (10%)	6.7%	RR 1.5 (0.27 to 8.34)	34 more per 1000 (from 49 fewer to 492 more)	MODERATE	IMPORTANT	

⁽a) Downgraded by one increment if the confidence interval crossed one MID

Combinations of interventions – pharmacological adjunct J.12.12

Low back pain without sciatica J.12.12.1

Table 288: Pharmacological (NSAID) plus manual therapy (massage) compared to manual therapy (massage)

Quality asses	ssment				No of pation	ents	Effect					
No of studies	Design	Risk of bias	Inconsistenc y	Indirectne ss	Imprecisio n	Other	Massage + NSAID	Massage	Relativ e (95%	Absolute	Qualit	Importanc
									CI)		У	е
Pain (VAS 0-:	100 converted t	o 0-10) - ≤	4 months (follo	w-up 2 weeks	; measured w	ith: VAS;	range of sc	ores: 0-10; B	etter indic	ated by lower value	es)	
1: majchrzyck i 2014	Randomised trials	Very serious a	No serious inconsistenc	No serious indirectnes	Seriousb	None	26	28	-	MD 1.16 lower (2.31 to 0.01 lower)	VERY LOW	CRITICAL

Quality asse	ssment				No of pation	ents	Effect					
No of studies	Design	Risk of bias	Inconsistenc Y	Indirectne ss	Imprecisio n	Other	Massage + NSAID	Massage	Relativ e (95% CI)	Absolute	Qualit y	Importanc e
Disability (R	oland Morris) -	≤4 months	(follow-up 2 w	eeks; measure	ed with: RMD	Q; range	of scores: 0	-24; Better in	ndicated b	y lower values)		
1: majchrzyck i 2014	Randomised trials	Very serious a	No serious inconsistenc	No serious indirectnes s	Seriousb	None	26	28	-	MD 0.3 lower (2.7 lower to 2.1 higher)	VERY LOW	CRITICAL
Disability (O	swestry Disabili	ity Index) -	≤4 months (fol	low-up 2 weel	ks; measured	with: OD	l; range of s	cores: 0-100	; Better in	dicated by lower va	lues)	
1: majchrzyck i 2014	Randomised trials	Very serious	No serious inconsistenc	No serious indirectnes	Seriousb	None	26	28	-	MD 4.4 lower (11.06 lower to 2.26 higher)	VERY LOW	CRITICAL

Table 289: Pharmacological (NSAID) + exercise (biomech) compared to electroacupuncture

Quality a	ssessment						No of patients		Effect			
No of studies	Design	Risk of bias	Inconsisten cy	Indirectness	Imprecisio n	Other	NSAID + exercise (biomec h)	Electroacupunctu re	Relativ e (95% CI)	Absolute	Quality	Importanc e
Pain (VA	S 0-10) - ≤4 ma	nths (follo	w-up 3 weeks;	range of scores	:: 0-10; Better	indicate	<u> </u>	alues)	,			
1: shanka r 2011	Randomise d trials	Very serious a	No serious inconsistenc y	No serious indirectness	Seriousb	None	30	30	-	MD 0.9 higher (0.04 to 1.76 higher)	VERY LOW	CRITICAL

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

⁽a) Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias (b) Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

⁽b) Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

Table 290: Opioid and paracetamol versus placebo- low back pain with/without sciatica

Qualit	y assessmer	nt		·	·	iiii witii/ witiio	No. of patients		Effect			
No. of studi es	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other considerations	Combination (opioid and paracetamol) ≤4 months, low back pain with/without sciatica	Con trol	Relativ e (95% CI)	Absolute	Quality	Importa nce
Advers	se events (fo	llow-up ≤	4 months)									
2	Randomi sed trials	No seriou s risk of bias	No serious inconsistenc y	No serious indirectne ss	No serious imprecisio n	None	116/150 (77.3%)	71/1 45 (49 %)	RR 1.57 (1.31 to 1.89)	279 more per 1000 (from 152 more to 436 more)	HIGH	IMPORT ANT
								39.1 %		223 more per 1000 (from 121 more to 348 more)		
Respoi	nder criteria	pain redu	iction >30% (fo	llow-up 2 wee	eks)							
1	Randomi sed trials	No seriou s risk of bias	No serious inconsistenc y	No serious indirectne ss	Seriousa	None	49/85 (57.6%)	37/9 0 (41. 1%)	RR 1.4 (1.03 to 1.91)	164 more per 1000 (from 12 more to 374 more)	MODER ATE	IMPORT ANT
								41.1 %		164 more per 1000 (from 12 more to 374 more)		
Function	on (Korean (DDI 0-100) (follow-up 2 w	veeks; range o	f scores: 0-10	00; Better indica	ted by lower values)					
1	Randomi sed	No seriou	No serious inconsistenc	No serious indirectne	Seriousa	None	83	87	-	MD 4.04 higher (0.16	MODER ATE	CRITICAL

Quality	assessmen	t					No. of patients		Effect			
No. of studi es	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	Combination (opioid and paracetamol) ≤4 months, low back pain with/without sciatica	Con trol	Relativ e (95% CI)	Absolute	Quality	Importa nce
	trials	s risk of bias	У	SS						to 7.91 higher)		
Korean	Short Form	-36 Bodil	y pain (follow-u	p 2 weeks; ra	nge of scores	: 0-100; Better i	ndicated by higher values)					
1	Randomi sed trials	No seriou s risk of bias	No serious inconsistenc y	No serious indirectne ss	Very serious	None	83	87	-	MD 1.6 higher (3.54 lower to 6.74 higher)	LOW	CRITICAL
Korean	Short Form	-36 Gene	ral health (follo	w-up 2 weeks	; range of sco	ores: 0-100; Bett	ter indicated by higher valu	ues)		T	T	T
1	Randomi sed trials	No seriou s risk of bias	No serious inconsistenc y	No serious indirectne ss	Seriousa	None	83	87	-	MD 4.59 higher (0.52 to 8.66 higher)	LOW	CRITICAL
Korean	Short Form	-36 Ment	al health (follow	w-up 2 weeks;	range of sco	res: 0-100; Bett	er indicated by higher valu	es)				
1	Randomi sed trials	No seriou s risk of bias	No serious inconsistenc	No serious indirectne ss	Very serious	None	83	87	-	MD 2.09 higher (5.1 lower to 9.28 higher)	LOW	CRITICAL
Korean	Short Form	-36 Physi	cal functioning	(follow-up 2 v	veeks; range	of scores: 0-100	; Better indicated by highe	r values	s)			
1	Randomi sed trials	No seriou s risk of bias	No serious inconsistenc y	No serious indirectne ss	Serious	None	83	87	-	MD 3.15 higher (2.03 lower to 8.33 higher)	MODER ATE	CRITICAL
Korean	Short Form	-36 Repo	rted health tran	nsition (follow	-up 2 weeks;	range of scores	: 0-100; Better indicated b	y higher	values)			
1	Randomi sed trials	No seriou s risk	No serious inconsistenc	No serious indirectne ss	Seriousa	None	83	87	-	MD 11.17 lower (19.63 to 2.71	MODER ATE	CRITICAL

Quality	y assessmen	t					No. of patients		Effect			
No. of studi es	Design	Risk of bias	Inconsisten cy	Indirectne ss	Imprecisi on	Other consideratio ns	Combination (opioid and paracetamol) ≤4 months, low back pain with/without sciatica	Con trol	Relativ e (95% CI)	Absolute	Quality	Importa nce
		of bias								lower)		
Korear	Short Form	-36 Role	emotional (follo	w-up 2 weeks	; range of sc	ores: 0-100; Bet	ter indicated by higher val	ues)				
1	Randomi sed trials	No seriou s risk of bias	No serious inconsistenc y	No serious indirectne ss	Very serious	None	83	87	-	MD 0.66 higher (7.94 lower to 9.26 higher)	LOW	CRITICAL
Korear	Short Form	-36 Role	physical (follow	-up 2 weeks; ı	range of scor	es: 0-100; Bette	r indicated by higher value	s)				
1	Randomi sed trials	No seriou s risk of bias	No serious inconsistenc y	No serious indirectne ss	Seriousa	None	83	87	-	MD 7.35 higher (0.35 to 14.35 higher)	MODER ATE	CRITICAL
Korear	Short Form	-36 Socia	I functioning (fo	ollow-up 2 we	eks; range of	scores: 0-100; B	Better indicated by higher v	/alues)				
1	Randomi sed trials	No seriou s risk of bias	No serious inconsistenc y	No serious indirectne ss	Seriousa	None	83	87	-	MD 5.14 higher (1.88 lower to 12.16 higher)	MODER ATE	CRITICAL
Korean	Short Form	-36 Vitali	ty (follow-up 2	weeks; range	of scores: 0-1	LOO; Better indic	ated by higher values)					
1	Randomi sed trials	No seriou s risk of bias	No serious inconsistenc y	No serious indirectne ss	Seriousa	None	83	87	-	MD 5.32 higher (0.63 lower to 11.27 higher)	MODER ATE	CRITICAL

⁽a) Downgraded by one increment if the confidence interval crossed one MID.

J.13 Combined interventions: multidisciplinary biopsychosocial rehabilitation (MBR) programmes

I.13.1 Population: overall with or without sciatica

Table 291: MBR programme 3 elements: physical + psychological + education vs. Usual care/waiting list control

			Quality as	sessment			No of patien	ts		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	MBR programme 3 elements: physical + psychological + education	care/waiting list	Relative (95% CI)	Absolute	Quality	Importance
Pain seve	erity (intensit	y), VAS 0)-10 (> 4 months)	(follow-up >4 m	nonths – 1 year	; range of scores	: 0-10; Better indicated by lo	ower values)				
1		,	no serious inconsistency		no serious imprecision	none	29	23	1	MD 2.5 lower (3.65 to 1.35 lower)	⊕⊕OO LOW	CRITICAL
Function	, ODI 0-100 (>	> 4 month	ns)(follow-up >4 ı	months – 1 year	r; range of sco	res: 0-100; Better	indicated by lower values)					
1		- ,	no serious inconsistency	no serious indirectness	Serious ^b	none	30	23	-	MD 16.4 higher (7.06 to 25.74 higher)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 292: MBR programme 3 elements: physical + psychological + education vs. Single intervention (aerobic exercise)

			Quality as:	sessment			No of patient	s		Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	MBR programme 3 elements: physical + psychological + education	Single intervention	Relative (95% CI)	Absolute	Quality	Importance

^b Downgraded by 1 increment if the confidence interval crossed one MID

					O : h		40			MD 4 0 I		ODITIOA
l	randomised trials	Serious	no serious inconsistency	no serious indirectness	Serious ^b	none	48	51	-	MD 1.0 lower (4.76 lower to 2.76 higher)	⊕⊕OO LOW	CRITICA
Qualit	y of life, SF-12	physical (0-100 (>4 month	s – 1 year) - Ex	ercise - aerobio	(follow-up >4 mo	onths – 1 year; range of	scores: 0-100; Be	etter indi	cated by lower v	alues)	
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	48	51	-	MD 1 lower (4.81 lower to 2.81 higher)	⊕⊕OO LOW	CRITICA
Qualit	y of life, SF-12	mental 0-	100 (≤4 months)	- Exercise - ae	robic (follow-u	o ≤4 months; ranç	ge of scores: 0-100; Bett	er indicated by lo	wer valu	ies)		
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	48	51	-	MD 1 higher (2.55 lower to 4.55 higher)	⊕⊕OO LOW	CRITICA
Qualit	y of life, SF-12	mental 0-	100 (>4 months	– 1 year) - Exer	cise - aerobic (follow-up >4 mon	iths– 1 year; range of sc	ores: 0-100; Bett	er indica	ted by lower val	ues)	
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	48	51	-	MD 1 higher (1.97 lower to 3.97 higher)	⊕⊕OO LOW	CRITICA
Pain s	everity, NRS 0-	-10 (≤4 mo	onths) - Exercise	- aerobic (folic	ow-up ≤4 month	ns; range of score	es: 0-10; Better indicated	l by lower values)			
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	48	51	-	MD 0 higher (0.87 lower to 0.87 higher)	⊕⊕⊕O MODERATE	CRITICA
1			ontho 1 year) I	Exercise - aero	bic (follow-up >	4 months- 1 year	r; range of scores: 0-10;	Better indicated	by lower	values)		
Pain s	everity, NRS 0-	·10 (>4 mo	Jillis - i year) - i									
Pain s		<u> </u>	no serious inconsistency	no serious indirectness	very serious ^c	none	48	51	-	MD 0 higher (0.72 lower to 0.72 higher)	#000 VERY LOW	CRITICA
1	randomised trials	Serious ^a	no serious inconsistency	indirectness			48 0-24; Better indicated by		-	(0.72 lower to		CRITICA

1	randomised trials				no serious imprecision	none	48	51	-	MD 0.10 lower (1.49 lower to 1.29 higher)	⊕⊕⊕O MODERATE	CRITICAL
Function	on, back perfo	rmance so	cale 0-15 (≤4 mor	nths) - Exercise	- aerobic (follo	ow-up ≤4 months	; range of scores: 0-15; Bet	ter indicated I	y lower	values)		
1	randomised trials		no serious inconsistency		no serious imprecision	none	49	51	-	MD 0 higher (1.1 lower to 1.1 higher)	⊕⊕⊕O MODERATE	CRITICAL

 ^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias
 ^b Downgraded by 1 increment if the confidence interval crossed one MID
 ^c Downgraded by 2 increments if the confidence interval crossed two MIDs

Table 293: MBR programme 3 elements: physical + psychological + education vs. Combined intervention (manual therapy + exercise + postural therapy + self management; manual therapy + exercise + advice)

			ileiit, iliailuai	шениру		100,							
			Quality ass	sessment			No of patien	ts		Effect			
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	MBR programme 3 elements: physical + psychological + education	Combined intervention	Relative (95% CI)	Absolute	Quality	Importance	
Pain severity, NRS 0-10 (≤ 4 months) (range of scores: 0-10; Better indicated by lower values)													
	randomised trials	,			no serious imprecision	none	75	75	-	MD 3.10 lower (3.59 to 2.61 lower)	⊕⊕OO LOW	CRITICAL	
Pain sev	erity, VAS 0-	10 (> 4 mo	onths)- manual +	exercise + adv	rice (follow-up	>4 months - 1 ye	ear; range of scores: 0-10;	Better indicate	d by lowe	er values)			
1	randomised trials	Serious ^b		no serious indirectness	Serious ^c	none	46	55	-	MD 0.40 lower (1.51 lower to 0.71 higher)	⊕⊕OO LOW	CRITICAL	
Pain sev values)	erity, NRS 0-	10 (>4 mo	onths– 1 year) - n	nanual + exerci	se + postural t	herapy + self mai	nagement (follow-up >4 m	onths – 1 year;	range of	scores: 0-10; Be	etter indicate	d by lower	
1	randomised	very	no serious	no serious	no serious	none	75	75	-	MD 1.8 lower	⊕⊕OO	CRITICAL	

	trials	serious ^a	inconsistency	indirectness	imprecision					(2.3 to 1.3 lower)	LOW	
Function	n, ODI 0-100 (≤4 month	s) manual + exe	cise + postura	l therapy + self	management (fo	llow-up >4 months; range	of scores: 0-10	0; Better	indicated by lov	ver values)	
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	75	75	-	MD 9.8 lower (11.45 to 8.15 lower)	⊕⊕OO LOW	CRITICAL
Function lower va		>4 month	ıs – 1 year) manı	ıal + exercise +	postural thera	py + self manage	ement (Copy) (follow-up >4	months – 1 yea	ar; range	of scores: 0-10	0; Better indi	cated by
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	75	75	-	MD 15.8 lower (17.48 to 14.12 lower)	⊕⊕OO LOW	CRITICAL
Function	, RMDQ 0-24	(>4 mon	ths – 1 year) - ma	anual + exercis	e + advice (foll	ow-up >4 months	- 1 year; range of scores:	0-24; Better in	dicated I	by lower values)		
1	randomised trials	Serious ^b	no serious inconsistency	no serious indirectness	Serious ^c	none	46	55	-	MD 2.3 lower (4.51 to 0.09 lower)	⊕⊕OO LOW	CRITICAL
Quality o	of life, EQ-5D	-0.5 to 1.	0 (>4 months – 1	year) (follow-u	p >4 months -	1 year; range of	scores: -0.5-1; Better indic	ated by higher	values)			
1	randomised trials	Serious ^b	no serious inconsistency	no serious indirectness	no serious imprecision	none	46	55	-	MD 0.00 higher (0.11 lower to 0.11 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality o	of life, SF-36	0-100 (≤ 4	months) - Physi	cal functioning	(range of sco	res: 0-100; Better	indicated by higher value	s)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	75	75	-	MD 20.8 higher (17.49 to 24.11 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life, SF-36	0-100 (≤ 4	months) - Emot	ional role (rang	e of scores: 0-	100; Better indica	ated by higher values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	75	75	-	MD 21.8 higher (15.3 to 28.3 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life, SF-36	0-100 (≤ 4	months) - Gene	ral health (rang	e of scores: 0-	100; Better indica	ated by higher values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^c	none	75	75	-	MD 16.7 higher (12.74 to 20.66 higher)	⊕OOO VERY LOW	CRITICAL

Quality	of life, SF-36	0-100 (≤ 4	months) - Ment	al health (range	of scores: 0-1	100; Better indicat	ted by higher values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	75	75	-	MD 23.8 higher (20.34 to 27.26 higher)	⊕⊕OO LOW	CRITICA
Quality	of life, SF-36	0-100 (≤ 4	months) - Phys	ical pain (range	of scores: 0-1	100; Better indicat	ted by higher values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	75	75	-	MD 17.8 higher (13.06 to 22.54 higher)	⊕⊕OO LOW	CRITICA
Quality	of life, SF-36	0-100 (≤ 4	months) - Phys	ical role (range	of scores: 0-1	00; Better indicate	ed by higher values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	75	75	-	MD 22.5 higher (16.9 to 28.1 higher)	⊕⊕OO LOW	CRITICA
Quality	of life, SF-36	0-100 (≤ 4	months) - Soci	al functioning (range of score	s: 0-100; Better in	dicated by higher values					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	75	75	-	MD 18.4 higher (14.8 to 22 higher)	⊕⊕OO LOW	CRITICA
Quality	of life, SF-36	0-100 (≤ 4	months) - Vital	ity (range of sco	ores: 0-100; Be	etter indicated by	higher values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	75	75	-	MD 15.2 higher (11.09 to 19.31 higher)	⊕⊕OO LOW	CRITICA
Quality	of life, SF-36	0-100 (> 4	months – 1 yea	ır) - Physical fu	nctioning (rang	ge of scores: 0-10	0; Better indicated by hig	her values)				•
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	75	75	-	MD 27.6 higher (24.64 to 30.56 higher)	⊕⊕OO LOW	CRITICA
Quality	of life, SF-36	0-100 (> 4	months – 1 yea	ır) - Emotional r	ole (range of s	scores: 0-100; Bet	ter indicated by higher va	lues)	•			
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	75	75	-	MD 34.4 higher (28.87 to 39.93 higher)	⊕⊕OO LOW	CRITICA
Quality	of life, SF-36	0-100 (> 4	months – 1 yea	ır) - General hea	alth (range of s	scores: 0-100; Bet	ter indicated by higher va	lues)				

1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	75	75	-	MD 25.9 higher (21.93 to 29.87 higher)	⊕⊕OO LOW	CRITICAL
Quality	of life, SF-36	0-100 (> 4	months- 1 year) - Mental healt	h (range of sco	ores: 0-100; Bette	r indicated by higher value	es)				
1		very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	75	75	-	MD 25.5 higher (22.13 to 28.87 higher)	⊕⊕OO LOW	CRITICAL
Quality	of life, SF-36	0-100 (> 4	months- 1 year) - Physical pai	n (range of sco	ores: 0-100; Bette	r indicated by higher value	es)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	75	75	-	MD 27 higher (22.68 to 31.32 higher)	⊕⊕OO LOW	CRITICAL
Quality	of life, SF-36	0-100 (> 4	l months- 1 year) - Physical role	e (range of sco	ores: 0-100; Better	r indicated by higher value	s)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	75	75	-	MD 25.8 higher (20.96 to 30.64 higher)	⊕⊕OO LOW	CRITICAL
Quality	of life, SF-36	0-100 (> 4	months- 1 year) - Social functi	ioning (range o	of scores: 0-100; I	Better indicated by higher	values)				
1		very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	75	75	-	MD 22.7 higher (19.08 to 26.32 higher)	⊕⊕OO LOW	CRITICAL
Quality	of life, SF-36	0-100 (> 4	months- 1 year) - Vitality (rang	je of scores: 0	-100; Better indic	ated by higher values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	75	75	-	MD 23 higher (19.36 to 26.64 higher)	⊕⊕OO LOW	CRITICAL

 ^a Downgraded by 2 increments if the majority of the evidence was at very high risk of bias
 ^b Downgraded by 1 increment if the majority of the evidence was at high risk of bias
 ^c Downgraded by 1 increment if the confidence interval crossed one MID

Table 294: MBR programme 2 elements: physical + psychological vs. Usual care/waiting list control

	•				<u>, , </u>							
			Quality as	sessment			No of patie	ents		Effect	Quality	l
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	MBR programme 2 elements: physical + psychological	Usual care/waiting list control	Relative (95% CI)	Absolute	Quality	Importance
Pain sev	n severity, VAS 0-10 (> 4 months)(follow-up >4 months – 1 year; range of scores: 0-10; Better indicated by lower values)											
1	randomised trials			no serious indirectness	Serious ^b	none	56	50	-	MD 0.82 lower (1.64 lower to 0.00 higher)	⊕⊕OO LOW	CRITICAL
Function	, Roland-Mo	rris (> 4 n	months)(follow-u	p >4 months –	1 year; range	of scores: 0-24; I	Setter indicated by low	er values)				
	randomised trials	Serious ^a			no serious imprecision	none	56	50	-	MD 2.56 lower (4.27 to 0.85 lower)	⊕⊕⊕O MODERATE	CRITICAL
Psycholo	ogical distres	ss, BDI 0-	-63 (>4 months)(follow-up >4 m	onths; range o	of scores: 0-63; B	etter indicated by lowe	r values)				
1	randomised trials			no serious indirectness	very serious ^c	none	56	50	-	MD 0.04 higher (1.71 lower to 1.79 higher)	⊕000 VERY LOW	CRITICAL
Return to	o work (>4 m	onths)(fo	llow-up >4 mont	ths)								
1	randomised trials	,		no serious indirectness	Serious ^b	none	20/22 (90.9%)	68.8%	RR 1.32 (1.05 to 1.67)	220 more per 1000 (from 34 more to 461 more)	⊕000 VERY LOW	IMPORTANT

 ^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias
 ^b Downgraded by 1 increment if the confidence interval crossed one MID
 ^c Downgraded by 2 increments if the confidence interval crossed both MIDs

NICE, 2016

Table 295: MBR programme 2 elements: physical + psychological vs. Single intervention (mixed modality exercise; individual biomechanical exercise; psychological – cognitive behavioural approaches)

				navioural ap	, ,							
			Quality as:	sessment			No of patier	nts		Effect	Quality.	I
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	MBR programme 2 elements: physical + psychological	Single intervention	Relative (95% CI)	Absolute	Quality	Importance
Pain seve	erity, VAS 0-	10 (≤4 mo	onths) - Mixed mo	odality exercise	(follow-up <4	months; range o	f scores: 0-10; Better ir	ndicated by lov	ver values)			
	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	27	27	-	MD 2.59 lower (3.28 to 1.9 lower)	⊕⊕OO LOW	CRITICAL
Pain seve	erity, VAS 0-	10 (≤4 mo	onths) - Mixed mo	odality exercise	(aerobic + bic	omechanical) (fol	low-up <4 months; rang	ge of scores: 0	-10; Better	indicated by lowe	er values)	
	randomised trials		no serious inconsistency	no serious indirectness	Serious ^c	none	55	52	-	MD 0.02 higher (0.88 lower to 0.92 higher)	⊕⊕OO LOW	CRITICAL
Pain seve	erity, VAS 0-	10 (≤4 mo	onths) - Psycholo	gical - cognitiv	e behavioural	approaches(folio	ow-up <4 months; range	e of scores: 0-	10; Better ir	ndicated by lower	· values)	
	randomised trials		no serious inconsistency	no serious indirectness	Serious ^c	none	55	55	-	MD 0.53 lower (1.42 lower to 0.35 higher)	⊕⊕OO LOW	CRITICAL
Pain seve	erity, VAS 0-	10 (>4 mc	onths) - Individua	I biomechanica	al exercise (fol	low-up >4 month	s; range of scores: 0-10); Better indica	ated by low	er values)		
			no serious inconsistency	no serious indirectness	Serious ^c	none	64	48	-	MD 0.70 lower (1.61 lower to 0.21 higher)	⊕OOO VERY LOW	CRITICAL
Pain severity, VAS 0-10 (>4 months) - Mixed modality exercie (aerobic + biomechanical) (follow-up >4 months; range of scores: 0-10; Better indicated by lower val											values)	
	randomised trials		no serious inconsistency	no serious indirectness	Serious ^c	none	53	51	-	MD 0.80 lower (1.71 lower to 0.1 higher)	⊕⊕OO LOW	CRITICAL
Pain seve	erity, VAS 0-	10 (>4 mc	onths) - Psycholo	gical - cognitiv	e behavioural	approaches(folio	ow-up >4 months; range	e of scores: 0-	10; Better in	ndicated by lower	values)	

1	randomised trials	Serious ^b	no serious inconsistency	no serious indirectness	Serious ^c	none	53	52	-	MD 0.89 lower (1.79 lower to 0.02 higher)	⊕⊕OO LOW	CRITICAL
Function	n, RMDQ 0-24	(≤4 mon	ths) - Psycholog	ical - cognitive	behavioural a	pproaches(follow	y-up <4 months; range o	of scores: 0-24	; Better ind	icated by lower v	/alues)	
1	randomised trials	Serious⁵	no serious inconsistency	no serious indirectness	Serious ^c	none	55	55	-	MD 0.57 lower (2.26 lower to 1.12 higher)	⊕⊕OO LOW	CRITICAL
Function	n, RMDQ 0-24	(≤4 mon	ths) - Mixed mod	dality exercise (aerobic + bion	nechanical) (follo	w-up <4 months; range	of scores: 0-2	4; Better in	dicated by lower	values)	
1	randomised trials	Serious ^b	no serious inconsistency	no serious indirectness	no serious imprecision	none	55	52	-	MD 0.05 higher (1.68 lower to 1.78 higher)	⊕⊕⊕O MODERATE	CRITICAL
Function	n, RMDQ 0-24	(>4 mon	ths) - Psycholog	jical - cognitive	behavioural a	pproaches(follow	r-up >4 months; range o	of scores: 0-24	; Better ind	icated by lower v	/alues)	
2	randomised trials	Serious ^b	no serious inconsistency	no serious indirectness	Serious ^c	none	109	104	-	MD 1.44 lower (2.64 to 0.24 lower)	⊕⊕OO LOW	CRITICAL
Function	n, RMDQ 0-24	(>4 mon	ths) - Mixed mod	dality exercise (aerobic + bion	nechanical) (follo	w-up >4 months; range	of scores: 0-2	4; Better in	dicated by lower	values)	
2	randomised trials	Serious ^b	no serious inconsistency	no serious indirectness	Serious ^c	none	109	103	-	MD 1.19 lower (2.43 lower to 0.04 higher)	⊕⊕OO LOW	CRITICAL
Function	n, RMDQ 0-24	. (≤4 mon	ths) - Mixed mod	dality exercise (aerobic + bion	nechanical) (follo	w-up <4 months; range	of scores: 0-2	4; Better in	dicated by lower	values)	
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	27	27	-	MD 4.55 lower (5.77 to 3.33 lower)	⊕⊕OO LOW	CRITICAL
Psychol	logical distres	ss, BDI 0-	63 (≤4 months) -	- Psychological	- cognitive be	havioural approa	ches(follow-up <4 mont	hs; range of s	cores: 0-63	; Better indicated	d by lower va	lues)
1	randomised trials	Serious ^b	no serious inconsistency	no serious indirectness	Serious ^c	none	55	55	-	MD 1.62 lower (3.56 lower to 0.32 higher)	⊕⊕OO LOW	CRITICAL
Psychol	logical distres	ss, BDI 0-	63 (>4 months) -	- Psychological	- cognitive be	havioural approa	ches(follow-up >4 mont	ths; range of s	cores: 0-63	; Better indicated	d by lower va	lues)
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	53	52	-	MD 0.09 higher (1.88 lower to	⊕⊕⊕O MODERATE	CRITICAL

										2.06 higher)		
Psychol	ogical distres	s BDI 0-	63 (<4 months) -	Mixed modality	v exercise (aeı	robic + biomechar	nical) (follow-up <4 mor	oths: range of	scores: 0-6	3. Better indicate	ed by lower v	ralues)
1	randomised trials			no serious indirectness	Serious ^c	none	53	52	-	MD 2.17 lower (4.13 to 0.21 lower)	⊕⊕OO LOW	CRITICAL
Psychol	ogical distres	ss, BDI 0-	63 (>4 months) -	Mixed modality	y exercise (aeı	robic + biomechai	nical) (follow-up >4 mor	nths; range of	scores: 0-6	3; Better indicate	ed by lower v	values)
1	randomised trials	Serious ^b	no serious inconsistency	no serious indirectness	Serious ^c	none	53	51	-	MD 1.06 lower (3.04 lower to 0.92 higher)	⊕⊕OO LOW	CRITICAL
Psychol	ogical distres	ss, HADS	0-21 (>4 months	s) - individual bi	iomechanical	exercise (follow-u	p >4 months; range of	scores: 0-21; I	Better indic	ated by lower val	lues)	
1	randomised trials	very serious ^b	no serious inconsistency	no serious indirectness	Serious ^c	none	42	41	-	MD 0.7 lower (3.63 lower to 2.23 higher)	⊕000 VERY LOW	CRITICAL
Healthca	are utilisation	, number	of GP visits (>4	months) - mixe	ed modality ex	ercise (aerobic +	biomechanical) (follow-	-up >4 months	; Better ind	icated by lower v	values)	
1	randomised trials	Serious ^b	no serious inconsistency	no serious indirectness	Serious ^c	none	56	52	-	MD 0.87 lower (2.52 lower to 0.78 higher)	⊕⊕OO LOW	IMPORTANT
Healthca	are utilisation	, number	of medical spec	ialist visits (>4	months) - mix	ed modality exerc	cise (aerobic + biomech	nanical) (follov	∕-up >4 mor	nths; Better indic	ated by lowe	er values)
1	randomised trials	Serious ^b	no serious inconsistency	no serious indirectness	no serious imprecision	none	56	52	-	MD 0.15 lower (1.18 lower to 0.88 higher)	⊕⊕⊕O MODERATE	IMPORTANT
Healthca	are utilisation	, number	of radiology vis	its (>4 months)	- mixed moda	ality exercise (aero	obic + biomechanical) (follow-up >4 n	nonths; Bet	ter indicated by I	ower values)
1	randomised trials	Serious ^b	no serious inconsistency	no serious indirectness	Serious ^c	none	56	52	-	MD 0.20 higher (0.19 lower to 0.59 higher)	⊕⊕OO LOW	IMPORTANT
Healthca	are utilisation	, number	of occupational	physician visit	ts (>4 months)	- mixed modality	exercise (aerobic + bio	mechanical) (follow-up >	4 months; Better	indicated by	lower
1	randomised trials	Serious ^b	no serious inconsistency	no serious indirectness	no serious imprecision	none	56	52	-	MD 0.02 higher (0.15 lower to 0.19 higher)	⊕⊕⊕O MODERATE	IMPORTANT

Healthca	re utilisation	, number	of psychologist	visits (>4 mon	ths) - mixed m	odality exercise (aerobic + biomechanic	al) (follow-up	>4 months;	Better indicated	by lower val	ues)
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	56	52	-	MD 0.23 lower (1.14 lower to 0.68 higher)	⊕⊕⊕O MODERATE	IMPORTANT
Healthca	re utilisation	, number	of therapist ses	sions (>4 mont	hs) - mixed me	odality exercise (a	aerobic + biomechanica	ıl) (follow-up >	4 months;	Better indicated b	oy lower valu	ies)
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^c	none	56	52	-	MD 2.95 higher (4.17 lower to 10.07 higher)	⊕⊕OO LOW	IMPORTANT
Healthca	re utilisation	, number	of alternative th	erapist visits (>	4 months) - m	ixed modality ex	ercise (aerobic + biome	chanical) (foll	ow-up >4 m	onths; Better ind	icated by lov	wer values)
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^c	none	56	52	-	MD 1.32 higher (2.15 lower to 4.79 higher)	⊕⊕OO LOW	IMPORTANT
Healthca	re utilisation	, number	of GP visits (>4	months) - psyc	chological (cog	nitive behaviour	al approaches) (follow-	up >4 months;	Better indi	cated by lower va	alues)	
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^c	none	56	52	-	MD 1.17 lower (2.58 lower to 0.24 higher)	⊕⊕OO LOW	IMPORTANT
Healthca	re utilisation	, number	of medical spec	ialist care visit	s (>4 months)	- psychological (cognitive behavioural a	pproaches) (fo	ollow-up >4	months; Better i	ndicated by	lower
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^c	none	56	52	-	MD 0.43 higher (0.44 lower to 1.3 higher)	⊕⊕OO LOW	IMPORTANT
Healthca	re utilisation	, number	of radiology vis	its (>4 months)	- psychologic	al (cognitive beh	avioural approaches) (f	ollow-up >4 m	onths; Bett	er indicated by lo	wer values)	
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	56	52	-	MD 0.10 higher (0.31 lower to 0.51 higher)	⊕⊕⊕O MODERATE	IMPORTANT
Healthca	re utilisation	, number	of occupational	physician visit	ts (>4 months)	- psychological (cognitive behavioural a	pproaches) (f	ollow-up >4	months; Better i	ndicated by	lower
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^c	none	56	52	-	MD 0.12 lower (0.41 lower to 0.17 higher)	⊕⊕OO LOW	IMPORTANT

Healthca	re utilisation	, number	of psychologist	visits (>4 mont	ths) - psycholo	gical (cognitive	behavioural approaches	s) (follow-up >	4 months;	Setter indicated I	y lower valu	es)
1	randomised trials			no serious indirectness	no serious imprecision	none	56	52	-	MD 0.05 higher (0.42 lower to 0.52 higher)	⊕⊕⊕O MODERATE	IMPORTANT
Healthca	re utilisation	, number	of therapist visi	ts (>4 months)	- psychologica	al (cognitive beha	avioural approaches) (fo	ollow-up >4 m	onths; Bette	er indicated by lo	wer values)	
1	randomised trials				no serious imprecision	none	56	52	-	MD 1.67 lower (9.97 lower to 6.63 higher)	⊕⊕⊕O MODERATE	IMPORTANT
Healthca	re utilisation	, number	of alternative th	erapist visits (>	-4 months) - ps	sychological (cog	gnitive behavioural appr	oaches) (folio	ow-up >4 m	onths; Better ind	icated by low	ver values)
1	randomised trials			no serious indirectness	Serious ^c	none	56	52	-	MD 1.67 higher (1.67 lower to 5.01 higher)	⊕⊕OO LOW	IMPORTANT
Return to	o work < 4 m	onths							•		•	
1		,	no serious inconsistency	no serious indirectness	Serious ^c	none	27/39 (69.2%)	0%	RR 1.04 (0.76 to 1.42)	-	⊕000 VERY LOW	IMPORTANT
Return to	o work > 4 m	onths										
1		- ,	no serious inconsistency	no serious indirectness	Serious ^c	none	60/64 (93.8%)	85.40%	RR 1.10 (0.96 to 1.25)	85 more per 1000 (from 34 fewer to 214 more)	⊕OOO VERY LOW	IMPORTANT

 ^a Downgraded by two increments if the majority of evidence was at very high risk of bias
 ^b Downgraded by 1 increment if the majority of the evidence was at high risk of bias
 ^c Downgraded by 1 increment if the confidence interval crossed one MID

Table 296: MBR programme 2 elements: physical + psychological vs. Combined intervention

			Quality as	sessment			No of patie	nts	I	Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	MBR programme 2 elements: physical + psychological	Combined intervention	Relative (95% CI)	Absolute	quanty	importance
Pain sev	erity, NRS 0-	10 (≤4 mo	nths) - Exercise	(biomechanica) + manual the	erapy (mobilisatio	n) (follow-up ≤4 months	; range of scor	es: 0-10; B	etter indicated b	y lower valu	es)
	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	45	45	-	MD 2.27 lower (2.74 to 1.8 lower)	⊕⊕⊕O MODERATE	CRITICAL
Pain sevo	erity, NRS 0-	10 (≤4 mo	nths) - Exercise	(biomechanica) + manual the	erapy (mobilisatio	on + manipulation) (follo	w-up ≤4 month	s; range of	scores: 0-10; B	etter indicate	ed by lower
	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	96	88	-	MD 2.22 lower (2.62 to 1.83 lower)	⊕⊕⊕O MODERATE	CRITICAL
	erity, NRS 0- dicated by lo			(biomechanica) + manual the	erapy (mobilisatio	on) + postural therapy (p	ostural control) (follow-up	o ≤4 months; rar	ge of scores	s: 0-10;
	randomised trials		no serious inconsistency	no serious indirectness	Serious°	none	10	10	-	MD 1 lower (2.39 lower to 0.39 higher)	⊕000 VERY LOW	CRITICAL
Pain sev	erity, NRS 0-	10 (> 4 mo	onths)- Exercise	(biomechanica) + manual the	erapy (mobilisatio	pn) (follow-up >4 months	- 1 year; rang	e of scores	: 0-10; Better in	dicated by lo	wer values)
	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	45	45	-	MD 3.95 lower (4.42 to 3.48 lower)	⊕⊕⊕O MODERATE	CRITICAL
Pain seve		10 (> 4 mo	onths)- Exercise	(biomechanica) + manual the	erapy (mobilisatio	on + manipulation) (follo	w-up >4 month	s – 1 year;	range of scores	: 0-10; Better	r indicated
	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious°	none	51	43	-	MD 1.50 lower (2.33 to 0.67 lower)	⊕⊕OO LOW	CRITICAL
Function	, RMDQ 0-24	(≤4 mont	hs) - Exercise (b	iomechanical) +	+ manual thera	apy (mobilisation)	(follow-up ≤4 months; ı	ange of scores	s: 0-24; Bet	ter indicated by	lower values	s)

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1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	45	45	-	MD 6.0 lower (6.89 to 5.11 lower)	⊕⊕⊕O MODERATE	CRITICAL
Function values)	ı, ODI 0-100 (≤4 month	s) - Exercise (bio	omechanical) +	manual therap	y (mobilisation +	manipulation) (follow-u	p ≤4 months; ra	inge of sco	ores: 0-100; Bett	er indicated l	by lower
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	51	43	-	MD 10.90 lower (13.94 to 7.86 lower)	⊕⊕⊕O MODERATE	CRITICAL
	ı, ODI 0-100 (d by lower va		s) - Exercise (bio	omechanical) +	manual therap	y (mobilisation) +	- postural therapy (postu	ural control) (fo	llow-up ≤4	months; range	of scores: 0-	100; Better
1	randomised trials	very serious ^b	no serious inconsistency	no serious indirectness	no serious imprecision	none	10	10	-	MD 7 lower (11.16 to 2.84 lower)	⊕⊕OO LOW	CRITICAL
Function	ı, RMDQ 0-24	(> 4 mon	ths)- Exercise (b	iomechanical)	+ manual thera	apy (mobilisation)	(follow-up >4 months –	1 year; range o	of scores: ()-24; Better indi	cated by lowe	er values)
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	45	45	-	MD 9.69 lower (10.44 to 8.94 lower)	⊕⊕⊕O MODERATE	CRITICAL
Function lower va		> 4 montl	ns)- Exercise (bio	omechanical) +	manual therap	y (mobilisation +	manipulation) (follow-u	p >4 months – '	1 year; ran	ge of scores: 0-	100; Better in	dicated by
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^c	none	51	43	-	MD 9.80 lower (14.21 to 5.39 lower)	⊕⊕OO LOW	CRITICAL
	of life, SF-36 of by higher v		months) - physic	cal functioning	- Exercise (bio	mechanical) + ma	anual therapy (mobilisati	ion) (follow-up	≤4 months	; range of score	s: 0-100; Bet	ter
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	45	45	-	MD 21.00 higher (12.78 to 29.22 higher)	⊕⊕⊕O MODERATE	CRITICAL
			months) - physic indicated by higl		- Exercise (bio	mechanical) + ma	anual therapy (mobilisati	ion) + postural	therapy (po	ostural control)	(follow-up ≤4	months;
1	randomised trials	very serious ^b	no serious inconsistency	no serious indirectness	no serious imprecision	none	10	10	-	MD 17 higher (9.77 to 24.23 higher)	⊕⊕OO LOW	CRITICAL

1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	45	45	-	MD 21.33 higher (9.49 to 33.17 higher)	⊕⊕⊕O MODERATE	CRITICAL
			months) - emot ted by higher va		rcise (biomech	anical) + manua	I therapy (mobilisation)	+ postural thera	py (postura	al control) (follow	-up ≤4 mont	:hs; range
1	randomised trials	very serious ^b	no serious inconsistency	no serious indirectness	no serious imprecision	none	10	10	-	MD 20 higher (5.98 to 34.02 higher)	⊕⊕OO LOW	CRITICA
	ry of life, SF-36 r values)	0-100 (≤4	months) - gene	ral health - Exe	cise (biomech	anical) + manua	l therapy (mobilisation)	(follow-up ≤4 m	onths; rang	e of scores: 0-10	0; Better inc	licated by
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^c	none	45	45	-	MD 29.00 higher (21.82 to 36.18 higher)	⊕⊕OO LOW	CRITICA
			months) - gene ted by higher va		cise (biomech	anical) + manua	l therapy (mobilisation)	+ postural thera	py (postura	al control) (follow	-up ≤4 mont	hs; range
1	randomised trials	very serious ^b	no serious inconsistency	no serious indirectness	no serious imprecision	none	10	10	-	MD 16 higher (10.15 to 21.85 higher)	⊕⊕OO LOW	CRITICA
	y of life, SF-36 r values)	0-100 (≤4	months) - ment	al health - Exer	cise (biomecha	nical) + manual	therapy (mobilisation)	(follow-up ≤4 mo	nths; range	e of scores: 0-100	; Better ind	cated by
			no serious	no serious	no serious imprecision	none	45	45	-	MD 26.31 higher (20.84 to 31.78 higher)	⊕⊕⊕O MODERATE	CRITICA
	randomised trials	Serious	inconsistency	indirectness	,					31.70 Higher)		
nighe I Qualit	trials by of life, SF-36	0-100 (≤4	,	al health - Exerc		nnical) + manual	therapy (mobilisation)	+ postural therap	y (postura	, ,	up ≤4 montl	ns; range o

i e			•					1		T		
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	45	45	-	MD 24.36 higher (18 to 30.72 higher)	⊕⊕⊕O MODERATE	CRITICAL
			months) - physic		ise (biomecha	nical) + manual th	nerapy (mobilisation) + p	ostural therapy	(postural	control) (follow	-up ≤4 month	s; range of
1	randomised trials	very serious ^b	no serious inconsistency	no serious indirectness	Serious ^c	none	10	10	-	MD 10 higher (1.39 to 18.61 higher)	⊕OOO VERY LOW	CRITICAL
Quality o		0-100 (≤4	months) - physic	cal role - Exerci	se (biomechar	nical) + manual th	erapy (mobilisation) (fol	low-up ≤4 mon	ths; range	of scores: 0-100	; Better indic	ated by
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	45	45	-	MD 21.66 higher (9.83 to 33.49 higher)	⊕⊕⊕O MODERATE	CRITICAL
			months) - physic l by higher value		se (biomechar	nical) + manual th	erapy (mobilisation) + p	ostural therapy	(postural	control) (follow-	up ≤4 month	s; range of
1	randomised trials	very serious ^b	no serious inconsistency	no serious indirectness	no serious imprecision	none	10	10	-	MD 21 higher (8.97 to 33.03 higher)	⊕⊕OO LOW	CRITICAL
Quality o		0-100 (≤4	months) - social	functioning - E	xercise (biom	echanical) + manı	ual therapy (mobilisation	n) (follow-up ≤4	months; ra	ange of scores:	0-100; Better	indicated
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	45	45	-	MD 22.77 higher (15.96 to 29.58 higher)	⊕⊕⊕O MODERATE	CRITICAL
			months) - social		xercise (biom	echanical) + manı	ual therapy (mobilisation	n) + postural the	erapy (post	ural control) (fo	llow-up ≤4 m	onths;
1	randomised trials	very serious ^b	no serious inconsistency	no serious indirectness	no serious imprecision	none	10	10	-	MD 20 higher (13.86 to 26.14 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life, SF-36	0-100 (≤4	months) - vitality	/ - Exercise (bid	omechanical) +	manual therapy	(mobilisation) (follow-up	o ≤4 months; ra	nge of sco	res: 0-100; Bette	er indicated b	y higher
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^c	none	45	45	-	MD 25.33 higher (19.01 to	⊕⊕OO LOW	CRITICAL

										31.65 higher)		
			months) - vitalit		omechanical) +	- manual therapy	(mobilisation) + postura	I therapy (post	ural contro	ol) (follow-up ≤4 n	nonths; ranç	ge of
1	randomised trials	very serious ^b	no serious inconsistency	no serious indirectness	no serious imprecision	none	10	10	-	MD 20 higher (11.57 to 28.43 higher)	⊕⊕OO LOW	CRITICA
	of life, SF-36 ed by higher v		months)- physic	cal functioning	- Exercise (bio	mechanical) + ma	nual therapy (mobilisat	ion) (follow-up	>4 months	s – 1 year; range	of scores: 0-	-100; Bette
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	45	45	-	MD 23.56 higher (15.49 to 31.63 higher)	⊕⊕⊕O MODERATE	CRITICA
	of life, SF-36 ed by higher v		months)- emoti	onal role - Exer	cise (biomech	anical) + manual t	herapy (mobilisation) (f	ollow-up >4 mo	nths – 1 ye	ear; range of sco	res: 0-100; E	Better
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	45	45	-	MD 32.59 higher (26.52 to 38.66 higher)	⊕⊕⊕O MODERATE	CRITICA
	/ of life, SF-36 ted by higher v		months)- gener	al health - Exer	cise (biomech	anical) + manual t	herapy (mobilisation) (fo	ollow-up >4 mo	nths – 1 ye	ear; range of sco	res: 0-100; E	Better
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^c	none	45	45	-	MD 28.56 higher (22.41 to 34.71 higher)	⊕⊕OO LOW	CRITICA
	of life, SF-36 ed by higher v		months)- menta	al health - Exerc	cise (biomecha	nical) + manual th	nerapy (mobilisation) (fo	llow-up >4 mor	nths – 1 ye	ar; range of scor	es: 0-100; B	etter
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious°	none	45	45	-	MD 35.65 higher (30.5 to 40.8 higher)	⊕⊕OO LOW	CRITICA
Qualit	of life, SF-36 ted by higher v		months)- physic	cal pain- Exerc	se (biomechar	nical) + manual the	erapy (mobilisation) (fol	low-up >4 mon	ths – 1 yea	r; range of score	s: 0-100; Be	tter
				no corious	no serious	none	45	45	-	MD 26.96	⊕⊕⊕О	CRITICAL
	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	imprecision					higher (20.57 to 33.35 higher)	MODERATE	

indicate	d by higher v	alues)										
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	45	45	-	MD 25.78 higher (17.85 to 33.71 higher)	⊕⊕⊕O MODERATE	CRITICAL
-	of life, SF-36 (d by higher v	•	months)- social	functioning - E	exercise (biom	echanical) + manu	ual therapy (mobilisation	n) (follow-up >4	l months –	1 year; range of	scores: 0-10	0; Better
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^c	none	45	45	-	MD 36.56 higher (32.05 to 41.07 higher)	⊕⊕OO LOW	CRITICAL
Quality of higher v		0-100 (> 4	months)- vitality	y - Exercise (bio	omechanical) +	- manual therapy	(mobilisation) (follow-up	>4 months –	1 year; ranç	ge of scores: 0-1	00; Better in	dicated by
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	no serious imprecision	none	45	45	-	MD 34.67 higher (29.98 to 39.36 higher)	⊕⊕⊕O MODERATE	CRITICAL
	are utilisation d by lower va		eking after interv	rention (> 4 mor	nths)- Exercise	(biomechanical)	+ manual therapy (mani	pulation + mol	oilisation) (follow-up >4 mo	nths – 1 year	; Better
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^c	none	51	43	-	MD 8.50 lower (12.74 to 4.26 lower)	⊕⊕OO LOW	CRITICAL
Healthca	re utilisation	, medicin	e use (≤4 month	s) - Exercise (bi	iomechanical)	+ manual therapy	(mobilisation) + postur	al therapy (pos	tural contr	ol) (follow-up >4	months)	
1		very serious ^b	no serious inconsistency	no serious indirectness	Serious ^c	none	0/10 (0%)	0%	RR 0.07 (0 to 1.03)	-	⊕OOO VERY LOW	CRITICAL

 ^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias
 ^b Downgraded by 2 increments if the majority of the evidence was at very high risk of bias
 ^c Downgraded by 1 increment if the confidence interval crossed one MID

Table 297: MBR programme 2 elements: physical + education vs. Single intervention (biomechanical exercise)

			Quality as	sessment		-	No of patie	nts		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	MBR programme 2 elements: physical + education	Single intervention	Relative (95% CI)	Absolute	quanty	importunio
Pain sev	erity, VAS 0-1	0 (≤4 mor	nths) (follow-up <	4 months; range	e of scores: 0-1	0; Better indicated	d by lower values)					
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	129	143	-	MD 0.53 higher (0.05 lower to 1.11 higher)	⊕OOO VERY LOW	CRITICAL
Pain seve	erity, VAS 0-1	0 (>4 moı	nths) - Biomechai	nical exercise (f	ollow-up >4 mo	enths; range of sco	ores: 0-10; Better indica	ted by lower v	alues)			
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	129	143	-	MD 0.66 higher (0.09 to 1.23 higher)	⊕000 VERY LOW	CRITICAL
Function	, RMDQ 0-24	(>4 montl	hs) - Biomechanio	cal exercise - co	re stability (foll	ow-up <4 months	; Better indicated by lov	ver values)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	129	143	-	MD 2.10 higher (0.81 to 3.39 higher)	⊕000 VERY LOW	CRITICAL
Quality o	f life, SF-36 (:	≤4 months	s) - physical func	tioning (follow-ı	ıp <4 months; ı	ange of scores: 0	-100; Better indicated b	y higher values	s)		,	
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	129	143	-	MD 6.20 higher (1.53 to 10.87 higher)	⊕⊕OO LOW	CRITICAL
Function	, RMDQ 0-24	(≤4 montl	hs) - Biomechanio	cal exercise - co	re stability (foll	ow-up <4 months	; range of scores: 0-24;	Better indicate	d by low	er values)		
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	129	143	-	MD 1.5 higher (0.34 to 2.66 higher)	⊕000 VERY LOW	CRITICAL
Quality o	f life, SF-36 (:	≤4 months	s) - emotional role	e (follow-up <4 r	months; Better	indicated by high	er values)					
1	randomised	very	no serious	no serious	very serious ^c	none	129	143	-	MD 3.10 higher (7	⊕OOO	CRITICAL

	trials	serious ^a	inconsistency	indirectness						lower to 13.2 higher)	VERY LOW	
Quality o	of life, SF-36 (:	≤4 month	s) - general healt	h (follow-up <4 ı	months; range	of scores: 0-100; E	Better indicated by highe	er values)				
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	129	143	-	MD 1.29 lower (5.69 lower to 3.11 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life, SF-36 (≤4 month	s) - mental health	n (follow-up <4 n	nonths; range o	f scores: 0-100; B	etter indicated by highe	r values)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	129	143	-	MD 0.10 lower (4.75 lower to 4.55 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life, SF-36 (:	≤4 month:	s) - physical pain	(follow-up <4 m	nonths; range o	f scores: 0-100; B	etter indicated by highe	r values)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	129	143	-	MD 5.70 higher (0.61 to 10.79 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life, SF-36 (≤4 month	s) - physical role	(follow-up <4 m	onths; Better ir	ndicated by higher	values)					
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	very serious ³	none	129	143	-	MD 3.2 higher (5.75 lower to 12.15 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	of life, SF-36 (:	≤4 month	s) - social function	oning (follow-up	<4 months; ran	ge of scores: 0-10	00; Better indicated by h	igher values)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	129	143	-	MD 0.40 higher (5.08 lower to 5.88 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life, SF-36 (≤4 month	s) - vitality (follow	v-up <4 months	; range of score	s: 0-100; Better in	dicated by higher value	s)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	129	143	-	MD 3.00 higher (2.04 lower to 8.04 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life, SF-36 (:	≤4 month	s) - physical com	ponent summa	ry score (follow	-up <4 months; ra	nge of scores: 0-100; Be	etter indicated	by highe	r values)		
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	129	143	-	MD 2.20 higher (0.41 to 3.99 higher)	⊕⊕OO LOW	CRITICAL

1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	129	143	-	MD 0.40 lower (2.89 lower to 2.09 higher)	⊕⊕OO LOW	CRITICA
Quality	y of life, SF-36 (>4 month	ıs) - physical fund	ctioning (follow	-up >4 months;	range of scores:	0-100; Better indicated b	y higher values	s)	•		
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	129	143	-	MD 10.10 higher (4.92 to 15.28 higher)	⊕⊕OO LOW	CRITICA
Quality	y of life, SF-36 (>4 month	s) - emotional ro	le (follow-up >4	months; range	of scores: 0-100;	Better indicated by high	er values)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	129	143	-	MD 8.30 higher (2.82 lower to 19.42 higher)	⊕⊕OO LOW	CRITICA
Quality	y of life, SF-36 (>4 month	s) - general healt	th (follow-up >4	months; range	of scores: 0-100;	Better indicated by high	er values)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	129	143	-	MD 2.34 lower (6.47 lower to 1.79 higher)	⊕⊕OO LOW	CRITICA
Quality	y of life, SF-36 (>4 month	s) - mental healtl	h (follow-up >4	months; range	of scores: 0-100; I	Better indicated by highe	er values)	_			
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	129	143	-	MD 2.90 higher (2.07 lower to 7.87 higher)	⊕⊕OO LOW	CRITICA
Quality	y of life, SF-36 (>4 month	s) - physical pair	າ (follow-up >4 ເ	months; range o	of scores: 0-100; E	Better indicated by highe	r values)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	129	143	-	MD 4.80 higher (0.42 lower to 10.02 higher)	⊕⊕OO LOW	CRITICA
Quality	y of life, SF-36 (>4 month	s) - physical role	(follow-up >4 n	nonths; range o	f scores: 0-100; B	setter indicated by highe	r values)				
	randomised	very serious ¹	no serious inconsistency	no serious	no serious	none	129	143	-	MD 8.30 higher (1.14 lower to	⊕⊕OO LOW	CRITICA

1	randomised trials	very serious¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	129	143	-	MD 4.40 higher (1.97 lower to 10.77 higher)	⊕⊕OO LOW	CRITICAL
Quality o	f life, SF-36 (>4 months	s) - vitality (follow	v-up >4 months;	range of score	s: 0-100; Better in	dicated by higher values	s)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	129	143	-	MD 6.50 higher (0.86 to 12.14 higher)	⊕⊕OO LOW	CRITICAL
Quality o	f life, SF-36 (>4 months	s) - physical com	ponent summar	y score (follow	-up >4 months; ra	nge of scores: 0-100; Be	tter indicated	by highe	r values)		
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	129	143	-	MD 3.20 higher (1.32 to 5.08 higher)	⊕⊕OO LOW	CRITICAL
Quality o	f life, SF-36 (>4 months	s) - mental compo	onent summary	score (follow-u	p >4 months; rang	ge of scores: 0-100; Bett	er indicated by	/ higher v	values)		
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	129	143	-	MD 1.60 higher (1.1 lower to 4.3 higher)	⊕⊕OO LOW	CRITICAL

¹ Downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Downgraded by 1 increment if the confidence interval crossed one MID ³ Downgraded by 2 increments if the confidence interval crossed both MIDs

Table 298: MBR programme 2 elements: physical (exercise + manipulation) + education vs. Single intervention (manual therapy - manipulation)

	·		Quality asse			,	No of patients			Effect		Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	2-MBR physical (manipulation + exercise) + education		Relative (95% CI)	Absolute	Quanty	importance	
Pain (McGill Present Pain Intensity 0-5) - <4 months (Better indicated by lower values)													
		, ,		no serious indirectness	serious²	none	24	22	-	MD 0.76 lower (1.43 to 0.09 lower)	⊕OOO VERY LOW	CRITICAL	

Pain (McC	Gill Pain Ratin	g Index 0	-79) - <4 months (Better indicated	by lower val	lues)							
1		very serious¹		no serious indirectness	serious ²	none	24	22		MD 2.26 lower (5.17 lower to 0.65 higher)		CRITICAL	
Disability (RMDQ 0-24) - <4 months (Better indicated by lower values)													
1	randomised trials	very serious ¹		no serious indirectness	serious ²	none	24	22	-	MD 1.32 lower (2.84 lower to 0.2 higher)	⊕OOO VERY LOW	CRITICAL	
Psychological distress (Anxiety, STAI 20-80) - <4 months (Better indicated by lower values)													
1	randomised trials	very serious¹		no serious indirectness	serious ²	none	24	22	-	MD 6.94 lower (11.31 to 2.57 lower)	⊕OOO VERY LOW	CRITICAL	

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Table 299: MBR programme 2 elements: physical (exercise) + education vs. Single intervention (manual therapy - manipulation)

			Quality asse	ssment			No of patients			Effect	Ovelite	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	2-MBR physical (ex) + education	Control	Relative (95% CI)	Absolute	Quanty	Importance
Pain (McG	ill Present Pai	n Intensity	/ 0-5) - <4 months (
		- ,		no serious indirectness	serious ²	none	21	22	-	MD 0.15 higher (0.56 lower to 0.86 higher)	⊕OOO VERY LOW	CRITICAL
Pain (McG	ill Pain Rating	Index 0-7	9) - <4 months (Co	py) (Better indica	ted by lower	values)						
				no serious indirectness	serious²	none	21	22	-	MD 0.64 higher (2.37 lower to 3.65 higher)	⊕OOO VERY LOW	CRITICAL

Disability	Disability (RMDQ 0-24) - <4 months (Copy) (Better indicated by lower values)														
1		- ,	no serious inconsistency	no serious indirectness	serious ²	none	21	22	-	MD 2.85 higher (0.42 to 5.28 higher)	⊕OOO VERY LOW	CRITICAL			
Psychological distress (Anxiety, STAI 20-80) - <4 months (Copy) (Better indicated by lower values)															
1		- ,	no serious inconsistency	no serious indirectness	serious ²	none	21	22	-	MD 1.92 lower (7.02 lower to 3.18 higher)	⊕OOO VERY LOW	CRITICAL			

Table 300: MBR programme 3 elements: physical + psychological (cognitive) + education vs. MBR programme 2 elements: physical + education

			Quality ass	essment			No of pati	ents		Effect	Qualifu	In a section of
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	MBR program 3 elements (psych=cognitive)	MBR program 2 elements: physical + education	Relative (95% CI)	Absolute	Quality	Importance
Pain Inte	nsity, pain ra	ting char	t (≤4 months) (fo	llow-up ≤4 mor	nths; measur	ed with: pain ration	ng chart; Better indicated	d by lower values)				
2		- ,	no serious inconsistency	no serious indirectness	serious²	none	17	18	1	MD 0.18 higher (0.33 lower to 0.69 higher)	⊕OOO VERY LOW	CRITICAL
Pain Inte	nsity, pain ra	ting char	t (> 4 months)(fo	ollow-up > 4 mo	nths; measu	red with: pain rati	ng chart; Better indicate	d by lower values)				
2		,	no serious inconsistency	no serious indirectness	serious ²	none	13	16	-	MD 0.34 higher (0.32 lower to 1 higher)	⊕OOO VERY LOW	CRITICAL
Psychological distress, BDI 0-63 (≤4 months) (follow-up ≤4 months; measured with: Beck Depression Inventory; Better indicated by lower values)												
2		,	no serious inconsistency	no serious indirectness	serious ²	none	17	18	-	MD 3.95 higher (0.31 lower to 8.2 higher)	⊕OOO VERY LOW	CRITICAL

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

Psychol	ogical distres	s, BDI 0-	63 (> 4 months)(follow-up > 4 m	onths; meas	sured with: Beck D	Depression Inventory ; Be	etter indicated by lo	wer valu	es)				
2		very serious ¹	no serious inconsistency	no serious indirectness	very serious ³	none	15	17	-	MD 0.36 lower (5.21 lower to 4.48 higher)	⊕OOO VERY LOW	CRITICAL		
Psychol	ogical distres	s, State-	Trait Inventory: §	State (≤4 month	s) (follow-up	≤4 months; meas	sured with: State-Trait Inv	ventory: State ; Bett	er indica	ted by lower valu	es)			
1		very serious ¹	no serious inconsistency	no serious indirectness	very serious ³	none	8	9	-	MD 2.24 higher (9.18 lower to 13.66 higher)	⊕OOO VERY LOW	CRITICAL		
Psychol	ogical distres	s, State-	Trait Inventory: S	State (> 4 montl	ns)(follow-up	> 4 months; mea	sured with: State-Trait In	ventory: State ; Bet	ter indic	ated by lower valu	ies)			
1		very serious ¹	no serious inconsistency	no serious indirectness	very serious ³	none	6	9	-	MD 0.61 higher (14.94 lower to 16.16 higher)	⊕OOO VERY LOW	CRITICAL		
Function	Function, Sickness Impact Profile (≤4 months) (follow-up ≤4 months; measured with: Sickness Impact Profile ; Better indicated by lower values)													
2		very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	17	18	-	MD 3.23 lower (10.84 lower to 4.39 higher)	⊕OOO VERY LOW	CRITICAL		
Function	n, Sickness In	npact Pro	ofile (> 4 months))(follow-up > 4	months; mea	sured with: Sickn	ness Impact Profile ; Bette	er indicated by lowe	er values)				
2		very serious ¹	no serious inconsistency	no serious indirectness	very serious ³	none	15	17	-	MD 1.95 lower (10.02 lower to 6.11 higher)	⊕OOO VERY LOW	CRITICAL		
Medicati	ion use (≤4 m	onths) (fe	ollow-up ≤4 mon	ths; Better indi	cated by low	er values)								
1		very serious ¹	no serious inconsistency	no serious indirectness	very serious ³	none	8	9	-	MD 0.02 higher (0.96 lower to 1 higher)	⊕OOO VERY LOW	IMPORTANT		
Medicat	ion use (> 4 m	nonths)(f	ollow-up >4 mon	ths - 1 year; Be	etter indicate	d by lower values	s)							
1		very serious ¹	no serious inconsistency	no serious indirectness	very serious ³	none	6	9	-	MD 0.23 higher (1.03 lower to 1.49 higher)	⊕OOO VERY LOW	IMPORTANT		

Table 301: MBR programme 3 elements: physical + psychological (behavioural) + education vs. MBR programme 2 elements: physical + education

	01. W.D.V P	. og. a	ine o cicinent	o. pilyoleal .	Payeriolog	icai (Bellavica	iai, · caacation voi ivib	n programme 2	· Cicilic	its. priysicar :	Caaca		
			Quality ass	essment			No of patie	nts		Effect			
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	MBR program 3 elements (psych=behavioural)	MBR program 2 elements: physical + education	Relative (95% CI)	Absolute	Quality	Importance	
Pain Inte	nin Intensity, pain rating chart (≤4 months) (follow-up ≤4 months; measured with: pain rating chart ; Better indicated by lower values)												
1	randomised trials	,	no serious inconsistency	no serious indirectness	serious ²	none	8	9	-	MD 0.8 lower (1.47 to 0.13 lower)	⊕OOO VERY LOW	CRITICAL	
Pain Inte	in Intensity, pain rating chart (> 4 months)(follow-up >4 months - 1 year; measured with: pain rating chart; Better indicated by lower values)												
		very serious ¹	no serious inconsistency	no serious indirectness	very serious ³	none	5	8	-	MD 0.14 lower (1.17 lower to 0.89 higher)	⊕OOO VERY LOW	CRITICAL	
Psycholo	sychological distress, BDI 0-63 (≤4 months) (follow-up ≤4 months; measured with: Beck Depression Inventory; Better indicated by lower values)												
	randomised trials	very serious¹	no serious inconsistency	no serious indirectness	serious ²	none	8	9	-	MD 5.02 higher (2.52 lower to 12.56 higher)	⊕000 VERY LOW	CRITICAL	
Psycholo	ogical distres	ss, BDI 0-	63 (> 4 months)(follow-up >4 m	onths - 1 yea	ar; measured witl	h: Beck Depression Invento	ry ; Better indicate	ed by low	er values)			
		very serious ¹	no serious inconsistency	no serious indirectness	very serious³	none	6	9	-	MD 8.11 higher (0.61 lower to 16.83 higher)	⊕000 VERY LOW	CRITICAL	
Psycholo	ogical distres	ss, State-	Trait Inventory: S	State (≤4 month	s) (follow-up	o ≤4 months; mea	sured with: State-Trait Inve	ntory: State ; Bette	er indicat	ed by lower valu	es)		
		very serious ¹	no serious inconsistency	no serious indirectness	very serious³	none	8	9	-	MD 1.49 higher (9.58 lower to	⊕000 VERY	CRITICAL	

¹ Downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Downgraded by 1 increment if the confidence interval crossed either the MID for benefit or the MID for harm ³ Downgraded by 2 increments if the confidence interval crossed both the MID for benefit and the MID for harm

										10 CC bimb :\	1014			
	<u> </u>	L								12.56 higher)	LOW	L		
Psycholo	ogical distres	s, State-	Trait Inventory: S	State (> 4 month	าร)(follow-up	> 4 months; mea	asured with: State-Trait Inve	entory: State ; Bett	er indica	ted by lower valu	ıes)			
		very serious¹			very serious³	none	6	9	-	MD 3.73 lower (14.38 lower to 6.92 higher)	⊕OOO VERY LOW	CRITICAL		
Function	, Sickness In	npact Pro	ofile (≤4 months)	(follow-up ≤4 n	nonths; mea	sured with: Sickn	ess Impact Profile ; Better i	ndicated by lower	values)					
		very serious ¹		no serious indirectness	serious²	none	8	9	-	MD 7.2 lower (17.52 lower to 3.12 higher)	⊕OOO VERY LOW	CRITICAL		
Function	unction, Sickness Impact Profile (> 4 months)(follow-up > 4 months; measured with: Sickness Impact Profile; Better indicated by lower values)													
	randomised trials		no serious inconsistency		very serious³	none	6	9	-	MD 4.91 higher (8.12 lower to 17.94 higher)	⊕OOO VERY LOW	CRITICAL		
Medicati	on use (≤4 m	onths) (fe	ollow-up ≤4 mon	ths; Better indi	cated by low	er values)								
	randomised trials		no serious inconsistency		very serious³	none	8	9	-	MD 0.02 higher (1.08 lower to 1.12 higher)	⊕000 VERY LOW	IMPORTANT		
Medicati	on use (> 4 m	nonths)(f	ollow-up > 4 mor	nths; Better ind	icated by lov	ver values)								
		very serious ¹	no serious inconsistency		very serious³	none	6	9	-	MD 0.27 lower (1.53 lower to 0.99 higher)	⊕OOO VERY LOW	IMPORTANT		

J.13.2 Population: Low back pain without sciatica

Table 302: MBR programme 3 elements: physical + psychological + education vs. Usual care/waiting list control

	Quality assessment	No of patients	Effect	Quality Importance
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¹ Downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Downgraded by 1 increment if the confidence interval crossed either the MID for benefit or the MID for harm

³ Downgraded by 2 increments if the confidence interval crossed both the MID for benefit and the MID for harm

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	MBR programme 3 elements: physical + psychological + education	Usual care/waiting list control	Relative (95% CI)	Absolute				
Pain seve	Pain severity, Aberdeen pain scale 0-100 (≤4 months) - Pain severity, Aberdeen pain scale 0-100 (≤4 months) (follow-up ≤4 months; range of scores: 0-100; Better indicated by low values)													
	randomised trials			no serious indirectness	serious²	none	85	94	1	MD 2.59 higher (0.37 to 4.81 higher)	⊕⊕OO LOW	CRITICAL		
Pain seve		en pain s	cale 0-100 (> 4 m	onths)- Pain se	-100 (> 4 months)(follow-up >	>4 months - 1 yea	ır; range	of scores: 0-100;	Better i	ndicated				
	randomised trials			no serious indirectness	serious ²	none	83	88	-	MD 4.44 higher (1.01 to 7.87 higher)	⊕⊕OO LOW	CRITICAL		
Function	, RMDQ 0-24	(≤4 mont	hs) - Function, R	MDQ (≤4 month	s) (follow-up	≤4 months; rang	je of scores: 0-24; Better ind	icated by lower va	alues)					
	randomised trials			no serious indirectness	serious²	none	85	94	1	MD 0.92 higher (0.02 lower to 1.86 higher)	⊕⊕OO LOW	CRITICAL		
Function	, RMDQ 0-24	(> 4 mon	ths)- Function, R	MDQ (> 4 mont	hs)(follow-up	>4 months - 1 y	ear; range of scores: 0-24; E	Setter indicated by	lower v	alues)				
	randomised trials			no serious indirectness	serious²	none	83	88	-	MD 1.42 higher (0.29 to 2.55 higher)	⊕⊕OO LOW	CRITICAL		

 $^{^{\}rm 1}$ Downgraded by 1 increment if the majority of the evidence was at high risk of bias $^{\rm 2}$ Downgraded by 1 increment if the confidence interval crossed one MID

Table 303: MBR programme 2 elements: physical + psychological vs. Usual care/waiting list control

			Quality ass	essment	,		No of patie	nts		Effect	Quality	Importance
No of	Design	Risk of	Inconsistency	Indirectness	Imprecision	Other	MBR programme 2	Usual	Relative	Absolute		

studies		bias				considerations	elements: physical + psychological	care/waiting list control	(95% CI)					
Psycholo	gical- BDI (≤₄	4 months) (follow-up ≤4 m	onths; range of	scores: 0-63	; Better indicated	by lower values)							
		very serious ¹	no serious inconsistency	no serious indirectness	serious²	none	27	25	-	MD 0.52 lower (7.37 lower to 6.33 higher)	⊕OOO VERY LOW	CRITICAL		
Psychological- STAI state (≤4 months) (follow-up ≤4 months; Better indicated by lower values)														
1		very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	27	25	-	MD 5.3 lower (9.32 to 1.28 lower)	⊕OOO VERY LOW	CRITICAL		
Psycholo	Psychological- STAI trait (≤4 months) (follow-up ≤4 months; Better indicated by lower values)													
		very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	27	25	-	MD 3.82 lower (9.88 lower to 2.24 higher)	⊕OOO VERY LOW	CRITICAL		
Pain seve	erity, VAS 0-1	0 (≤4 mo	nths) (follow-up :	≦4 months; rang	e of scores:	0-10; Better indic	ated by lower values)							
1		very serious ¹	no serious inconsistency	no serious indirectness	serious²	none	27	25	-	MD 1.41 lower (2.85 lower to 0.03 higher)	⊕OOO VERY LOW	CRITICAL		
Function	, RMDQ 0-24	(≤4 mont	hs) (follow-up ≤4	months; range	of scores: 0	-24; Better indicat	ed by lower values)							
		very serious ¹	no serious inconsistency	no serious indirectness	serious²	none	27	25	-	MD 2.85 lower (5.88 lower to 0.18 higher)	⊕OOO VERY LOW	CRITICAL		

¹ Downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Downgraded by 1 increment if the confidence interval crossed one MID

J.14 Return to work programmes

Individually delivered return to work programme (multidisciplinary) versus usual care in low back pain with or without sciatica J.14.1

Quality assessment	No of patients	Effect	Quality	Importance

	-	1	1		1	1								
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Individual multidisciplinary RTW programme	Usual care	Relative (95% CI)	Absolute				
Quality o	Quality of life (EQ-5D 0-1, change score) ≤ 4 months (range of scores: 0-1; Better indicated by lower values)													
1		no serious risk of bias	no serious inconsistency	no serious indirectness	no serious imprecision	none	94	92	-	MD 0.05 lower (0.13 lower to 0.03 higher)	⊕⊕⊕⊕ HIGH	CRITICAL		
Pain (NR	Pain (NRS 0-10, change score) ≤ 4 months (range of scores: 0-10; Better indicated by lower values)													
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	94	94	-	MD 0.21 higher (0.55 lower to 0.97 higher)	⊕⊕⊕O MODERATE	CRITICAL		
Pain (NR	Pain (NRS 0-10) >4 months (range of scores: 0-10; Better indicated by lower values)													
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^b	none	58	59	-	MD 0.21 lower (0.34 to 0.8 lower)	⊕⊕⊕O MODERATE	CRITICAL		
Pain (NR	S 0-10) >4 m	onths (rang	je of scores: 0-1(); Better indica	ted by lower va	alues)								
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	89	52	-	MD 1.16 lower (2.12 to 0.2 lower)	⊕⊕OO LOW	CRITICAL		
Function	(RMDQ 0-24	, change so	core) ≤ 4 months	(range of score	es: 0-24; Bette	r indicated by low	ver values)							
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	94	94	-	MD 0.91 higher (0.8 lower to 2.62 higher)	⊕⊕⊕O MODERATE	CRITICAL		
Function	(RMDQ 0-24	, change so	core) >4 months	(range of score	es: 0-24; Better	indicated by low	er values)							
1		no serious risk of bias	no serious inconsistency	no serious indirectness	very serious ^b	none	58	59	-	MD 2.73 higher (2.47 to 2.99 higher)	⊕⊕OO LOW	CRITICAL		
Psycholo	ogical distres	s (BDI, 0-6	3) > 4 months (ra	nge of scores:	0-63; Better in	dicated by lower	values)							
1	randomised	Serious ^a	no serious	no serious	no serious	none	89	52	-	MD 1.3 lower	⊕⊕⊕О	CRITICAL		

	T			1	I			1	I	272 fower nor		1			
										272 fewer per 1000 (from 111					
										fewer to 377 fewer)					
										iewei)					
Healthca	Healthcare utilisation (graded activity therapist, n of patients) > 4 months														
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	very serious ^b	none	55/66 (83.3%)	0%	RR 114.31 (7.21 to 1813.19)	-	⊕⊕OO LOW	IMPORTANT			
Healthca	Healthcare utilisation (manual therapist, n of patients) > 4 months														
4		· .					0/00	00.40/	DD 0.04	000 (IMPORTANT.			
	randomised trials	no serious risk of bias	inconsistency	no serious indirectness	no serious imprecision	none	6/66 (9.1%)	29.4%	RR 0.31 (0.13 to 0.72)	203 fewer per 1000 (from 82 fewer to 256 fewer)	⊕⊕⊕⊕ HIGH	IMPORTANT			
Healthca	Healthcare utilisation (cesar therapist, n of patients) > 4 months														
1	randomised	no serious	no corious	no serious	very serious ^b	none	3/66	7.4%	RR 0.62	28 fewer per 1000	⊕⊕00	IMPORTANT			
1	trials		inconsistency	indirectness	very serious	none	(4.5%)	7.470	(0.15 to 2.48)	(from 63 fewer to 110 more)	LOW	IIVIFORTANT			
Healthca	re utilisation	(physiothe	rapist, n of patie	ents) > 4 months	S										
1	randomised	no serious	no serious	no serious	very serious ^b	none	2/66	7.40%	RR 0.41	44 fewer per 1000	$\oplus\oplus$ OO	IMPORTANT			
	trials		inconsistency	indirectness	rely comedo		(3%)		(0.08 to 2.05)	(from 68 fewer to 78 more)	LOW				
Healthca	are utilisation	(psycholog	gist, n of patients	s) > 4 months											
1	randomised	no serious	no serious	no serious	very serious ^b	none	2/66	7.40%	RR 0.41	44 fewer per 1000	⊕⊕00	IMPORTANT			
·	trials		inconsistency	indirectness	very defidud	none	(3%)	7.4070	(0.08 to 2.05)	(from 68 fewer to 78 more)	LOW	IIVII OTTIVI			
Healthca	re utilisation	(alternative	therapist, n of	patients) > 4 mg	onths										
1	randomised	no serious	no sorious	no serious	very serious ^b	none	12/66	22 50/	DD 0 77 /0 4	54 fewer per 1000	⊕⊕00	IMPORTANT			
1	trials		inconsistency	indirectness	very serious	none	(18.2%)	23.5%		(from 141 fewer to 120 more)	LOW	IIVIFORTAINT			
Healthca	are utilisation	(medical s	pecialist, n of pa	tients) > 4 mon	ths										
1	randomised	no serious	no serious	no serious	Serious ^b	none	13/66	42.6%	RR 0.46	230 fewer per	⊕⊕⊕О	IMPORTANT			
L <u>'</u>	Landonniscu	ino ocinous	110 0011000	110 0011000	0011000	110110	10/00	72.070	11110.70	-oo lower per	\Box	IVII OKLANI			

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

NICE. 2016 Individually delivered return to work programme (multidisciplinary) versus usual care in low back pain without sciatica

			<u> </u>	0 (,,						
		Quality assessment No of patients Effect										
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Individual multidisciplinary RTW programme	Usual care	Relative (95% CI)	Absolute	Quality	Importance
Pain severity (NRS, 0-10 change score) ≤ 4 months (range of scores: 0-10; Better indicated by lower values)												
1	randomised trials	serious	no serious inconsistency	no serious indirectness	no serious imprecision	none	61	63	-	MD 0.30 lower (1.22 lower to 0.62 higher)	⊕⊕⊕O MODERATE	CRITICAL
Pain seve	erity (NRS, 0-	10 change s	score) > 4 months	(range of score	es: 0-10; Better	indicated by lowe	er values)		_			
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	60	59	-	MD 0.20 lower (1.3 lower to 0.9 higher)	⊕⊕⊕O MODERATE	CRITICAL
Function	(RMDQ, 0-24) ≤ 4 month	s (range of score	s: 0-24; Better i	ndicated by lov	ver values)			•			
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	62	64	-	MD 1.4 lower (3.66 lower to 0.86 higher)	⊕⊕OO LOW	CRITICAL
Function	(RMDQ, 0-24) > 4 months	s (range of score	s: 0-24; Better ii	ndicated by low	ver values)						
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	60	60	-	MD 0.6 lower (2.88 lower to 1.68 higher)	⊕⊕⊕O MODERATE	CRITICAL
Healthca	re utilisation	(consultatio	on with GP) > 4 m	onths (Better in	dicated by lowe	er values)						
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	67	67	-	MD 2.3 lower (4.22 to 0.38 lower)	⊕⊕OO LOW	IMPORTANT
Healthca	re utilisation	(Consultation	on with occupation	nal physician) >	>4 months (Bet	ter indicated by lo	ower values)					
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	Serious ^b	none	67	67	-	MD 0.9 lower (2.19 lower to 0.39	⊕⊕⊕O MODERATE	IMPORTANT

										higher)			
Healthca	ro utilisation	(CT scans/N	IRI scans) >4 mo	nths (Battar ind	icated by lowe	r values)							
1		no serious	no serious inconsistency	,	Serious ^b	none	67	67	-	MD 0.17 higher (0.05 lower to 0.39 higher)		IMPORTANT	
Healthca	re utilisation	(X-ray lumb	ar back) >4 mont	hs (Better indica	ated by lower v	values)							
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	67	67	-	MD 0.1 higher (0.43 lower to 0.63 higher)	⊕⊕⊕⊕ HIGH	IMPORTANT	
Healthca	Healthcare utilisation (Physio/paramedical therapy) >4 months (Better indicated by lower values)												
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	67	67	-	MD 7.5 higher (5.29 lower to 20.29 higher)	⊕⊕⊕⊕ HIGH	IMPORTANT	
Healthca	re utilisation	(Consultatio	ons to specialist)	>4 months (Bet	ter indicated b	y lower values)							
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	67	67	-	MD 0 higher (0.36 lower to 0.36 higher)	⊕⊕⊕⊕ HIGH	IMPORTANT	
Healthca	re utilisation	(Consultatio	ons to alternative	therapist) >4 m	onths (Better i	ndicated by lower	values)						
1		no serious	no serious inconsistency		no serious imprecision	none	67	67	-	MD 0.7 lower (2.38 lower to 0.98 higher)	⊕⊕⊕⊕ HIGH	IMPORTANT	
Healthca	re utilisation	(Pain medic	ation) >4 months	(Better indicate	ed by lower val	ues)							
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ^b	none	67	67	-	MD 0.4 lower (1.2 lower to 0.4 higher)	⊕⊕⊕O MODERATE	IMPORTANT	

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

NICE. 2016 Individually delivered return to work programme (unidisciplinary) versus usual care in low back pain without sciatica

Quality assessment						No of patients			Quality	Importance			
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	RTW individual unidisciplinary	Usual care	Relative (95% CI)	Absolute			
Quality of life (SF-36 Bodily Pain, 0-100) ≤ 4 months (range of scores: 0-100; Better indicated by higher values)													
	randomised trials	,			no serious imprecision	none	110	114	-	MD 6.2 higher (0.79 to 11.61 higher)	⊕⊕OO LOW	CRITICAL	
Quality of life (SF-36 Physical functioning, 0-100) ≤ 4 months (follow-up 3 months; range of scores: 0-100; Better indicated by higher values)													
	randomised trials	- ,			no serious imprecision	none	110	114	-	MD 5.6 higher (1.48 to 9.72 higher)	⊕⊕OO LOW	CRITICAL	
Pain (NRS	6 0-10, chang	e score) ≤	4 months (range	of scores: 0-10;	Better indicated	by lower values)							
	randomised trials	,			no serious imprecision	none	110	114	-	MD 0.7 lower (1.46 lower to 0.06 higher)	⊕⊕OO LOW	CRITICAL	
Function	(RMDQ 0-24,	change so	core) ≤ 4 months (range of scores:	0-24; Better inc	dicated by lower v	alues)						
	randomised trials	- ,			no serious imprecision	none	110	114	-	MD 1 lower (2.3 lower to 0.3 higher)	⊕⊕OO LOW	CRITICAL	
Sick leave	e ≤ 4 months												
	randomised trials			no serious indirectness	Serious ^b	none	17/150 (11.3%)	29/150 (19.3%)	RR 0.59 (0.34 to 1.02)	79 fewer per 1000 (from 128 fewer to 4 more)	⊕⊕OO LOW	CRITICAL	

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or downgraded by 2 increments if the confidence interval crossed both MIDs

NICE. 2016 Individually delivered return to work programme versus combination of interventions in low back pain without sciatica

			Quality asse	essment			No of pa	atients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Return to work programme (individual)	Combination of interventions	Relative (95% CI)		Quanty	Importance
Pain (NRS	6 0-10, final v	alue) ≤ 4 n	nonths (range of	scores: 0-10; Be	tter indicated	d by lower values)						
1	randomised trials	Serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	24	23	-	MD 0.72 lower (1.96 lower to 0.52 higher)	⊕⊕OO LOW	CRITICAL
Function	(RMDQ 0-24,	final value	e) ≤ 4 months (ran	ge of scores: 0-	24; Better in	dicated by lower v	/alues)					
1	randomised trials	Seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	24	23	-	MD 0.76 lower (3.65 lower to 2.13 higher)	⊕⊕OO LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID

J.14.5 Mixed group and individually delivered return to work programme versus usual care in low back pain with or without sciatica

			Quality ass	essment			No of patients			Effect	O. alifa	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Return to work programme (group and individual)	usual care	Absolute		Quanty	Importance
Return to	work >4 mor	nths										
		no serious risk of bias	no serious inconsistency		no serious imprecision	none	71/142 (50%)	47/81 (58%)	RR 0.86 (0.67 to 1.1)	81 fewer per 1000 (from 191 fewer to 58 more)	⊕⊕⊕⊕ HIGH	CRITICAL

Mixed group and individually delivered return to work programme (graded activity, cognitive behavioural approaches and education) versus return to work programme (graded activity and education) in low back pain without sciatica Ouglity assessment No of patients Fiffect

			Quality asso	essment			No of patient	s		Effect	Ovelito	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	RTW (group and individual, multidisciplinary)	RTW programme	Absolute		Quality	Importance
Return to	work >4 mo	nths (asse	essed with: Van d	len Hout)								
1	randomised trials			no serious indirectness	Serious ^b	none	35/41 (85.4%)	22/35 (62.9%)	RR 1.36 (1.02 to 1.8)	226 more per 1000 (from 13 more to 503 more)	⊕⊕OO LOW	CRITICAL

^a Downgraded by 1 increment if the majority of evidenc was at hight risk of bias

J.15 Spinal injections

J.15.1 Image-guided facet join injection

Table 304: Steroid versus saline for management of low back pain

			Quality as	sessment			No of patien	ts		Effect	0	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Image-guided FJI: Steroid	Saline	Relative (95% CI)	Absolute	Quality	Importance
Pain Seve	rity(VAS,0-10)	≤ 4 month	s (range of scores:	0-10; Better indic	ated by lower val	lues)						
	randomised trials				no serious imprecision ²	none	48	48	-	MD 0.2 lower (1.14 lower to 0.74 higher)	⊕⊕OO LOW	

^b Downgraded by 1 increment if the confidence interval crossed one MID

Pain Seve	rity(VAS,0-10)	>4 months	s - 1 year (follow-up	>4 months - 1 ye	ear; range of scor	es: 0-10; Better ind	licated by lower v	alues)					
1	randomised trials		no serious inconsistency	no serious indirectness	serious³	none	48	47	-	MD 1 lower (1.94 to 0.06 lower)	⊕OOO VERY LOW	CRITICAL	
Function(I	Function(MSIP) ≤ 4 month) (follow-up ≤4 months; range of scores: 0-100; Better indicated by lower values)												
1	randomised trials	- ,	no serious inconsistency		no serious imprecision²	none	48	48	-	MD 0.5 lower (2.72 lower to 1.72 higher)	⊕⊕OO LOW	CRITICAL	
Function(I	MSIP) >4 mont	h) (follow-	up >4 months - 1 y	ear; range of sco	es: 0-100; Better	indicated by lower	values)						
1	randomised trials	- ,	no serious inconsistency	no serious indirectness	serious²	none	48	47	-	MD 3 lower (6.16 lower to 0.16 higher)	⊕OOO VERY LOW	CRITICAL	

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

Table 305: Steroid versus hyaluronans for management of low back pain

			Quality as	sessment			No of pa	atients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Image-guided FJI: Steroid	Hyaluronans	Relative (95% CI)	Absolute		
Pain Seve	rity(VAS,0-10)) ≤ 4 mont	hs (follow-up ≤4 m	onths; range of s	scores: 0-10; Bet	ter indicated by lo	wer values)					
	randomised trials		no serious inconsistency	no serious indirectness	serious ²	none	29	30	-	MD 1.07 higher (0.18 lower to 2.32 higher)	⊕000 VERY LOW	CRITICAL
Pain Seve	rity(VAS,0-10)	>4 month	ns - 1 year (follow-	up >4 months - 1	year; range of s	cores: 0-10; Better	indicated by lo	wer values)				
	randomised trials			no serious indirectness	serious ²	none	29	30	-	MD 0.46 higher (0.73 lower to 1.65 higher)	⊕OOO VERY	CRITICAL

LOW

											LOW	
ınctio	n(ODI) ≤ 4 mon	th) (follow	-up ≤4 months; ra	nge of scores: 0-	100; Better indic	cated by lower valu	es)					
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision²	none	29	30	-	MD 0.95 higher (1.41 lower to 3.31 higher)	⊕⊕OO LOW	CRITICAL
unctio	n(RMQ) ≤ 4 mo	nth) (follow	w-up ≤4 months; r	ange of scores: (0-24; Better indic	ated by lower valu	es)		-			
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision ²	none	29	30	-	MD 1.20 higher (1.48 lower to 3.88 higher)	⊕⊕OO LOW	CRITICAL
unctio	n(LBOS)≤4 moi	nth (follow	y-up ≤4 months; ra	nge of scores: 0-	-75; Better indica	ated by lower value	s)					
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	29	30	-	MD 0.4 higher (30.53 lower to 31.33 higher)	⊕OOO VERY LOW	CRITICAL
unctio	n(ODI)>4 montl	h) (follow-	up >4 months - 1 y	/ear; range of sco	ores: 0-100; Bett	er indicated by low	er values)					
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision ²	none	29	30	-	MD 0.20 lower (2.37 lower to 1.97 higher)	⊕⊕OO LOW	CRITICAL
unctio	n(RMQ)>4 mon	th (range	of scores: 0-24; Be	etter indicated by	lower values)							
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision ²	none	29	30	-	MD 1.22 lower (3.83 lower to 1.39 higher)	⊕⊕OO LOW	CRITICAL
unctio	n(LBOS)>4 mor	nth (range	of scores: 0-10; B	Better indicated b	y lower values)							
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	29	30	-	MD 1.9 lower (32.39 lower to 28.59 higher)	⊕OOO VERY LOW	CRITICAL

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

Table 306: Steroid plus biomechanical exercise versus biomechanical exercise

		P.0.5		and the term	<u></u>	ilallical exercis						
			Quality ass	essment			No of patie	ents		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Image-guided FJI:steroid+excercise	Biomechanical Exercise	Relative (95% CI)	Absolute		
Pain sev	erity(VAS,0-1	0) ≤ 4 mo	nths (Better indi	cated by lower	values)						*	
1	trials		no serious inconsistency	no serious indirectness	Serious ²	none	36	34	-	MD 0.5 lower (1.38 lower to 0.38 higher)	⊕000 VERY LOW	CRITICAL
Function	n(MVAS,0-150) ≤ 4 mon	ths (Better indic	ated by lower v	alues)							
1	randomised trials		no serious inconsistency	no serious indirectness	Serious ²	none	36	34	-	MD 6.6 lower (17.58 lower to 4.38 higher)	⊕OOO VERY LOW	CRITICAL
Positive	Responders(Pain VAS	>50%) ≤4 month	s								
1		very serious ¹	no serious inconsistency	no serious indirectness	very serious	none	19/36 (52.8%)	17/34 (50%)	RR 1.06 (0.67 to 1.67)	30 more per 1000 (from 165 fewer to 335 more)	⊕000 VERY LOW	IMPORTANT
								50%		30 more per 1000 (from 165 fewer to 335 more)		
Positive	Responders(Disability	MVAS>50%) ≤4	months								
1		very serious ¹	no serious inconsistency	no serious indirectness	very serious	none	26/36 (72.2%)	23/34 (67.6%)	RR 1.07 (0.78 to 1.45)	47 more per 1000 (from 149 fewer to 304 more)	⊕000 VERY LOW	IMPORTANT
								67.7%		47 more per 1000 (from 149 fewer to 305 more)		

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ²Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MID

Table 307: Steroid plus anaesthetic versus biomechanical exercise for management of low back pain (cohort)

			Quality asses	sment			No of pati	ents		Effect	Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations		Back education and physiotherapy		Absolute	Quanty	importance	
QoL(EQ	QoL(EQ5D) (range of scores: 0-1; Better indicated by lower values)												
1		- , -			very serious¹	none	17	19	-	MD 0.02 lower (0.55 lower to 0.51 higher)	⊕OOO VERY LOW	CRITICAL	
Pain Se	verity(McGill) ≤ 4 r	nonths ((follow-up ≤4 m	nonths; range	of scores:	0-78; Better indi	cated by lower value	s)					
1		٠,			no serious imprecision ¹	none	19	17	-	MD 7.6 lower (16.22 lower to 1.02 higher)	⊕⊕OO LOW	CRITICAL	
Function	unction(ODI) ≤ 4 month (follow-up ≤4 months; range of scores: 0-80; Better indicated by lower values)												
1		- ,			no serious imprecision¹	none	17	19	-	MD 3.5 higher (5.23 lower to 12.23 higher)	⊕⊕OO LOW	CRITICAL	

¹ Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

J.15.2 Other Image-guided Injections

Table 308: Steroid versus saline for management of low back pain

Quality assessment	No of patients	Effect	Quality	Importance	į.
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² Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Other Image-guided Injections: Steroid	Saline	Relative (95% CI)	Absolute		
Pain Seve	erity(VAS,0-10) ≤4 month	ns (Better indicate	d by lower values	s)							
3	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	63	62	-	MD 4.19 lower (4.55 to 3.82 lower)	⊕⊕OO LOW	
Pain Seve	erity(VAS,0-10) ≤4 month	ns - Injection agen	t: Betamethason	e (Better indicat	ed by lower values)					
2	randomised trials	very serious ¹	serious ²	no serious indirectness	no serious imprecision	none	40	40	-	MD 5.2 lower (5.66 to 4.74 lower)	⊕000 VERY LOW	
Pain Seve	erity(VAS,0-10) ≤4 montl	ns - Injection agen	t: Dexamethason	e (Better indica	ted by lower values	5)					
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	23	22	-	MD 2.44 lower (3.04 to 1.84 lower)	⊕⊕OO LOW	
Pain Seve	erity(VAS,0-10) >4 montl	ns - 1 year (Better	indicated by low	er values)							
3	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	63	62	-	MD 3.38 lower (3.76 to 3.01 lower)	⊕⊕OO LOW	
Pain Seve	erity(VAS,0-10) >4 monti	ns - 1 year - Injecti	on agent: Betam	ethasone (Bette	r indicated by lowe	r values)					
2	randomised trials	very serious ¹	serious ¹	no serious indirectness	no serious imprecision	none	40	40	-	MD 4.76 lower (5.2 to 4.31 lower)	⊕000 VERY LOW	
Pain Seve	erity(VAS,0-10) >4 monti	ns - 1 year - Injecti	on agent: Dexam	ethasone (Bette	er indicated by lowe	er values)					
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	23	22	-	MD 0.28 lower (0.95 lower to 0.39 higher)	⊕⊕OO LOW	
Function(ODI), 0-100 ≤4	1 months (Better indicated b	y lower values)				-				

Low back pain and sciatica in over 16s Quality assessment

		1				1	1	1		T		
3	randomised	very	no serious	no serious	no serious	none	63	62	-	MD 21.4 lower (24.09		
	trials	serious1	inconsistency	indirectness	imprecision					to 18.71 lower)	LOW	
Function	(ODI), 0-100 ≤4	4 months	- Injection agent:	Betamethasone	(Better indicated	by lower values)		· ·		+	ļ	
	`		,		`	•						
2	randomised	very	serious ²	no serious	no serious	none	40	40	_	MD 27.95 lower (31.72	⊕000	
	trials	serious ¹		indirectness	imprecision					to 24.19 lower)	VERY	
		00000								10 2 0 .0 0	LOW	
											LOVV	
Function	(ODI) 0-100 </td <td>1 months</td> <td>- Injection agent:</td> <td>Dovamothasono</td> <td>(Retter indicate)</td> <td>d by lower values)</td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td>	1 months	- Injection agent:	Dovamothasono	(Retter indicate)	d by lower values)						
unction	(ODI), 0-100 <u>-</u>	+ 1110111115	- injection agent.	Dexamethasone	(Detter marcate)	u by lower values,						
1	randomised	very	no serious	no serious	no serious	none	23	22	_	MD 14.6 lower (18.44	⊕⊕00	
•	trials	serious ¹	inconsistency	indirectness	imprecision					to 10.76 lower)	LOW	
	triais	ocnodo	inconsistency	indirectiness	Impredictor					10 10.70 lower)	LOVV	
Eunction	(ODL 0 100) >4	months	⊥ · 1 year (Better ind	licated by lower	values)						<u> </u>	
runction	(ODI,0-100) >4	- 1110111115 -	· i year (Better ind	ilcated by lower	values							
4	randomised	very	no serious	no serious	serious ³	none	109	114	_	MD 12.02 lower (14.79	⊕000	
•	trials	serious ¹	inconsistency	indirectness	Conodo	110110	100	1		to 9.24 lower)	VERY	
	uiais	3011003	litoonsistency	indirectiness						(0 5.24 lower)	LOW	
											LOVV	
Function	(ODI 0-100) > <i>4</i>	months -	1 year - Injection	agent: Betameth	nasono (Bottor ir	ndicated by lower va	luos)					
unction	(001,0-100) - 4	- 1110111113 -	i year - mjection	agent. Detamet	iasone (Better ii	idicated by lower ve	iiucs)					
2	randomised	verv	serious ²	no serious	no serious	none	40	40	_	MD 24.06 lower (28.13	⊕000	
_	trials	serious ¹	00000	indirectness	imprecision			'		to 20 lower)	VERY	
	triais	Jenous		indirectiness	Impredictor					10 20 10WC1)	LOW	
											LOW	
Eunotion	(ODL0 100) >4	months	1 year Injection	agant: Mathynr	odnicolone cost	oto (Bottor indicator	hy lower values)	ļ				
unction	(ODI,0-100) >4	monuis -	1 year - injection	agent: Methypr	ednisolone acet	ate (Better indicated	i by lower values)					
1	randomised	very	no serious	no serious	no serious	none	46	52	_	MD 1.1 lower (7.11	⊕⊕00	
-	trials	serious ¹	inconsistency	indirectness	imprecision					lower to 4.91 higher)	LOW	
	1315	0011000			ipi colololi					ionor to 4.01 mgmcr)	LOW	
Function	(ODI.0-100) >4	months -	1 vear - Injection	agent: Dexamet	hasone (Better i	ndicated by lower v	alues)			<u> </u>		
23	(-2.,0 100) - 4		. ,	and a solution	(2010)							
1	randomised	very	no serious	no serious	no serious	none	23	22	-	MD 1.8 lower (6.7	⊕⊕00	
	trials	serious ¹	inconsistency	indirectness	imprecision					lower to 3.1 higher)	LOW	
		3003								12.10. to 0.1.1.g/101)	2011	
		1	1					1		1		

¹Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

NICE, 2016

Table 309: Steroid plus anaesthetic versus anaesthetic for management of low back pain

			Quality as	sessment			No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Other image-guided injections: Steroid+Anaesthetic		Relative (95% CI)	Absolute		
Pain Sev	erity(NRS,0-	10)≤ 4 mo	onths (follow-up	<4 months; Bet	ter indicated b	y lower values)						
	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	135	135	-	MD 0.19 lower (0.49 lower to 0.1 higher)	⊕⊕⊕O MODERATE	CRITICAL
Pain Sev	erity(NRS,0-	10) >4 mc	onths (follow-up	>4 months; Bet	ter indicated b	y lower values)						
	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	125	123	-	MD 0.24 lower (0.59 lower to 0.12 higher)	⊕⊕⊕O MODERATE	CRITICAL
Function	 n(ODI,0-100)	≤ 4 month	ls (follow-up <4 r	months; Better	indicated by lo	wer values)						
_	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	135	135	-	MD 0.41 lower (1.67 lower to 0.85 higher)	⊕⊕⊕O MODERATE	CRITICAL
Function	(ODI,0-100)	>4 month	ns (follow-up >4 r	months; Better	indicated by lo	wer values)						
	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	125	123	-	MD 0.00 higher (1.4 lower to 1.4 higher)	⊕⊕⊕O MODERATE	CRITICAL
Pain imp	rovement(>5	0%) ≤ 4 n	 nonths (follow-นุ	o <4 months)								

²Downgraded by 1 or 2 increments because of Heterogeneity, I2=50%, p=0.04, unexplained by subgroup analysis.

³ Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

2	randomised trials			no serious indirectness	no serious imprecision	none	63/75 (84%)	85%		43 fewer per 1000 (from 136 fewer to 77 more)		IMPORTANT
Pain imp	Pain improvement(>50%) >4 months (follow-up >4 months)											
2	randomised trials	serious ¹	serious ²	no serious indirectness	no serious imprecision	none	56/75 (74.7%)	75.8%		23 fewer per 1000 (from 144 fewer to 121 more)	⊕⊕OO LOW	IMPORTANT

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Downgraded by 1 or 2 increments because heterogeneity, I2=50%, p=0.04, unexplained by subgroup analysis.

Table 310: Steroid plus anaesthetic versus mixed modality exercise

			Quality as:	sessment			No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Image-guided FJI: Steroid+Anaesthetic	Back education and physiotherapy	Relative (95% CI)	Absolute	Quanty	importance
QoL(EQ	5D) (range of	scores: (0-1; Better indica	ted by lower va	ılues)							
1	randomised trials		no serious inconsistency	no serious indirectness	very serious ²	none	17	19	ı	MD 0.02 lower (0.55 lower to 0.51 higher)	⊕OOO VERY LOW	CRITICAL
Pain Sev	erity(McGill)	≤ 4 mont	hs (follow-up ≤4	months; range	of scores: 0-7	8; Better indicate	d by lower values)					
1		- ,	no serious inconsistency		no serious imprecision	none	19	17	-	MD 7.6 lower (16.22 lower to 1.02 higher)	⊕⊕OO LOW	CRITICAL
Function	action(ODI) ≤ 4 month (follow-up ≤4 months; range of scores: 0-80; Better indicated by lower values)											
1			no serious inconsistency		no serious imprecision	none	17	19	-	MD 3.5 higher (5.23 lower to 12.23 higher)	⊕⊕OO LOW	CRITICAL

¹ Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

. 2016 **Prolotherapy Injections**

Table 311: Sclerosant versus anaesthetic for management of low back pain

			Quality asse	essment			No of patients Effect			Effect	Quality	Importance
No of studies Design Risk of bias Inconsistency Indirectness Imprecision Other considerations Prolotherapy Injections: Sclerosant Anaesthetic (95% CI)												
Pain Seve	in Severity(VAS,0-10)≤ 4 months (follow-up ≤4 months; range of scores: 0-10; Better indicated by lower values)											
	randomised trials	- ,		no serious indirectness	very serious²	none	9	2	-	MD 0.10 lower (8.06 lower to 7.86 higher)	⊕OOO VERY LOW	CRITICAL

Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 312: Sclerosant plus anaesthetic versus saline for management of low back pain

		р.т			·	cite of fore but	- P					
			Quality as	sessment			No of patients Ef		Effect	.		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Prolotherapy Injections: Sclerosant+Anaesthetic	Saline	Relative (95% CI)	Absolute	Quality	Importance
Pain Severity(VAS,0-7.5)≤ 4 months (follow-up ≤4 months; range of scores: 0-7.5; Better indicated by lower values)												
1	randomised trials			no serious indirectness	serious²	none	40	41	-	MD 1.16 lower (1.81 to 0.51 lower)	⊕⊕OO LOW	CRITICAL

² Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

² Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

Pain Sev	erity(VAS,0-7	.5)>4 moı	nths - 1 year (follo	ow-up >4 month	s - 1 year; rang	e of scores: 0-7.5	; Better indicated by lower value	es)				
1	randomised trials		no serious inconsistency	no serious indirectness	serious²	none	40	41	-	MD 1.58 lower (2.26 to 0.9 lower)	⊕⊕OO LOW	CRITICAL
Function(RMQ)≤ 4 months (follow-up ≤4 months; range of scores: 0-33; Better indicated by lower values)												
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	40	41	-	MD 3.79 lower (6.28 to 1.3 lower)	⊕⊕⊕O MODERATE	CRITICAL
Function	(RMQ)>4 mor	nths - 1 ye	ear (follow-up >4	months - 1 year	; range of scor	es: 0-33; Better in	dicated by lower values)					
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision²	none	40	41	-	MD 4.86 lower (7.44 to 2.28 lower)	⊕⊕⊕O MODERATE	CRITICAL

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

Table 313: Sclerosant plus anaesthetic versus anaesthetic for management of low back pain

			Quality asse	essment			No of patients	Effect	Quality	Importance		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Prolotherapy Injections: Sclerosant+Anaesthetic	Anaesthetic	Relative (95% Absolute CI)		Quanty	importance
Pain Sev	erity(VAS,0-8	3)>4 month	ıs - 1 year (follov	v-up >4 months	- 1 year; rar	nge of scores: 0-8	s; Better indicated by lower va	lues)				
	trials	no serious risk of bias		no serious indirectness	serious ¹	none	39	40	-	MD 0.56 lower (1.34 lower to 0.22 higher)	⊕⊕⊕O MODERATE	CRITICAL
Function	(RMQ)>4 mo	nths - 1 ye	ar (follow-up >4	months - 1 yea	r; range of s	cores: 0-24; Bette	er indicated by lower values)					
1	trials	no serious risk of bias		no serious indirectness	serious¹	none	39	40	-	MD 0.34 lower (2.05 lower to 1.37 higher)	⊕⊕⊕O MODERATE	_

¹ Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

NICE, 2016 Other non-image guided injections

Table 314: Botox versus saline for management of low back pain

			Quality asses	ssment			No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Other Non-Image guided Injections: Botox	Saline	Relative (95% CI)	Absolute		
Responde	er Criteria(VA	S>50%) ≤4 ı	months (follow-u	o ≤4 months)				!!			-	
				no serious indirectness	serious ¹	none	9/15 (60%)	13.3%	RR 4.50 (1.16 to 17.44)	465 more per 1000 (from 21 more to 1000 more)	⊕⊕⊕O MODERATE	IMPORTAN [*]

Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 315: Steroid plus anaesthetic versus steroid for management of low back pain

			Quality asse	essment			No of patients Effect				0!!	
No of studies Design Risk of bias Inconsistency Indirectness Imprecision Other considerations Other Hon-Image guided Injections: Steroid+Anaesthetic Steroid Relative (95% CI)								Quality	Importance			
Pain Seve	erity(First Blo	ck NRS,0	-10) ≤4 month (fol	low-up ≤4 mont	hs; range of	scores: 0-10; Bet	ter indicated by lower values)					
	randomised very no serious inconsistency indirectness serious ² none 30 30 - MD 0.44 higher (0.72 lower to 1.6 higher) CRITICAL VERY LOW											
Pain Seve	rain Severity(Second Block NRS,0-10) ≤4 month (follow-up ≤4 months; range of scores: 0-10; Better indicated by lower values)											

1				no serious indirectness	serious²	none	30	30	-	MD 0.44 higher (0.77 lower to 1.66 higher)	⊕OOO VERY LOW	CRITICAL	
Pain Sev	Pain Severity(First Block VAS,0-10) ≤4 month (follow-up ≤4 months; range of scores: 0-10; Better indicated by lower values)												
1		- ,	no serious inconsistency	no serious indirectness	serious ²	none	30	30	-	MD 0.57 higher (0.61 lower to 1.75 higher)	⊕OOO VERY LOW	CRITICAL	
Pain Sev	erity(Second	Block VA	S,0-10) ≤4 month	(follow-up ≤4 m	onths; range	of scores: 0-10; E	Better indicated by lower values)						
1				no serious indirectness	serious ²	none	30	30	-	MD 0.25 higher (0.94 lower to 1.44 higher)	⊕OOO VERY LOW	CRITICAL	

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

Table 316: Botox versus steroid plus anaesthetic (injections into the paraspinous muscle) (cohort)

			Quality ass	essment			No of patients Effect Other Non-image-				Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Other Non-image- guided Injections: COHORT: Botox	Steroid+ Anaesthetic	Relative (95% CI) Absolute			
Respond	ler Criteria(Pair	(McGill)	improvement) >4	months - 1 yea	ar (follow-up 12	2 months)						
			no serious inconsistency		no serious imprecision	none	0/10 (0%)	77.8%	OR 0.04 (0.01 to 0.26)	655 fewer per 1000 (from 301 fewer to 744 fewer)	⊕OOO VERY LOW	IMPORTANT
Respond	ler Criteria(Pair	(McGill)	worsening) >4 m	onths - 1 year (follow-up 12 m	ionths)						
			no serious inconsistency		no serious imprecision	none	0/10 (0%)	77.8%	OR 0.04 (0.01 to 0.26)	655 fewer per 1000 (from 301 fewer to 744 fewer)	⊕000 VERY LOW	IMPORTANT

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Respond	der Criteria(Fur	oction (OI	OI) improved) >4	months - 1 yea	r (follow-up 12	months)						
1	observational	very	no serious	no serious	Serious ²	none	1/10	55.6%	RR 0.18	456 fewer per	⊕ООО	IMPORTANT
	studies	serious ¹	inconsistency	indirectness			(10%)		(0.03 to	1000 (from 539	VERY	
									1.26)	fewer to 145 more)	LOW	
Respond	der Criteria(Fur	ction (OI	OI) worsened) >4	months - 1 yea	ır (follow-up 12	months)						
1	observational	very	no serious	no serious	very serious ²	none	5/10	11.1%	RR 4.5	389 more per 1000	⊕ООО	IMPORTANT
	studies	serious ¹	inconsistency	indirectness			(50%)		(0.64 to	(from 40 fewer to	VERY	
									31.6)	1000 more)	LOW	

¹Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

²Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

J.16 Radiofrequency denervation

Table 317: Radiofrequency denervation versus placebo/sham for low back pain

			Quality ass	essment		·	No of	patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	RF denervation	placebo/sham	Relative (95% CI)	Absolute		
Pain (VA	S) 0-10 - <4 m	nonths (Bet	ter indicated by le	ower values)								
4	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	53	43	-	MD 1.46 lower (2.06 to 0.86 lower)	⊕⊕⊕O MODERATE	CRITICAL
Pain (VA	S) 0-10 - >4 m	onths (Bet	ter indicated by le	ower values)								
3		very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	80	80	-	MD 1.57 lower (2.2 to 0.95 lower)	⊕⊕OO LOW	CRITICAL
Pain (Mc	Gill) - <4 mon	ths (Better	indicated by low	er values)								
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	18	12	-	MD 7 lower (14.11 lower to 0.11 higher)	⊕⊕OO LOW	CRITICAL
Pain (Mc	Gill) - >4 mon	ths (Better	indicated by low	er values)								
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	very serious ²	none	18	12	-	MD 5 lower (20.43 lower to 10.43 higher)	⊕OOO VERY LOW	CRITICAL
Function	ODI 0-100 (c	hange and	final values) - <4	months (Better	indicated by lo	wer values)	<u> </u>					
3	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious²	none	35	31	-	MD 4.38 lower (7.31 to 1.45 lower)	⊕⊕OO LOW	CRITICAL

	randomised	very	no serious	no serious	serious ²	none	20	20	-	MD 5.6 lower (9.59	\oplus OOO	CRITICA
	trials	serious ¹	inconsistency	indirectness						to 1.61 lower)	VERY LOW	
unctio	n RMDQ 0-100) (change a	and final values) -	<4 months (Be	tter indicated b	y lower values)	<u> </u>					
	randomised	very	no serious	no serious	serious ²	none	36	34	T -	MD 2.6 higher (6.21	⊕OOO	CRITIC
	trials	serious ¹	inconsistency	indirectness						lower to 11.41 higher)	VERY LOW	
uality	of life (SF-36)	- General	health - <4 month	s (range of sco	res: 0-100; Bett	er indicated by	lower values)					
	randomised	serious ¹	no serious	no serious	no serious	none	40	41	T -	MD 3.1 higher (3.72	⊕⊕⊕О	CRITICA
	trials		inconsistency	indirectness	imprecision					lower to 9.92 higher)		
uality	of life (SF-36)	- Mental h	ealth - <4 months	(range of score	es: 0-100; Bette	r indicated by le	ower values)					
	randomised	serious ¹	no serious	no serious	serious ²	none	40	41	-	MD 2 higher (9.07	⊕⊕00	CRITIC
	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	40	41	-	MD 2 higher (9.07 lower to 13.07 higher)	⊕⊕OO LOW	CRITIC
uality	trials			indirectness				41	-	lower to 13.07		CRITIC
uality	trials of life (SF-36)		inconsistency	indirectness				41	-	lower to 13.07	LOW	
uality	trials of life (SF-36)	- Pain - <4	inconsistency months (range o	indirectness f scores: 0-100;	Better indicate	d by lower valu	ues)		-	lower to 13.07 higher)	±000	CRITICA
	trials of life (SF-36) randomised trials	- Pain - <4	months (range o	indirectness f scores: 0-100; no serious indirectness	Better indicate very serious ²	none	ues) 40	41	-	lower to 13.07 higher) MD 0.2 higher (9.29	±000	
	trials of life (SF-36) randomised trials of life (SF-36)	- Pain - <4	months (range o	indirectness f scores: 0-100; no serious indirectness	Better indicate very serious ²	none	ues) 40	41	-	lower to 13.07 higher) MD 0.2 higher (9.29	⊕000 VERY LOW	
	trials of life (SF-36) randomised trials of life (SF-36)	- Pain - <4 serious ¹ - Physical	months (range o	indirectness f scores: 0-100; no serious indirectness months (range of	very serious ²	none petter indicat	40 ted by lower value	41 es)	-	Iower to 13.07 higher) MD 0.2 higher (9.29 lower to 9.69 higher)	⊕000 VERY LOW	CRITIC
uality	randomised trials of life (SF-36) randomised trials randomised trials	- Pain - <4 serious¹ - Physical serious¹	months (range on the serious inconsistency) functioning - <4 in the serious inconsistency	indirectness f scores: 0-100; no serious indirectness months (range of the control of the con	very serious ² of scores: 0-100 serious ²	none ped by lower value none ped by lower value none	ted by lower value	41 es)	-	MD 0.2 higher (9.29 lower to 9.69 higher) MD 3.1 lower (11.09	⊕000 VERY LOW	CRITIC
uality	randomised trials of life (SF-36) randomised trials randomised trials	- Pain - <4 serious¹ - Physical serious¹ - Social fu	months (range on serious inconsistency) functioning - <4 in the serious inconsistency inconsistency	indirectness f scores: 0-100; no serious indirectness months (range of the control of the con	very serious ² of scores: 0-100 serious ²	none ped by lower value none ped by lower value none	ted by lower value	41 es)	-	MD 0.2 higher (9.29 lower to 9.69 higher) MD 3.1 lower (11.09	⊕OOO VERY LOW	CRITICA

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1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	40	41	-	MD 7.7 higher (0.64 to 14.76 higher)	⊕⊕OO LOW	CRITICAL
AEs: tre	eatment relate	d pain (mo	derate or severe)	- no. of patient	s - <4 months	•				•		•
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	23/39 (59%)	35.9%	RR 1.64 (1 to 2.69)	230 more per 1000 (from 0 more to 607 more)	⊕⊕OO LOW	IMPORTANT
AEs: ch	ange of sensi	bility (irrita	ting or evident d	ysaesthesia or a	allodynia) - no.	of patients - <	<4 months			L		
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	very serious	none	2/39 (5.1%)	0/40 0%	RR 5.12 (0.25 to 103.45)	-	⊕OOO VERY LOW	IMPORTANT
AEs: lo	ss of motor fu	nction (irri	tating or evident	motor loss) - no	o. of pts - <4 mo	onths						
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	very serious	none	0/38 (0%)	2.4%	RR 0.36 (0.02 to 8.55)	15 fewer per 1000 (from 24 fewer to 181 more)	⊕OOO VERY LOW	IMPORTANT
HC utili	sation: analge	sic use (no	o. of tablets/4 day	s) - <4 months	(Better indicate	d by lower va	lues)					
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	15	16	-	MD 3.24 lower (6.6 lower to 0.12 higher)	⊕OOO VERY LOW	IMPORTAN ⁻
HC utili	sation: analge	sic use (gl	obal perception of	of improvement	, 0-6) - >4 mont	hs (Better ind	icated by lower val	lues)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	40	40	-	MD 0.8 lower (1.56 to 0.04 lower)	⊕000 VERY LOW	IMPORTANT
Respon	der criteria (p	ercentage	of patients with >	•50% pain reduc	tion - global pe	rceived effec	t) - <4 months			l.		
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	-	-	OR 9.53 (1.05 to	-	⊕000 VERY LOW	IMPORTANT
								0%	86.28)	-		
Respon	der criteria (n	umber of p	atients with >50%	6 back pain or p	ain reduction -	global percei	ived effect) - <4 mo	onths	•			

Low back pain and sciatica in over 16s Quality assessment

2		no serious risk of bias	no serious inconsistency	no serious indirectness	serious ²	none	33/54 (61.1%)	39%	RR 1.74 (1.15 to 2.63)	289 more per 1000 (from 58 more to 636 more)	⊕⊕⊕O MODERATE	IMPORTANT
Respond	ler criteria (nu	umber of pa	tients with >50%	back pain or pa	ain reduction -	global perceived	effect) - >4 mo	onths (Copy)				
1	trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	7/15 (46.7%)	39%	RR 3.73 (0.92 to 15.21)	1000 more per 1000 (from 31 fewer to 1000 more)	⊕OOO VERY LOW	IMPORTANT
Respond	ier criteria (ni	umber of pa	tients with >50%	back pain redu	ction - VAS) - <	4 months						
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	very serious ²	none	13/40 (32.5%)	34.2%	RR 0.95 (0.51 to 1.76)	17 fewer per 1000 (from 168 fewer to 260 more)	⊕⊕OO LOW	IMPORTANT

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MID's.

Table 318: Radiofrequency denervation versus medial branch block for low back pain

			Quality asse	ssment			No of p	patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	RF denervation	medial branch block	Relative (95% CI)	Absolute	L uuy	
Pain (VNS) 0-10 - <4 mor	nths (Bette	er indicated by lowe	er values)	L							
1				no serious indirectness	serious ²	none	50	50	-	MD 1.2 lower (1.79 to 0.61 lower)	⊕000 VERY LOW	CRITICAL
Pain (VNS) 0-10 - >4 mor	nths (Bette	er indicated by lowe	er values)								
1		l ' .	no serious inconsistency	no serious indirectness	serious ²	none	50	50	-	MD 2.3 lower (3.42 to 1.18 lower)	⊕000 VERY	CRITICAL

											LOW	
Quality of	life (EQ-5D) 5	-15 scale -	<4 months (Bette	er indicated by lo	ower values)	<u> </u>	1		!			!
I	randomised trials	very serious ¹	no serious inconsistency	serious ³	serious ²	none	50	50	-	MD 0.4 lower (0.97 lower to 0.17 higher)	⊕000 VERY LOW	CRITICAL
Quality of	life (EQ-5D) 5	-15 scale -	>4 months (Bette	er indicated by l	ower values)							
I	randomised trials	very serious ¹	no serious inconsistency	serious ³	serious ²	none	50	50	-	MD 1.3 lower (2.87 lower to 0.27 higher)	⊕000 VERY LOW	CRITICAL

Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

J.17 Epidural injections for sciatica

Table 319: Image-guided Anaesthetic versus sham/placebo for Sciatica (>70% disc prolapse)

			Quality asses	sment			No of patients	3		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Anaesthetic versus sham/placebo	Control	Relative (95% CI)	Absolute		
Leg pain	(0-10, final va	lue) <4 mont	hs (Better indicat	ed by lower valu	es)							
1	randomised trials			no serious indirectness	serious ²	none	27	37	-	MD 1.2 higher (0.15 lower to 2.55 higher)	⊕⊕OO LOW	CRITICAL
Responde	er criteria: >5	0% reduction	n in pain <4 month	ıs								
1					very serious³	none	2/27 (7.4%)	18.9%	RR 0.39 (0.09 to 1.74)	115 fewer per 1000 (from 172 fewer to 140 more)		IMPORTANT

² Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MID's.

³ Downgraded by 1 or 2 increments because the majority of the evidence had indirect outcomes

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- 1 Downgraded by 1 increment if the majority of the evidence was at high risk of bias
- 2 Downgraded by 1 increment if the confidence interval crossed one MID
- 3 Downgraded by 2 increments if the confidence interval crossed both MIDs

Table 320: Image-guided Anti-TNF (mean of 3 doses) versus sham/placebo for Sciatica (>70% disc prolapse

			Quality as:	sessment			No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Anti-TNF (mean of 3 doses) versus sham/placebo	Control	Relative (95% CI)	Absolute	Quanty	Importance
Mean dail	ly worst leg p	ain (0-10,	change score) <4	months (Better	indicated by lo	wer values)						
1		very serious ¹	no serious inconsistency	no serious indirectness	very serious ²	none	27	10	-	MD 1.32 lower (3.3 lower to 0.66 higher)	⊕OOO VERY LOW	CRITICAL
AEs <4 m	onths											
1		- ,			no serious imprecision	none	0/18 (0%)	0%	not pooled	not pooled	⊕⊕OO LOW	IMPORTANT
AEs >4 m	onths											
1					no serious imprecision	none	0/18 (0%)	0/6 (0%)	not pooled	not pooled	⊕⊕OO LOW	IMPORTANT
			the majority of the					0%		not pooled		

¹ Downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 321: Image-guided Steroid + anaesthetic versus Sham/placebo for Sciatica (>70% disc prolapse)

	<u> </u>				7 1		·					
	Quality assessment						No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Steroid + anaesthetic versus Sham/placebo	Control	Relative (95% CI)	Absolute	•	·

² Downgraded by 2 increments if the confidence interval crossed both MIDs

Intensity	of leg pain -	Intensity of	leg pain <4 mon	ths (Better indic	cated by lower	values)						
1		no serious risk of bias	no serious inconsistency	no serious indirectness	serious ¹	none	28	37	-	MD 1.40 lower (2.79 to 0.01 lower)	⊕⊕⊕O MODERATE	CRITICAL
Oswestr	y disability in	dex - Oswe	stry disability inc	lex <4 months (Better indicate	d by lower values)					
1	randomised trials		no serious inconsistency	no serious indirectness	serious ¹	none	80	80	-	MD 1.3 lower (8.6 lower to 6 higher)	⊕⊕OO LOW	CRITICAL
Oswestr	y disability in	dex - Oswe	stry disability inc	lex >4 months (Better indicate	d by lower values)					
1	randomised trials		no serious inconsistency		no serious imprecision	none	80	80	-	MD 0.4 lower (7 lower to 6.2 higher)	⊕⊕⊕O MODERATE	CRITICAL
Respond	ler criteria: >5	50% reducti	on in pain <4 mo	nths		•						
1		no serious risk of bias	no serious inconsistency	no serious indirectness	no serious imprecision	none	15/28 (53.6%)	18.9%	RR 2.83 (1.34 to 6)	346 more per 1000 (from 64 more to 945 more)	⊕⊕⊕⊕ HIGH	IMPORTANT

Table 322: Image-guided Steroid+ anaesthetic versus anaesthetic for Sciatica (>70% prolanse)

	zz. muge į	Suided 3	Quality ass		unucstrictic	TOT SCIALICA (>)	No of patients			Effect	Quality	Importance
No of studies	Design Inconsistency Indirectness Imprecision					Other considerations	Steroid+ anaesthetic versus anaesthetic (>70% prolapse)	Control	Relative (95% CI)	Absolute		
Pain (0-1	0, change/fin	al scores) <	<4 months transf	oraminal epidu	ral (follow-up <	4 months; Better	indicated by lower valu	ies)				
	randomised trials				no serious imprecision ²	none	116	117	-	MD 0.52 lower (1.04 lower to 0 higher)	⊕⊕⊕O MODERATE	CRITICAL

¹ Downgraded by 1 increment if the confidence interval crossed one MID ² Downgraded by 1 increment if the majority of the evidence was at high risk of bias

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Pain (0-	10, change/fir	al scores)	<4 months caud	al epidural (foll	ow-up <4 mon	ths; Better indic	cated by lower values)					
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	176	177	-	MD 0.70 lower (1.33 to 0.07 lower)	⊕⊕OO LOW	CRITICAL
Pain (0-	10, change/fin	nal scores)	>4 months - trar	sforminal appr	oach (Better in	idicated by lowe	er values)					
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	no serious imprecision	none	60	60	-	MD 0.2 higher (0.37 lower to 0.77 higher)	⊕⊕⊕⊕ HIGH	CRITICAL
Pain (0-	10, change/fin	al scores)	>4 months - cau	dal epidural (Bo	etter indicated	by lower values	5)					
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	60	60	-	MD 0.6 lower (1.24 lower to 0.04 higher)	⊕⊕OO LOW	CRITICAL
ODI sco	ore (0-100, cha	nge/final so	core) <4 months	(Better indicate	ed by lower va	lues)						
2	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	120	120	-	MD 2.46 lower (4.16 to 0.75 lower)	⊕⊕OO LOW	CRITICAL
ODI sco	ore (0-100, fina	ıl score) >4	months (Better	indicated by lo	wer values)							
2	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	120	120	-	MD 1.4 lower (3.16 lower to 0.36 higher)	⊕⊕⊕O MODERATE	CRITICAL
Respon	der criteria: >	50% reduct	ion in pain <4 m	onths - transfo	 raminal approa	ach						
3	randomised trials	serious ¹	serious	no serious indirectness	serious ²	none	80/116 (69%)	76.7%	RR 1.29 (1.06 to 1.57)	222 more per 1000 (from 46 more to 437 more)	⊕000 VERY LOW	IMPORTANT
Respon	der criteria: >	50% reduct	ion in pain <4 m	onths - caudal	epidural							
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	48/60 (80%)	76.7%	RR 1.04 (0.86 to	31 more per 1000 (from 107 fewer to	⊕⊕OO LOW	IMPORTANT

	1											
									1.26)	199 more)		
spond	ler criteria: >	50% reduct	ion in pain <4 m	onths - interlam	inar (parisagg	jital approach)						
	randomised	very	no serious	no serious	serious ²	none	30/35	65%	RR 1.71	462 more per 1000	⊕OOO	IMPORTA
	trials	serious ¹	inconsistency	indirectness			(85.7%)		(1.19 to	(from 124 more to		
			,				,		2.46)	949 more)		
espond	ler criteria: >	 50% reduct	 ion in pain >4 m	onths - transfor	 aminal approa	ach						
	randomised	no serious	no corious	no serious	serious ²	none	43/88	65%	RR 0.84	92 fewer per 1000	###O	IMPORTA
	trials		inconsistency	indirectness	Serious	none	(48.9%)	05%	(0.64 to	(from 208 more to	0000	
	liiais	lisk of blas	linconsistency	lituliectiless			(40.970)		1.1)	58 more)	MODERATE	
									1.1)	56 more)		
espond	ler criteria: >	50% reduct	ion in pain >4 m	onths - caudal	epidural							
	randomised	serious ¹	no serious	no serious	serious ²	none	41/60	65%	RR 1.08	52 more per 1000	⊕⊕00	IMPORTAI
	trials		inconsistency	indirectness	Scrious	none	(68.3%)	0370	(0.83 to	(from 111 fewer to	LOW	IIVII OITIA
	uiais		linconsistency	indirectiness			(00.370)		1.4)	260 more)	LOW	
									1.4)	200 more)		
espond	ler criteria: >	50% reduct	ion in pain >4 m	onths - interlam	inal (parisagg	ital) approach		•				!
	randomised	very	no serious	no serious	serious ²	none	31/35	65%	RR 1.51	331 more per 1000	⊕ООО	IMPORTA
	trials	serious1	inconsistency	indirectness			(88.6%)		(1.11 to	(from 72 more to	VERY LOW	
			,						2.04)	676 more)		
espond	ler criteria: >	50% reduct	ion in ODI <4 mo	onths - transfor	aminal approa	ch						
	randomised	serious ¹	no serious	no serious	serious ²	none	41/60	75%	RR 0.91	67 fewer per 1000	⊕⊕OO	IMPORTAI
	trials		inconsistency	indirectness			(68.3%)		(0.73 to	(from 202 fewer to	LOW	
									1.14)	105 more)		
espond	ler criteria: >	50% reduct	ion in ODI <4 mo	onths - caudal e	pidural							
					<u> </u>							
	randomised	no serious	no serious	no serious	serious ²	none	44/60	61.7%	RR 1.19	117 more per 1000	$\oplus \oplus \oplus O$	
	trials	risk of bias	inconsistency	indirectness			(73.3%)		(0.93 to	(from 43 fewer to	MODERATE	
			1						1.53)	327 more)		1
									1.00)	027 111010)		

Low back pain and sciatica in over 16s Quality assessment

Respond	er criteria: >	50% reduc	tion in ODI >4 mo	nths								
	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	81/120 (67.5%)	65.8%	RR 1.03 (0.86 to 1.23)	20 more per 1000 (from 92 fewer to 151 more)	⊕⊕⊕O MODERATE	IMPORTANT
HC use: S	Surgery >4 m	onths										
	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	8/28 (28.6%)	66.7%	RR 0.43 (0.23 to 0.82)	380 fewer per 1000 (from 120 fewer to 514 fewer)	⊕⊕OO LOW	IMPORTANT
HC use: o	opioid intake	, mg dose	in last 12 months	<4 months (Be	etter indicated I	by lower values)						
	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	120	120	-	MD 4.73 lower (13.53 lower to 4.08 higher)	⊕⊕⊕O MODERATE	IMPORTANT
HC use: o	opioid intake	, mg dose	in last 12 months	>4 months (Be	etter indicated I	by lower values)	l					
	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	120	120	-	MD 3.98 lower (12.8 lower to 4.84 higher)	⊕⊕⊕O MODERATE	IMPORTANT
HC use: r	number of pa	l ntients hav	ing additional inj	ections>4 mont	ths (follow-up >	•4 months)						
		very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	20/35 (57.1%)	66.7%	RR 0.84 (0.58 to 1.22)	107 fewer per 1000 (from 280 fewer to 147 more)		IMPORTANT
			l	1				l .			l .	l

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, or 2 increments if at very high risk of bias ² Downgraded by 1 increment if the confidence interval crossed one MID

Table 323: Image-guided Steroid+ anaesthetic versus anaesthetic for Sciatica (non disc lesion)

· ·	•			
Quality assessment	No of patients	Effect	Quality	Importance

No of

Risk of

Other

Steroid+

Relative

	randomised trials	serious ²	no serious inconsistency	no serious indirectness	very serious ⁴	none	31/50 (62%)	66%	RR 0.94 (0.7 to 1.26)	40 fewer per 1000 (from 198 fewer to	⊕000 VERY LOW	IMPORTAN
			,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,				(3=73)			172 more)	VERT EOT	
espo	nder criteria: >	50% redu	ction in pain >4 r	nonths								
	randomised	serious ²	no serious	no serious	very serious ⁴	none	22/50	42%	RR 1.05	21 more per 1000		IMPORTAN
	trials		inconsistency	indirectness			(44%)		(0.67 to 1.65)	(from 139 fewer to 273 more)	VERY LOW	
spo	nder criteria: >3	30% reduc	ction in RMDQ <	4 months								
	randomised	very	no serious	no serious	serious ³	none	61/193	37.3%	RR 0.85	56 fewer per 1000		IMPORTAN
	trials	serious ¹	inconsistency	indirectness			(31.6%)		(0.64 to 1.12)	(from 134 fewer to 45 more)	VERY LOW	
espo	nder criteria: >	50% reduc	ction in ODI <4 m	nonths								
	randomised	serious ²	no serious	no serious	serious ³	none	25/50	58%	RR 0.86 (0.6	81 fewer per 1000		IMPORTAN'
	trials		inconsistency	indirectness			(50%)		to 1.24)	(from 232 fewer to 139 more)	LOW	
espo	nder criteria: >	50% reduc	ction in ODI >4 m	nonths								
	randomised	serious ²	no serious	no serious	very serious ⁴	none	23/50	42%	RR 1.1 (0.7	42 more per 1000		IMPORTAN
	trials		inconsistency	indirectness			(46%)		to 1.71)	(from 126 fewer to 298 more)	VERY LOW	
C us	e: opioid intake	, mg dose	in last 12 montl	ns <4 months (B	 Setter indicated b	y lower value	s)					
	randomised	serious ²	no serious	no serious	no serious	none	50	50	-	MD 0.2 lower (12.69	⊕⊕⊕О	IMPORTAN'
	trials		inconsistency	indirectness	imprecision					lower to 12.29 higher)	MODERATE	

1	randomised trials		no serious inconsistency	no serious indirectness	serious ³	none	50	50	-	MD 3.2 lower (18.6 lower to 12.2 higher)	⊕⊕OO LOW	IMPORTANT
SAEs <4	months	I							l			
2			no serious inconsistency	no serious indirectness	very serious ⁴	none	4/250 (1.6%)	1.3%	RR 0.8 (0.22 to 2.94)	3 fewer per 1000 (from 10 fewer to 25 more)	⊕000 VERY LOW	IMPORTANT
SAEs >4	months	•										
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision ⁵	none	0/50 (0%)	0%	not pooled	not pooled	⊕⊕⊕O MODERATE	IMPORTANT

¹ Downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 324: Image-guided Steroid+ anaesthetic versus anaesthetic for Sciatica (mixed population / unclear spinal pathology)

			Quality ass	essment			No of pa	atients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Steroid+ anaesthetic	Anaesthetic	Relative (95% CI)	Absolute		
Pain <4 n	nonths-transf	oraminal ep	pidural (follow-up	<4 months; Bet	tter indicated by	y lower values)		,		,		
2		very serious ¹	serious ²	no serious indirectness	serious ³	none	168	164		MD 0.06 lower (0.34 lower to 0.22 higher)	⊕000 VERY LOW	CRITICAL
Pain <4 n	nonths-appro	ach not spe	ecified (follow-up	<4 months; Bet	ter indicated by	/ lower values)						
1		very serious¹	serious ²	no serious indirectness	no serious imprecision	none	168	164		MD 0.07 lower (1.11 lower to 1.25 higher)	⊕OOO VERY LOW	CRITICAL
Pain, PPI	(0-5, change	score) <4 m	nonths (Better inc	licated by lower	r values)							

² Downgraded by 1 increment if the majority of the evidence was at high risk of bias

³ Downgraded by 1 increment if the confidence interval crossed one MID

⁴ Downgraded by 2 increments if the confidence interval crossed both MIDs

⁵ Zero events in both arms

	•		•		1	,					•	
1	randomised trials	very serious¹	no serious inconsistency	no serious indirectness	serious ³	none	34	35	-	MD 0.04 higher (0.35 lower to 0.43 higher)	⊕OOO VERY LOW	CRITICAL
ODI scor	e (0-100, chai	nge/final sc	ore) <4 months (E	Better indicated	by lower value	s)						
2	randomised trials	very serious ¹	very serious ²	no serious indirectness	no serious imprecision ⁴	none	134	129	-	MD 0.01 higher (2.83 lower to 2.85 higher)	⊕000 VERY LOW	CRITICAL
HC use:	Surgery <4 m	onths										
2	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	very serious ⁴	none	9/62 (14.5%)	18.3%	RR 0.79 (0.36 to 1.74)	38 fewer per 1000 (from 117 fewer to 135 more)	⊕000 VERY LOW	IMPORTANT
HC use:	Surgery >4 m	onths										
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	very serious ⁴	none	9/64 (14.1%)	21.5%	RR 0.65 (0.3 to 1.4)	75 fewer per 1000 (from 150 fewer to 86 more)	⊕000 VERY LOW	IMPORTANT
HC use:	medication re	eduction (>2	0% opioid use or	cessation non-	-opioids) <4 mo	nths						
1	randomised trials		no serious inconsistency	no serious indirectness	serious ³	none	17/28 (60.7%)	46.7%	RR 1.3 (0.8 to 2.11)	140 more per 1000 (from 93 fewer to 518 more)	⊕⊕⊕O MODERATE	IMPORTANT
HC use:	medication re	eduction (>2	0% opioid use or	cessation non-	-opioids) >4 mo	nths						
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	serious ³	none	11/12 (91.7%)	75%	RR 1.22 (0.85 to 1.77)	165 more per 1000 (from 112 fewer to 577 more)	⊕⊕⊕O MODERATE	IMPORTANT
AEs: con	nplications >4	4 months										
1	randomised trials	very serious¹	no serious inconsistency	no serious indirectness	no serious imprecision ⁵	none	0/64 (0%)	0/65 (0%)	not pooled	not pooled	⊕⊕OO LOW	IMPORTANT
								0%		not pooled		
AEs: con	nplications <	4 months										
1	randomised	very	no serious	no serious	no serious	none	0/65	0/59	not pooled	not pooled	⊕⊕00	IMPORTANT

trials	serious ¹	inconsistency	indirectness	imprecision ⁵	(0%)	(0%)		LOW	
						0%	not pooled		

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Table 325: Image guided: Steroid + anaesthetic epidural versus combination of non-invasive interventions for Sciatica (>7	70% prolai	pse)
rabic 323. illiage guided. Steroid - allaestrictic epidaral versus combination of non-illvasive interventions for sciatica (*/	7070 prota	pscj

			Quality ass	essment			No of patients			Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Steroid + anesthetic versus combination of non invasive interventions	Control	Relative (95% CI)	Absolute	- Quality	Importance
HRQoL (f	follow-up >4	months; B	l Setter indicated b	y lower values)								
	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	50	50	-	MD 2.24 lower (2.76 to 1.72 lower)	⊕⊕⊕O MODERATE	CRITICAL
Pain (foll	ow-up > 4; B	etter indica	ated by lower val	lues)								
	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	50	50	-	MD 3.39 lower (3.65 to 3.13 lower)	⊕⊕⊕O MODERATE	CRITICAL
Function	(follow-up >	4 months;	; Better indicated	by lower value	es)							
	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	50	50	-	MD 12.59 lower (13.42 to 11.76 lower)	⊕⊕⊕O MODERATE	CRITICAL
Psycholo	gical distres	รร (follow-น	up >4 months; Be	etter indicated b	y lower value	s)						
1	randomised	serious ¹	no serious	no serious	no serious	none	50	50	-	MD 4.67 lower	⊕⊕⊕О	CRITICAL

	trials		inconsistency	indirectness	imprecision					(5.44 to 3.9 lower)	MODERATE	
Respond	der criteria (c	omplete rel	lief of pain) >4 n	nonths (follow-	up >4 months)							
1	randomised	no serious	no serious	no serious	no serious	none	43/52	12/50	RR 3.45	588 more per	$\oplus \oplus \oplus \oplus$	IMPORTANT
	trials	risk of bias	inconsistency	indirectness	imprecision		(82.7%)	(24%)	(2.07 to 5.73)	1000 (from 257 more to 1000 more)	HIGH	
								0%		-		

Low back pain and sciatica in over 16s Quality assessment

Table 326: Image-guided Anti-TNF + anaesthetic versus anaesthetic for Sciatica (>70% disc prolapse)

			Quality asse	ssment			No of patients	S		Effect	Quality	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Anti-TNF + anaesthetic versus anaesthetic	Control	Relative (95% CI)	Absolute	Quality	Importance
Pain (0-10), change/fina	al scores) <	4 months (Better	indicated by lov	wer values)							
	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	very serious¹	none	26	30	-	MD 0.22 lower (1.76 lower to 1.32 higher)	⊕⊕OO LOW	CRITICAL
ODI score	e (0-100, final	score) <4 r	nonths (Better in	dicated by lowe	r values)					·		
	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	serious ²	none	26	30	-	MD 10.26 higher (0.69 to 19.83 higher)	⊕⊕⊕O MODERATE	CRITICAL
HC use: S	Surgery <4 m	onths										
	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	very serious ¹	none	6/26 (23.1%)	16.7%	RR 1.38 (0.48 to 4.01)	63 more per 1000 (from 87 fewer to 503 more)		IMPORTANT
Responde	er criteria: >5	60% reduction	on in pain <4 mor	nths								

a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

1	I .	no serious risk of bias	no serious inconsistency	no serious indirectness	very serious ¹	none	11/26 (42.3%)	43.3%	RR 0.98 (0.53 to 1.79)	9 fewer per 1000 (from 204 fewer to 342 more)	⊕⊕OO LOW	IMPORTANT
Respond	er criteria: >5	60% reduction	on in pain >4 mo	nths								
1		no serious risk of bias	no serious inconsistency	no serious indirectness	very serious ¹	none	10/26 (38.5%)	40%	RR 0.96 (0.5 to 1.85)	16 fewer per 1000 (from 200 fewer to 340 more)	⊕⊕OO LOW	CRITICAL
HC use:	medication re	duction (>2	20% opioid use o	cessation non-	-opioids) <4 ı	months						
1		no serious risk of bias	no serious inconsistency	no serious indirectness	very serious ¹	none	9/26 (34.6%)	46.7%	RR 0.74 (0.39 to 1.42)	121 fewer per 1000 (from 285 fewer to 196 more)	⊕⊕OO LOW	IMPORTANT
HC use:	medication re	duction (>2	20% opioid use o	cessation non-	-opioids) >4 ı	months						
1	I .	no serious risk of bias	no serious inconsistency	no serious indirectness	very serious ¹	none	7/11 (63.6%)	75%	RR 0.85 (0.49 to 1.48)	112 fewer per 1000 (from 382 fewer to 360 more)	⊕⊕OO LOW	IMPORTANT

¹ Downgraded by 2 increments if the confidence interval crossed both MIDs ² Downgraded by 1 increment if the confidence interval crossed one MI

Table 327: Image-guided Steroid + anaesthetic versus Anti-TNF + anaesthetic for Sciatica (>70% disc prolapse)

			Quality asse				No of patients		<u> </u>	Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Steroid + anaesthetic versus Anti-TNF + anaesthetic	Control	Relative (95% CI)	Absolute	Quality	Importance
Pain (0-1	0) <4 months	(Better ind	icated by lower v	alues)								
1		no serious risk of bias		no serious indirectness	serious ¹	none	28	26		MD 1.02 lower (2.63 lower to 0.59 higher)		CRITICAL
ODI scor	e (0-100, final	score) <4 ı	months (Better in	dicated by lowe	er values)							

						1						1
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	serious ¹	none	28	26	-	MD 16.16 lower (26.15 to 6.17 lower)	⊕⊕⊕O MODERATE	CRITICAL
Respond	ler criteria: >5	60% reducti	on in pain <4 mo	nths								
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	very serious ²	none	14/28 (50%)	42.3%	RR 1.18 (0.66 to 2.11)	76 more per 1000 (from 144 fewer to 470 more)	⊕⊕OO LOW	IMPORTANT
Respond	ler criteria: >5	50% reducti	on in pain >4 mo	nths								
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	very serious ²	none	8/28 (28.6%)	38.5%	RR 0.74 (0.35 to 1.59)	100 fewer per 1000 (from 250 fewer to 227 more)	⊕⊕OO LOW	IMPORTANT
HC use:	Surgery <4 m	onths										
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	very serious ²	none	6/28 (21.4%)	23.1%	RR 0.93 (0.34 to 2.52)	16 fewer per 1000 (from 152 fewer to 351 more)	⊕⊕OO LOW	IMPORTANT
HC use:	medication re	eduction (>2	20% opioid use o	r cessation non	ı-opioids) <4	months						
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	serious ¹	none	17/28 (60.7%)	34.6%	RR 1.75 (0.96 to 3.22)	259 more per 1000 (from 14 fewer to 768 more)	⊕⊕⊕O MODERATE	IMPORTANT
HC use:	medication re	eduction (>2	20% opioid use o	r cessation non	ı-opioids) >4	months						
1	trials		no serious inconsistency	no serious indirectness		none	11/12 (91.7%)	63.6%	RR 1.44 (0.89 to 2.32)	280 more per 1000 (from 70 fewer to 840 more)	⊕⊕⊕O MODERATE	IMPORTANT

¹ Downgraded by 1 increment if the confidence interval crossed one MID

Table 328: Non image guided: Steroid epidural versus placebo/sham for Sciatica

Quality assessment	No of patients	Effect	Quality	Importance

² Downgraded by 2 increments if the confidence interval crossed both MIDs

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Steroid versus placebo/sham	Control	Relative (95% CI)	Absolute		
Function	(follow-up 3-	12 months;	measured with: O	DI/RMDQ; Bette	r indicated by I	ower values)						
2	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	112	109	ı	SMD 0.1 lower (0.37 lower to 0.16 higher)	⊕⊕OO LOW	CRITICAL
Pain (VAS	S) (follow-up	3-4 months;	measured with: \	/AS; Better indi	cated by lower	values)						
2	randomised trials	serious ^a			no serious imprecision	none	65	109	-	MD 0.41 lower (1.39 lower to 0.56 higher)	⊕⊕⊕O MODERATE	CRITICAL
Pain McG	ill: present p	ain intensity	(follow-up 3 mor	nths; measured	with: McGill sca	ale; Better indicate	ed by lower values)					
1		no serious risk of bias			no serious imprecision	none	77	79	-	MD 0 higher (0.49 lower to 0.49 higher)	⊕⊕⊕⊕ HIGH	CRITICAL
Pain (McC	Gill score: pa	in raiting inc	dex) (follow-up 3 r	months; measur	ed with: McGill	score ; Better ind	icated by lower val	ues)				
1		no serious risk of bias			no serious imprecision	none	77	79	-	MD 0 higher (5.93 lower to 5.93 higher)	⊕⊕⊕⊕ HIGH	CRITICAL
adverse e	events- morbi	dity (follow-	up 2-27 weeks; as	ssessed with: no	o of minor even	its)						
2	randomised trials	serious ^a		no serious indirectness	serious ^c	none	25/113 (22.1%)	19/119 (16%)	RR 1.36 (0.81 to 2.3)	48 more per 1000 (from 25 fewer to 172 more)	⊕⊕OO LOW	CRITICAL
						af biss and day			:f.th			

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 329: Non image guided: Steroid epidural versus usual care for Sciatica

Quality assessment	No of patients	Effect	Quality	Importance
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^b Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

^cDowngraded by 1 increment I²>50%, and point estimates vary widely.

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Steroid versus usual care	•	Relative (95% CI)	Absolute		
Pain score	>4months - N	IRS back _I	pain (follow-up 52 v	veeks; measured	with: VAS; Bette	r indicated by lowe	r values)					
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	33 3	30	-	MD 0.7 lower (1.92 lower to 0.52 higher)	⊕⊕OO LOW	CRITICAL
Quality of	life (SF-36) 0-	100 ≤4 moi	nths - Mental comp	osite (Better indic	cated by lower va	ilues)						
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	25 2	25	-	MD 3.8 higher (2.65 lower to 10.25 higher)	⊕⊕OO LOW	CRITICAL
Quality of	life (SF-36) 0-	100 ≤4 moi	nths - Physical con	nposite (Better inc	licated by lower	values)						
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	25 2	25	-	MD 9.5 higher (2.32 to 16.68 higher)	⊕⊕OO LOW	CRITICAL
Quality of	life (SF-36) 0-	100 ≤4 moi	nths - Physical fun	ctioning (Better in	dicated by lower	values)						
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	25 2	25	-	MD 8.7 higher (1.03 to 16.37 higher)	⊕⊕OO LOW	CRITICAL
Quality of	life (SF-36) 0-	100 ≤4 moi	nths - Physical role	limitations (Bette	er indicated by lo	wer values)		•				
1	randomised	serious ¹	no serious	no serious	serious ²	none	25 2	25	-	MD 14 higher (5.68 lower	⊕⊕ОО	CRITICAL

	trials		inconsistency	indirectness						to 33.68 higher)	LOW	
Quality of	life (SF-36) 0-	100 ≤4 mo	nths - Social funct	ioning (Better ind	icated by lower v	alues)		11-				<u>'</u>
	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	25	25	-	MD 4.4 higher (3.32 lower to 12.12 higher)	⊕⊕OO LOW	CRITICAI
uality of	ilfe (SF-36) 0-	100 ≤4 mo	nths - Emotional r	ole limitations (Be	etter indicated by	lower values)		<u> </u>				
	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	25	25	-	MD 13.5 higher (2.69 lower to 29.69 higher)	⊕⊕OO LOW	CRITICAL
Quality of	life (SF-36) 0-	100 ≤4 mo	nths - Emotional w	vell-being (Better	indicated by lowe	r values)						
	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	25	25	-	MD 1.2 lower (9.33 lower to 6.93 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality of	life (SF-36) 0-	100 ≤4 mo	nths - Energy/fatig	ue (Better indicat	ed by lower value	es)		<u> </u>				
	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	25	25	-	MD 2.4 lower (11.24 lower to 6.44 higher)	⊕⊕OO LOW	CRITICAL
Quality of	life (SF-36) 0-	100 ≤4 mo	nths - Pain (Better	indicated by lowe	er values)							
	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	25	25	-	MD 3.1 higher (2.14 lower to 8.34 higher)	⊕⊕⊕O MODERATE	CRITICAI
uality of	life (SF-36) 0-	100 ≤4 mo	nths - General hea	Ith perceptions (E	Better indicated b	y lower values)						

1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	25	25	-	MD 6.8 higher (0.72 lower to 14.32 higher)	⊕⊕OO LOW	CRITICAL
Quality of	life (SF-36) 0-	100 ≤4 mo	nths - Change in pe	erceived help (Bet	ter indicated by l	lower values)						
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	very serious ²	none	25	25	-	MD 2.6 higher (10.99 lower to 16.19 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	life (SF-36) 0-	100 >4 mo	nths - Mental comp	osite (Better indic	cated by lower va	alues)						
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	25	25	-	MD 1.8 higher (4.92 lower to 8.52 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality of	life (SF-36) 0-	100 >4 mo	nths- Physical com	posite (Better ind	icated by lower \	/alues)						
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	25	25	-	MD 11.9 higher (4.64 to 19.16 higher)	⊕⊕OO LOW	CRITICAL
Quality of	life (SF-36) 0-	100 >4 mo	nths - Physical fun	ctioning (Better in	dicated by lower	rvalues)						
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	25	25	-	MD 7.5 higher (0.36 lower to 15.36 higher)	⊕⊕OO LOW	CRITICAL
Quality of	life (SF-36) 0-	100 >4 mo	nths - Physical role	limitations (Bette	er indicated by lo	wer values)						
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	25	25	-	MD 29.1 higher (8.55 to 49.65 higher)	⊕⊕OO LOW	CRITICAL

Quality of	ilfe (SF-36) 0-	100 >4 mo	nths – 1 year - Soc	ial functioning (B	etter indicated by	/ lower values)						
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	25	25	-	MD 4.6 higher (3.26 lower to 12.46 higher)	⊕⊕OO LOW	CRITICAL
Quality of	life (SF-36) 0-	100 >4 mo	nths – 1 year - Em	otional role limita	tions (Better indi	cated by lower valu	ues)	-				•
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	25	25	-	MD 9.1 higher (7.57 lower to 25.77 higher)	⊕⊕OO LOW	CRITICAL
Quality of	life (SF-36) 0-	100 >4 mo	nths – 1 year - Emo	otional well-being	(Better indicated	by lower values)		-				•
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	25	25	-	MD 4.8 lower (13.13 lower to 3.53 higher)	⊕⊕OO LOW	CRITICAL
Quality of	life (SF-36) 0-	100 >4 mo	nths – 1 year - Ene	rgy/fatigue (Bette	r indicated by lov	ver values)						
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	25	25	-	MD 1.4 lower (10.2 lower to 7.4 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality of	life (SF-36) 0-	100 >4 mo	nths – 1 year - Pair	n (Better indicated	l by lower values)						
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	25	25	-	MD 1.5 lower (6.81 lower to 3.81 higher)	⊕⊕⊕O MODERATE	CRITICAL
Quality of	life (SF-36) 0-	100 >4 mo	nths – 1 year - Gen	eral health perce	otions (Better inc	licated by lower va	lues)					
1	randomised	serious ¹	no serious	no serious	serious ²	none	25	25	-	MD 4.7 higher (3.16 lower	⊕⊕ОО	CRITICAL

	trials		inconsistency	indirectness						to 12.56 higher)	LOW	
Quality of	life (SF-36) 0-	100 >4 mo	nths – 1 year - Cha	nge in perceived	help (Better indic	ated by lower value	es)	<u> </u>		-		1
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	25	25	-	MD 14.5 higher (0.53 to 28.47 higher)	⊕⊕OO LOW	CRITICAL
Pain score	e ≤4 months - ∣	NRS back	pain (follow-up me	an 13 weeks; Bet	ter indicated by l	ower values)						•
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	33	30	-	MD 0.9 lower (2.27 lower to 0.47 higher)	⊕⊕OO LOW	CRITICAL
Pain score	e ≤4 months - I	NRS total	pain (follow-up 13	weeks; Better ind	icated by lower v	alues)						
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	33	30		MD 0.7 lower (2.02 lower to 0.62 higher)	⊕⊕OO LOW	CRITICAL
Pain score	e ≤4 months - l	NRS pain	during night (follow	v-up 13 weeks; Be	etter indicated by	lower values)		•				,
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	33	30	-	MD 0.9 lower (2.27 lower to 0.47 higher)	⊕⊕OO LOW	CRITICAL
Pain score	e ≤4 months - l	NRS pain	during day (follow-	up 13 weeks; Bet	ter indicated by le	ower values)						
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	33	30	-	MD 0.7 lower (2.09 lower to 0.69 higher)	⊕⊕OO LOW	CRITICAL
Pain score	e ≤4 months - l	NRS leg pa	ain (follow-up 13 w	eeks; Better indic	ated by lower val	lues)						

1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	33	30	-	MD 1.1 lower (2.42 lower to 0.22 higher)	⊕⊕OO LOW	CRITICAL		
Pain scor	e >4 months –	1 year - Ni	RS leg pain (follow	up 52 weeks; Bet	ter indicated by	ower values)								
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	33	30	-	MD 0.4 lower (1.44 lower to 0.64 higher)	⊕⊕OO LOW	CRITICAL		
Pain scor	e >4 months –	1 year - Ni	RS pain during day	(follow-up 52 wee	eks; Better indica	ated by lower value	s)							
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	33	30	-	MD 1 lower (2.27 lower to 0.27 higher)	⊕⊕OO LOW	CRITICAL		
Pain scor	Pain score >4 months – 1 year - NRS pain during night (follow-up 52 weeks; Better indicated by lower values)													
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	33	30	-	MD 1 lower (2.19 lower to 0.19 higher)	⊕⊕OO LOW	CRITICAL		
Pain scor	e >4 months –	1 year - Ni	RS total pain (follow	w-up 52 weeks; Bo	etter indicated by	/ lower values)								
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	33	30	-	MD 0.8 lower (2.07 lower to 0.47 higher)	⊕⊕OO LOW	CRITICAL		
Function	score ≤ 4 mont	ths (follow	-up mean 13 weeks	s; measured with:	ODI; Better indic	cated by lower valu	es)							
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	33	30	-	MD 2.3 lower (5.32 lower to 0.72 higher)	⊕⊕OO LOW	CRITICAL		

Function s	score >4 mont	hs – 1 yea	r (follow-up mean 5	2 weeks; measure	ed with: ODI; Bet	ter indicated by lov	ver values)					
1	randomised trials			no serious indirectness	serious ²	none	33	30	-	MD 1.8 lower (4.35 lower to 0.75 higher)	⊕⊕OO LOW	CRITICAL

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

Table 330: Non image guided: Steroid + anaesthetic epidural versus placebo for Sciatica

			Quality as	sessment			No of patient	s		Effect	Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Steroid + anaesthetic versus placebo	Control	Relative (95% CI)	Absolute	Quality	importance	
Function	score - Disab	oility (ODI))≤4 months (follow	w-up mean 12 w	eeks; measure	d with: ODI; Bette	r indicated by lower	values)					
	randomised trials				no serious imprecision	none	120	108	-	MD 0 higher (5.22 lower to 5.22 higher)	⊕⊕⊕O MODERATE	CRITICAL	
Function	nction score - (ODI) >4 months – 1 year (follow-up mean 52 weeks; measured with: ODI; Better indicated by lower values)												
1	randomised trials			no serious indirectness	serious ^b	none	120	108	-	MD 2 lower (8.12 lower to 4.12 higher)	⊕⊕OO LOW	CRITICAL	
Pain ≤4 m	nonths - VAS	leg pain (follow-up mean 1	2 weeks; measu	red with: VAS;	Better indicated I	by lower values)						
1	randomised trials			no serious indirectness	no serious imprecision	none	120	108	-	MD 0.5 lower (1.36 lower to 0.36 higher)	⊕⊕⊕O MODERATE	CRITICAL	
Pain ≤4 m	nonths - VAS	back pair	ı (follow-up mean	12 weeks; mea	sured with: VA	S; Better indicated	d by lower values)						
1	randomised trials				no serious imprecision	none	120	108	-	MD 0.3 lower (1.08 lower to 0.48 higher)	⊕⊕⊕O MODERATE	CRITICAL	
Pain> 4 m	nonths – 1 ye	ar - VAS I	eg pain (follow-u	mean 52 week	s; measured wi	th: VAS; Better in	dicated by lower val	ues)					

						_						
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	120	108	-	MD 0.3 lower (1.21 lower to 0.61 higher)	⊕⊕⊕O MODERATE	CRITICAL
Pain> 4 r	nonths – 1 ye	ar - VAS I	back pain (follow	-up mean 52 we	eks; measured	with: VAS; Better	indicated by lower v	values)				
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	120	108	-	MD 0.1 lower (0.93 lower to 0.73 higher)	⊕⊕⊕O MODERATE	CRITICAL
Psycholo	gical distres	s ≤ 4mont	ths - HAD anxiety	(follow-up mea	n 12 weeks; me	easured with: HAD	; Better indicated by	lower v	alues)			
1	randomised trials		no serious inconsistency	no serious indirectness	serious ^b	none	120	108	-	MD 1 higher (0.04 lower to 2.04 higher)	⊕⊕OO LOW	IMPORTANT
Psycholo	gical distress	s ≤ 4mont	hs - HAD depres	sion (follow-up ı	mean 12 weeks	; measured with: I	HAD; Better indicate	d by low	ver values)			
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	120	108	-	MD 0 higher (1.04 lower to 1.04 higher)	0000	IMPORTANT
Psycholo	gical distress	s >4 mont	ths – 1 year - HAI	O depression (fo	llow-up mean 5	52 weeks; measure	ed with: HAD; Better	indicate	ed by lower v	alues)		
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	106	108	-	MD 0 higher (1.21 lower to 1.21 higher)		IMPORTANT
Psycholo	gical distress	s >4 mont	ths – 1 year - HAI	D anxiety (follow	v-up mean 52 w	eeks; measured w	ith: HAD; Better ind	icated b	y lower value	es)		
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	106	97	-	MD 0 higher (1.38 lower to 1.38 higher)		IMPORTANT
Healthca	re utilisation	(further p	hysiotherapy) (fo	ollow-up mean 5	2 weeks; asses	sed with: No. unde	ertaking further phy	siothera	phy)			
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	37/120 (30.8%)	27/108 (25%)	RR 1.34 (0.75 to 2.4)	59 more per 1000 (from 50 fewer to 194 more)	⊕⊕OO LOW	IMPORTANT
Healthca	re utilisation	(referal to	pain manageme	nt services) (fol	low-up mean 52	2 weeks; assessed	l with: No. refered to	pain m	anagement)			

1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	0/120 (0%)	2/108 (1.9%)	RR 0.12 (0.01 to 1.94)	17 fewer per 1000 (from 19 fewer to 17 more)	⊕⊕OO LOW	IMPORTANT
Healthca	re utilisation	(further e	pidurals) (follow-	up mean 52 wee	ks; assessed w	rith: No. referred f	or further epidurals)					
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	19/120 (15.8%)	13/108 (12%)	RR 1.37 (0.64 to 2.94)	37 more per 1000 (from 40 fewer to 166 more)	⊕⊕OO LOW	IMPORTANT
Healthca	re utilisation	(analgesi	cs) - ≤4 months (f	ollow-up mean	12 weeks; meas	sured with: Mean a	analgesic use/week;	Better i	ndicated by	ower values)		
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	120	108	-	MD 7 lower (16.26 lower to 2.26 higher)	0000	IMPORTANT
Healthca	re utilisation	(analgesi	cs) - >4 months (f	ollow-up mean	52 weeks; meas	sured with: Mean	analgesic use/week;	Better i	ndicated by	lower values)		
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	120	108	ı	MD 2 lower (12.35 lower to 8.35 higher)	0000	IMPORTANT
Healthca	re utilisation	(surgery)	(follow-up mean	52 weeks; asses	ssed with: 75%	improvement on I	pack pain likert)					
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	18/120 (15%)	15/108 (13.9%)	RR 1.09 (0.52 to 2.29)	11 more per 1000 (from 62 fewer to 131 more)	⊕⊕⊕O MODERATE	IMPORTANT

Responder criteria - Improvement on leg pain (follow-up mean 52 weeks; assessed with: 75% improvement on leg pain likert)

1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	67/120 (55.8%)	51/108 (47.2%)	RR 1.41 (0.84 to 2.38)	86 more per 1000 (from 43 fewer to 208 more)	⊕⊕OO LOW	IMPORTANT
Respor	ıder criteria - Ir	nproveme	ent on back pain (follow-up mean	52 weeks; asse	essed with: 75% ir	nprovement on back	pain lik	ert)			
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	58/120 (48.3%)	47/108 (43.5%)	RR 1.21 (0.72 to 2.05)	47 more per 1000 (from 78 fewer to 177 more)	⊕⊕OO LOW	IMPORTANT
Advers	e events- morb	idity (follo	ow-up mean 52 w	eeks; assessed	with: minor adv	verse events)						
1	randomised trials	serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	11/120 (9.2%)	11/108 (10.2%)	RR 0.9 (0.41 to 1.99)	10 fewer per 1000 (from 60 fewer to 101 more)	⊕⊕OO LOW	IMPORTANT

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

Table 331: Non image guided :Steroid + Anaesthetic epidural versus combination of non-invasive interventions for Sciatica

			Quality as:	sessment			No of patients			Effect	Quality	
No of studies							Steroid + Anaesthetic versus usual care	Control	Relative (95% CI)	Absolute	Quality	Importance
Pain (VAS) (follow-up mean 2 weeks; measured with: VAS; Better indicated by lower values)												
1	randomised trials				no serious imprecision	none	120	19		MD 0.97 lower (11.95 lower to 10.01 higher)	⊕⊕⊕О	CRITICAL

					MODE	RATE	

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 332: Non image guided: Steroid + anaesthetic epidural versus pharmacological treatment (NSAIDS) for Sciatica

			Quality ass	essment			No of patients			Effect	Ouality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Steroid + anaesthetic versus pharmacological treatment (NSAIDS)	Control	Relative (95% CI)	Absolute	quanty	importance
Function	≤4 months (f	ollow-up	mean 3 months;	measured with:	ODI; Better	indicated by lowe	er values)					
	randomised trials			no serious indirectness	serious ^b	none	34	30	-	MD 4.1 lower (8.9 lower to 0.7 higher)		CRITICAL
Pain ≤4 n	nonths (follow	v-up mea	n 3 months; mea	sured with: VAS	S; Better indi	cated by lower va	alues)					
	randomised trials		no serious inconsistency	no serious indirectness	serious ^b	none	34	30	-	MD 0.8 lower (1.49 to 0.11 lower)	⊕⊕OO LOW	CRITICAL
Healthca	re utilisation	(analgesi	cs) (follow-up me	ean 3 months; a	ssessed wit	h: No. using para	cetamol)					
	randomised trials		no serious inconsistency	no serious indirectness	serious ^b	none	5/34 (14.7%)	8/30 (26.7%)	RR 0.47 (0.14 to 1.65)	121 fewer per 1000 (from 218 fewer to 108 more)		IMPORTANT

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

Table 333: Non image guided: Steroid + anaesthetic epidural versus pharmacological treatment (combination) for Sciatica

Quality assessment	No of patients	Effect	Quality	Importance
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No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Steroid + anaethetic versus pharmacological treatment (combination)	Control	Relative (95% CI)	Absolute		
Pain - ≤ 4	months (foll	ow-up me	ean 3 months; me	easured with: V	AS ; Better ii	ndicated by lower	values)					
1		,		no serious indirectness	serious ^b	none	25	25	-	MD 0.5 lower (1.23 lower to 0.23 higher)	⊕OOO VERY LOW	CRITICAL
Pain -> 4	months – 1 y	ear (follo	w-up mean 6 mo	nths; measured	with: VAS;	Better indicated b	y lower values)					
1	randomised trials			no serious indirectness	serious ^b	none	25	25	-	MD 0.5 lower (1.26 lower to 0.26 higher)	⊕⊕OO LOW	CRITICAL
Adverse	events - morl	bidity (fol	low-up mean 6 m	onths; assesse	d with: No. r	ninor adverse eve	ents)					
1	randomised trials			no serious indirectness	serious ^b	none	5/25 (20%)	4/25 (16%)	RR 1.25 (0.38 to 4.12)	40 more per 1000 (from 99 fewer to 499 more)	⊕⊕OO LOW	IMPORTANT

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

Table 334: Non image guided: Steroid + anaesthetic epidural versus anaesthetic epidural for Sciatica caused by (>70%) disc prolapse

			Quality ass	sessment			No of patients			Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Steroid + anaesthetic versus anaesthetic for sciatica caused by (>70%) disc prolapse	Control	Relative (95% CI)	Absolute	Quality	Importance
Pain ≤ 4 r	months - Met	thyl predr	nisolone versus l	bupivacaine (fo	l ollow-up 3 mon	ths; measured w	ith: VAS; Better indicated by	y lower v	alues)			

CRITICAL

 $\oplus \oplus \oplus O$

randomised serious1

no serious

no serious

no serious

none

50

55

MD 1.28 lower

									0.78)	fewer)		
Healthca	re utilisation	- physiot	herapy - Dexame	thasone + Bup	ivicaine versus	s anaesthetic (fol	low-up 3 months)	1				
1	randomised trials			no serious indirectness	serious ^b	none	12/40 (30%)	19/42 (45.2%)	`	154 fewer per 1000 (from 285 fewer to 81 more)	LOW	IMPORTANT

Table 335: Non image guided: Steroidand anesthetic epidual versus anaesthetic for sciatica caused by (>70%) spinal stenosis

			Quality ass	essment			No of patients		Effect			Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Steroid + anaesthetic versus anaesthetic for sciatica caused by (>70%) spinal stenosis	Control	Relative (95% CI)	Absolute	Quality	Importance
Respond	ler criteria <4	months:	spinal stenosis (follow-up 1 day	/s)							
		- ,	no serious inconsistency	no serious indirectness	serious ^b	none	10/18 (55.6%)	6/12 (50%)	RR 1.11 (0.55 to 2.24)	55 more per 1000 (from 225 fewer to 620 more)	⊕OOO VERY LOW	IMPORTANT
Respond	ler criteria >4	months:	spinal stenosis (follow-up 20.8	months)	l				<u> </u>		
		- ,		no serious indirectness	serious ^b	none	7/18 (38.9%)	4/12 (33.3%)	RR 1.17 (0.43 to 3.13)	57 more per 1000 (from 190 fewer to 710 more)	⊕OOO VERY LOW	IMPORTANT
HC use-	surgery: spin	al stenos	sis (follow-up 20.	8 months)	<u> </u>			,				
		very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	none	8/18 (44.4%)	7/12 (58.3%)	RR 0.76 (0.38 to	140 fewer per 1000 (from 362 fewer to	⊕OOO VERY	IMPORTANT

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

_								
					1.54)	315 more)	LOW	

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

Table 336: Non image guided: Steroid + epidural versus anaesthetic epidural for Sciatica in a population with unclear spinal pathology

			Quality ass	essment			No of patients			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Steroid + anaesthetic versus anaesthetic for sciatica in a population with unclear spinal pathology	Control	Relative (95% CI)	Absolute	quanty	importance
Reduced	analgesic in	take (follo	ow-up 1 months)									
		,	no serious inconsistency	no serious indirectness	serious ^b	none	8/15 (53.3%)	6/14 (42.9%)	`	104 more per 1000 (from 221 fewer to 403 more)	⊕OOO VERY LOW	IMPORTANT
healthca	re use - surg	ery (follo	w-up 1 months)									
		,	no serious inconsistency	no serious indirectness	serious ^b	none	4/15 (26.7%)	4/15 (26.7%)	RR 1 (0.31 to 3.28)	0 fewer per 1000 (from 184 fewer to 608 more)		IMPORTANT

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

Table 337: Non image guided: Steroid + epidural versus anaesthetic epidural for Sciatica in a population with unclear spinal pathology

			Quality asse	essment			No of patients			Effect	Quality	Importance
No of	Design	Risk of	Inconsistency	Indirectness	Imprecision	Other	Anaesthetic versus steroid with unclear	Control	Relative	Absolute		

studies		bias				considerations	spinal pathology	(95% CI)			
healthcar	e use (surge	ry) (follow	/-up 1 months)								
		,		no serious indirectness	serious ^b	none	0/19 (0%)		110 fewer per 1000 (from 124 fewer to 77 more)		IMPORTANT
										2011	

^a Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

Referral for surgery J.18

J.18.1 Low back pain

Table 338: Smoking for Referral for surgery (low back pain and/or Sciatica) - surgery: open decompressive laminectomy

Quality asse	essment						Adjusted effects	Quality
Number of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other consideration s, including publication bias where possible	Pooled effect with 95% CIs	
_	rsus non-smokin nd/or Sciatica]	g for predicting t	he treatment effect(1	TE=change in ODI(su	ırgery) – Change in	ODI(non-operati	ve) (Adjusted MDs) [adults	with low
1	Prospective cohort	very serious ^a	no serious inconsistency	no serious indirectness	No serious imprecision	None	Adjusted Mean Difference[Standard Error]: 10.1 (3.055)a	LOW

^a Downgraded by 1 increment if the majority of the evidence had serious limitations

Table 339: BMI>30 for Referral for surgery (patients with back or leg pain)-surgery not defined

Quality assess	ent	Adjusted effects	Quality

Quality asse	ssment						Adjusted effects	Quality
Number of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other consideration s, including publication bias where possible	Pooled effect with 95% CIs [if meta-analysed]	
BMI>30 vers	us BMI< 25 for բ	oredicting the eff	ect on Function (RDQ	(≤4) at 3 months(Ac	ljusted ORs) [adults	aged 18-65 with	back or leg pain]	
	Prospective cohort	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	None	Adjusted OR: 0.79 [0.21, 2.94]	VERY LOW

^a Downgraded by 1 increment if the majority of the evidence had serious limitations ^b95% Cl around the median crosses null line.

Table 340: Psychological Distress for Referral for surgery (patients with back or leg pain)-surgery not defined

Quality asse	ssment						Adjusted effects	Quality
Number of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other consideration s, including publication bias where possible	Pooled effect with 95% Cls [if meta-analysed]	
Psychologic	al Distress (Nega	tive Affectivity (N	NEM>1-≤4 versus NEM	⁄I ≤1) on Back Pain	(VAS≤10mm) (Adju	sted ORs) [adults	aged 18-65 with back or le	g pain]
1	Prospective cohort	serious ^a	no serious inconsistency	no serious indirectness	No serious imprecision	None	Adjusted OR: 0.55 [0.19, 1.61]	MODERATE
Psychologic	al Distress (Nega	tive Affectivity (N	NEM>4 versus NEM ≤	1) on Back Pain (VA	AS≤10mm) (Adjuste	d ORs) [adults ag	ed 18-65 with back or leg pa	ain]
1	Prospective cohort	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	None	Adjusted OR: 0.21 [0.06, 0.78]	VERY LOW

 $^{^{\}it a}$ Downgraded by 1 increment if the majority of the evidence had serious limitations $^{\it b}$ 95% CI around the median crosses null line.

J.18.2 Sciatica CE. 2016 Table 34

Table 341: Risk factor for Radicular Symptoms (continuous outcome) for Referral for surgery (low back pain and/or Sciatica population)-surgery: open decompressive laminectomy

Quality asse	ssment						Adjusted effects	Quality
Number of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other consideration s, including publication bias where possible	Pooled effect with 95% CIs [if meta-analysed]	
		n versus pre-op p in and/or Sciatica		in predicting the tre	eatment effect(TE=c	change in ODI sur	gery – Change in ODI non-o	perative)
1	Cohort studies	very serious ^a	no serious inconsistency	no serious indirectness	No serious imprecision	None	Adjusted Mean Difference[Standard Error]: -4.2 (1.088)	LOW

^a Downgraded by 1 increment if the majority of the evidence had serious limitations

Table 342: Risk factor for Radicular symptoms for Referral for surgery (patients with back or leg pain)-surgery not defined

Quality asse	essment						Adjusted effects	Quality
Number of studies	Imber of Study Risk of bias Inconsistency Indirectness Imprecision Other Pooled consideration CIs [if something publication bias where possible e-operative Leg Pain(VAS >43) versus Leg Pain (VAS ≤43)on Leg Pain(VAS≤10 mm) at 3 months (Adjusted ORs) Adjusted ORs) [again]							
Pre-operation leg pain]	ve Leg Pain(VAS	>43) versus Leg	Pain (VAS ≤43)on Leg	Pain(VAS≤10 mm)	at 3 months (Adjust	ted ORs) Adjusted	d ORs) [adults aged 18-65 w	ith back or
1	Cohort studies	very serious ^a	no serious inconsistency	no serious indirectness	No serious imprecision	None	Adjusted OR: 0.24 [0.10, 0.58]	LOW
Pre-operation leg pain]	ve Leg Pain(VAS	>43) versus Leg	Pain (VAS ≤43)on Leg	Pain(VAS≤10 mm)	at 12 months (Adju	sted ORs) Adjusto	ed ORs) [adults aged 18-65 v	vith back or

Quality ass	sessment		Adjusted effects	Quality				
1	Cohort studies	very serious ^a	no serious inconsistency	no serious indirectness	No serious imprecision	None	Adjusted OR: 0.38 [0.16, 0.75]	LOW

^a Downgraded by 1 increment if the majority of the evidence had serious limitations

Table 343: Risk factor for Radicular Symptoms (Categorical outcome) for Referral for surgery (Sciatica population)-surgery: dissection of the paravertebral muscles down to the laminae and resection of the interlaminar

Quality asse	ssment		Adjusted effects	Quality				
Number of studies	Study design	Risk of bias	Inconsistency Indirectne on (ODI>10) at 1 year (Adjusted ORs)		Imprecision	Other consideration s, including publication bias where possible	Pooled effect with 95% CIs [if meta-analysed]	
Effects of Pr	e-op Leg Pain(V/	AS) on Function (ODI>10) at 1 year (Ad	ljusted ORs) [adults	aged 15-83 with page	atients with Sciat	ica]	
1	Cohort studies	very serious ^a	no serious inconsistency	no serious indirectness	serious ^b	None	Adjusted OR: 0.523 [0.135, 2.028]	VERY LOW

 $^{^{\}it a}$ Downgraded by 1 increment if the majority of the evidence had serious limitations

Table 344: Risk factor for Radicular Symptoms (dichotomous outcome) for Referral for surgery (Sciatica population)-surgery: discectomy

Quality ass	essment			Adjusted effects	Quality			
Number of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other consideration s, including publication bias where possible	Pooled effect with 95% Cls [if meta-analysed]	
Effects for l	eg pain greater t	han back pain on	50% improvement in	pain assessed by \	/AS in one year (Ad	justed ORs) [adul	ts with Sciatica]	
1	Cohort studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	None	Adjusted OR: 1.02 [0.70, 1.48]	LOW

b95% CI around the median crosses null line.

^a Downgraded by 1 increment if the majority of the evidence had serious limitations

Table 345: Risk factor for Radicular Symptoms (dichotomous outcome) for Referral for surgery (Sciatica population)-surgery: discectomy

Quality asse	ssment			Adjusted effects	Quality			
Number of studies	Study design	Risk of bias	Inconsistency Indirectness on 30% improvement in function ass		Imprecision	Other consideration s, including publication bias where possible	Pooled effect with 95% Cls [if meta-analysed]	
Effects for le	g pain greater t	han back pain on	30% improvement in	function assessed	by ODI in one year	(Adjusted ORs) [a	adults with Sciatica]	
1	Cohort studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	None	Adjusted OR: 1.71[1.18, 2.47]	LOW

^a Downgraded by 1 increment if the majority of the evidence had serious limitations

Table 346: Risk factor for Radicular Symptoms (dichotomous outcome) for Referral for surgery (Sciatica population)-surgery: discectomy

Quality asse	ssment						Adjusted effects	Quality
Number of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other consideration s, including publication bias where possible	Pooled effect with 95% CIs [if meta-analysed]	
Effects for lo	eg pain greater t	han back pain on	50% improvement in	function assessed	by ODI in one year	(Adjusted ORs) [a	adults with Sciatica]	
1	Cohort studies	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	None	Adjusted OR: 1.93 [1.35,2.77]	LOW

^a Downgraded by 1 increment if the majority of the evidence had serious limitations

J.19 Disc replacement

Table 347: Clinical evidence profile: Disc replacement vs Spinal fusion (low back pain with/without sciatica)

			Quality as	sessment			No of pati	ents		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Disc replacement	Spinal fusion	Relative (95% CI)	Absolute		
Quality of	life (SF-36 m	ental com	ponent summary	score, 0-100) ≤ 4	months (3 mo	nths)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	393	166	-	MD 2.8 higher (0.65 to 4.95 higher)	⊕⊕OO LOW	CRITICAL
Quality of	life (SF-36 p	hysical co	mponent summar	y score, 0-100) ≤	4 months (3 m	onths)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	393	166	-	MD 4.5 higher (2.75 to 6.25 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	life (SF-36 m	ental com	ponent summary	score, 0-100) >4	months (1 year)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	393	163	-	MD 2 higher (0.09 lower to 4.09 higher)	⊕⊕OO LOW	CRITICAL
Quality of	life (SF-36 p	hysical co	mponent summai	y score, 0-100) >	4 months (1 yea	ar)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	393	163	-	MD 3.1 higher (0.96 to 5.24 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	life (SF-36 m	ental com	ponent summary	score, 0-100) > 4	l months (2 year	rs)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	379	145	-	MD 1.4 higher (0.71 lower to 3.51 higher)	⊕⊕OO LOW	CRITICAL
Quality of	life (SF-36 p	hysical co	mponent summar	y score, 0-100) >	4 months (2 ye	ars)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	379	145	-	MD 3 higher (0.68 to 5.32 higher)	⊕OOO VERY	CRITICAL

											LOW	
Quality o	of life (EQ-5D,	0-1) >4 mc	onths (1 year)									
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	80	72	1	MD 0.08 higher (0.01 lower to 0.17 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	of life (EQ-5D,	0-1) > 4 m	onths (2 years)									
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	80	72	-	MD 0.02 lower (0.11 lower to 0.07 higher)	⊕⊕OO LOW	CRITICAL
Function	(ODI, 0-100) ≤	4 months	s (3 months)									
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	393	166	-	MD 8.6 lower (11.76 to 5.44 lower)	⊕OOO VERY LOW	CRITICAL
Function	(ODI, 0-100) >	4 months	(1 year)	•		·						
2	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	473	235	-	MD 5.9 lower (8.87 to 2.92 lower)	⊕OOO VERY LOW	CRITICAL
Function	(ODI, 0-100) >	4 months	s (2 years)	•	•	•						
2	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	459	217	-	MD 4.69 lower (7.86 to 1.52 lower)	⊕OOO VERY LOW	CRITICAL
Pain sev	erity (Back pa	in NRS, 0-	10) ≤ 4 months (3	months)								
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	393	166	-	MD 0.92 lower (1.35 to 0.49 lower)	⊕OOO VERY LOW	CRITICAL
Pain sev	erity (Back pa	in VAS/NF	RS, 0-10) >4 mont	hs (1 year)	•	•						
2	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	473	235	-	MD 0.73 lower (1.15 to 0.31 lower)	⊕OOO VERY LOW	CRITICAL
Pain sev	erity (Back pa	in VAS/NR	RS, 0-10) > 4 mon	ths (2 years)								

	ı	1	1	1		1			1		1	
2	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	459	217	-	MD 0.51 lower (0.96 to 0.06 lower)	⊕⊕OO LOW	CRITICAL
Pain seve	rity (Leg pain	NRS, 0-1	0) ≤ 4 months (3 n	nonths)								
1		very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	393	166	-	MD 0.06 higher (0.37 lower to 0.49 higher)	⊕⊕OO LOW	CRITICAL
Pain seve	rity (Leg pain	VAS/NRS	6, 0-10) >4 months	(1 year)								
2	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	473	235	-	MD 0.57 lower (0.97 to 0.18 lower)	⊕⊕OO LOW	CRITICAL
Pain seve	rity (Leg pain	VAS/NRS	6, 0-10) > 4 months	s (2 years)								
2	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	459	217	-	MD 0.38 lower (0.82 lower to 0.05 higher)	⊕⊕OO LOW	CRITICAL
Adverse e	events (numb	er of patie	ents) ≤ 4 months (operative)								
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	59/405 (14.6%)	8.7%	RR 1.67 (0.98 to 2.86)	58 more per 1000 (from 2 fewer to 162 more)	⊕OOO VERY LOW	IMPORTAN T
Adverse e	events (possil	oly device	-related; number	of patients) ≤ 4 n	nonths (operativ	/e)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^c	none	2/405 (0.49%)	0%	RR 2.13 (0.10 to 44.15)	-	⊕OOO VERY LOW	IMPORTAN T
Reoperati	ons (number	of patient	s) > 4 months (2 y	ears)								
2	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^c	none	45/459 (9.8%)	10%	RR 0.97 (0.59 to 1.57)	3 fewer per 1000 (from 41 fewer to 57 more)	⊕OOO VERY LOW	IMPORTAN T
Reoperati	ons (number	of patient	:s) > 4 months (5 y	rears) - reoperati	ions at 5 years		<u>'</u>		!			
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	very serious ^c	none	5/80 (6.3%)	8.3%	RR 0.75 (0.24 to 2.35)	21 fewer per 1000 (from 63 fewer to 112 more)	⊕OOO VERY LOW	IMPORTAN T
Device-re	lated reopera	tions (nur	nber of events) >	4 months (5 year	rs)							

		- 3		no serious indirectness	Serious ^b	none	9/80 (11.3%)	27.8%	RR 0.41 (0.2 to 0.83)	164 fewer per 1000 (from 47 fewer to 222 fewer)	0000	IMPORTAN T
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Table 348: Clinical evidence profile: Disc replacement vs 3-elements MBR (low back pain without sciatica)

			Quality as	sessment			No of pa	ntients		Effect	Overlife.	Importanc
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Disc replacement	3-elements MBR	Relative (95% CI)	Absolute	Quality	e
Quality of	life (EQ-5D, 0)-1) >4 moı	nths (1 year)									
1	randomised trials	very seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	86	86	-	MD 0.13 higher (0.03 to 0.23 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	life (EQ-5D, 0)-1) > 4 mo	onths (2 years)									
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	none	none	86	86	-	MD 0.06 higher (0.03 lower to 0.15 higher)	⊕000 VERY LOW	CRITICAL
Quality of	life (SF-36 me	ental com	ponent summary s	core, 0-100) >4 m	onths (1 year)							
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	86	86	-	MD 1 higher (2.77 lower to 4.77 higher)	⊕⊕OO LOW	CRITICAL
Quality of	life (SF-36 ph	ysical cor	mponent summary	score, 0-100) >4	months (1 year)							
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	86	86	-	MD 5.5 higher (2.03 to 8.97 higher)	⊕OOO VERY LOW	CRITICAL
Quality of	life (SF-36 me	ental comp	ponent summary s	core, 0-100) > 4 n	nonths (2 years)							

 ^a Downgraded by 2 increments if the majority of the evidence was at very high risk of bias
 ^b Downgraded by 1 increment if the confidence interval crossed one MID
 ^c Downgraded by 2 increments if the confidence interval crossed both MIDs

	1	1	1		1	1	1			1		
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	no serious imprecision	none	86	86	-	MD 2.1 higher (1.55 lower to 5.75 higher)	⊕⊕OO LOW	CRITICAL
Quality o	f life (SF-36 pl	nysical cor	mponent summary	score, 0-100) > 4	months (2 year	s)						
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	86	86	-	MD 5.6 higher (2.33 to 8.87 higher)	⊕OOO VERY LOW	CRITICAL
Pain seve	erity (Back pai	n VAS, 0-1	10) >4 months (1 ye	ear)								
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	86	86	-	MD 1.76 lower (2.61 to 0.91 lower)	⊕⊕OO LOW	CRITICAL
Pain seve	erity (Back pai	n VAS, 0-1	(2 y	ears)								
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	86	86	-	MD 1.43 lower (2.29 to 0.57 lower)	⊕OOO VERY LOW	CRITICAL
Function	(ODI, 0-100) ≤	4 months	(3 months)									
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	86	86	1	MD 9.1 lower (13.17 to 5.03 lower)	⊕⊕OO LOW	CRITICAL
Function	(ODI, 0-100) >	4 months	(1 years)		_							
1	randomised trials	very serious ^a	no serious inconsistency	no serious indirectness	Serious ^b	none	86	86	-	MD 8.9 lower (13.88 to 3.92 lower)	⊕OOO VERY LOW	CRITICAL
Function	(ODI, 0-100) >	4 months	(2 years)									
1	randomised trials	very seriousª	no serious inconsistency	no serious indirectness	Serious ^b	none	86	86	-	MD 6.9 lower (11.57 to 2.23 lower)	⊕OOO VERY LOW	CRITICAL

^a Downgraded by 2 increments if the majority of the evidence was at very high risk of bias ^b Downgraded by 1 increment if the confidence interval crossed one MID

NICE. 2016 **Spinal fusion**

Table 349: Clinical evidence profile: Fusion versus Usual Care

			promer rusion v									
			Quality as	sessment			No of pa	atients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Spinal Fusion	Usual Care	Relative (95% CI)	Absolute		
Pain Seve	rity(VAS,0-10)	>4 months	s (2 years) (follow-	up 2 years; Better	indicated by low	ver values)						
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	201	63	-	MD 1.51 lower (2.09 to 0.93 lower)	⊕OOO VERY LOW	CRITICAL
Function(0	DDI,0-100) >4 ı	months (2	years) (Better indic	cated by lower va	lues)							
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	201	63	-	MD 9.9 lower (14.59 to 5.21 lower)	⊕OOO VERY LOW	CRITICAL
Adverse e	vents-Compli	cations (2	years)									
	randomised trials	very serious¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	48/211 (22.7%)	0/72 (0%)	OR 5 (2.45 to 10.19)	-	⊕OOO LOW	CRITICAL
Function(0	General Funct	ion Score,	GFS,0-100) >4 mon	ths (2 years) (Be	tter indicated by	lower values)						
	randomised trials	very serious¹	no serious inconsistency	no serious indirectness	serious ²	none	201	63	-	MD 11.4 lower (17.29 to 5.51 lower)	⊕000 VERY LOW	CRITICAL
Function(N	MillionVAS,MV	/AS,0-100)	>4 months (2 years	s) (Better indicate	ed by lower value	s)						
	randomised trials	very serious¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	201	63	-	MD 14.8 lower (20.11 to 9.49 lower)	⊕OOO LOW	CRITICAL
Reoperation	ons (2 years)											
1	randomised	very	no serious	no serious	no serious	none	16/211	0/72	OR 4.12 (1.3	-	⊕000	IMPORTANT

	trials	serious ¹	inconsistency	indirectness	imprecision	(7.6%)	(0%)	to 13.1)	LOW	

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MID's.

Table 350: Clinical evidence profile: Fusion versus Usual Care (cohort)

			Quality asses	sment			No of patient	ts		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Spinal Fusion versus Usual Care	Control	Relative (95% CI)	Absolute		
Quality of	life, SF-36(PCS,	0-100) >4	months - 1 year (fo	llow-up >4 montl	ns - 1 year; B	etter indicated by	lower values)		1			
	observational studies	very serious ¹		no serious indirectness	serious ²	none	53	43	-	MD 1.9 higher (1.12 lower to 4.92 higher)	⊕000 VERY LOW	CRITICAL
Quality of	life, SF-36(MCS,	0-100) >4	months - 1 year (fo	ollow-up >4 mont	hs - 1 year; E	Setter indicated by	lower values)					
1	observational studies	very serious ¹		no serious indirectness	serious ²	none	53	43	-	MD 2.6 lower (6.96 lower to 1.76 higher)	⊕000 VERY LOW	CRITICAL
Pain Seve	rity(NRS,0-10) >4	4 months -	· 1 year (follow-up >	>4 months - 1 yea	ır; Better ind	icated by lower va	lues)					
1	observational studies	very serious ¹		no serious indirectness	serious ²	none	53	43	-	MD 0.8 lower (1.94 lower to 0.34 higher)	⊕OOO VERY LOW	CRITICAL
Function (ODI,0-100)>4 mo	onths - 1 y	ear (follow-up >4 m	onths - 1 year; B	etter indicate	ed by lower values)				<u> </u>	
1	observational studies	very serious ¹		no serious indirectness	serious ²	none	53	43	-	MD 1.1 higher (7.87 lower to 10.07 higher)	⊕000 VERY LOW	CRITICAL

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MID's.

Table 351: Clinical evidence profile: Fusion versus Other treatment

			Quality as	sessment			No of	patients		Effect	Quality	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Spinal Fusion	Other Treatment	Relative (95% CI)	Absolute	Quality	Importance
Pain Seve	rity(VAS,0-10) >4 montl	ns - 1 year (1 year) (MBR) (Better i	ndicated by lowe	er values)						
2	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	63	55	-	MD 0.4 lower (1.29 lower to 0.48 higher)	⊕OOO VERY LOW	CRITICAL
Pain Seve	rity(VAS,0-10	, Mixed Mo	odality exercise: a	naerobic +biome	chanical) >4 mo	nths - 1 year (1 ye	ar) (Better	indicated by	lower va	lues)		
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	21	20	-	MD 2.83 lower (5.68 lower to 0.02 higher)	⊕OOO VERYLOW	CRITICAL
Pain Seve	erity(VAS,0-10	, Mixed Mo	odality exercise: a	naerobic +biome	chanical) >4 mo	nths (2 year) (Bett	er indicate	ed by lower v	alues)			
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	21	20	-	MD 3.06 lower (6.08 to 0.04 lower)	⊕OOO VERY LOW	CRITICAL
Function(ODI,0-100, 3 e	element MI	BR) >4 months - 1	year (1 year) (Be	etter indicated b	y lower values)						
2	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	63	55	-	MD 0.83 higher (6.03 lower to 7.7 higher)	⊕000 LOW	CRITICAL
Function(ODI, 0-100, M	ixed Moda	lity: aerobic+ bion	nechanical exerc	ise) >4 months -	- 1 year (1 year) (Be	etter indica	ated by lower	values)			
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	21	20	-	MD 26.06 lower (47.47 to 4.65 lower)	⊕OOO VERY LOW	CRITICAL
Function(ODI,0-100, 3 e	element Mi	BR) >4 months (2	year) (Better ind	icated by lower	values)						
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	176	173	_	MD 2.1 lower (6.47 lower to 2.27 higher)	⊕000 VERY LOW	CRITICAL

	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	21	20	-	MD 26.59 lower (44.82 to 8.36 lower)	⊕OOO VERY LOW	CRITICA
nctior	(General Fund	tion Score	e, GFS,, 0-100) >4	months - 1 year	(1 year) (Better i	ndicated by lower	values)					
	randomised trials	very serious ¹	serious inconsistency ³	no serious indirectness	very serious ²	none	63	55	-	MD 0.93 higher (10.12 lower to 11.97 higher))	⊕000 VERY LOW	CRITICA
ain Sev	erity(Japanes	e Orthopa	edic Association	Score,JOAS,0-3,	, Mixed Modality:	aerobic + biomech	anical exe	rcise) >4 mon	ths - 1 y	ear (1 year) (Better ind	icated by lowe	r values)
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	21	20	-	MD 0.96 higher (0.36 to 1.56 higher)	⊕OOO VERY LOW	CRITICA
ain Sev	erity(Japanes	e Orthopa	edic Association	Score,JOAS,0-3,	, Mixed Modality:	aerobic + biomech	anical exe	rcise) >4 mon	ths (2 ye	ear) (Better indicated by	lower values	
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	21	20	-	MD 1.16 higher (0.4 to 1.92 higher)	⊕000 VERY LOW	CRITICA
F36 at 2	2 years - Physi	ical compo	onent score, PCS	(Better indicated	d by lower values	3)						
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	115	131	1	MD 1.2 higher (2.5 lower to 4.9 higher)	⊕OOO LOW	CRITICA
F36 at 2	2 years - Menta	al compon	ent score, MSC (Better indicated	by lower values)							
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	115	131	1	MD 0.7 lower (3.79 lower to 2.39 higher)	⊕000 LOW	CRITICA
F36 at 2	2 years - Doma	ain-Genera	l health perception	on (Better indica	ted by lower valu	es)						
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	115	131	-	MD 3.9 higher (2.12 lower to 9.92 higher)	⊕OOO LOW	CRITICA
F36 at 2	2 years - Doma	ain-Physic	al functioning (Be	etter indicated by	y lower values)							
	randomised	very serious ¹	no serious inconsistency	no serious indirectness	no serious	none	115	131	-	MD 0.2 higher (6.92 lower to 7.32 higher)	⊕OOO LOW	CRITICA

			,							T	T	
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	115	131	-	MD 0.2 lower (10.98 lower to 10.58 higher)	⊕OOO LOW	CRITICAL
SF36 at 2	years - Doma	ain-Role lir	mitation(physical)	(Better indicated	by lower values	5)						
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	115	131	-	MD 1 higher (9.61 lower to 11.61 higher)	⊕000 LOW	CRITICAL
SF36 at 2	years - Doma	ain-Pain (B	Setter indicated by	lower values)								
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	115	131	-	MD 3.2 higher (3.26 lower to 9.66 higher)	⊕OOO LOW	CRITICAL
SF36 at 2	years - Doma	ain-Social	functioning (Bette	r indicated by lo	wer values)							
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	115	131	-	MD 2 lower (8.56 lower to 4.56 higher)	⊕OOO LOW	CRITICAL
SF36 at 2	years - Doma	ain-Mental	Health (Better ind	icated by lower	/alues)							
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	115	131	-	MD 1.9 lower (7.48 lower to 3.68 higher)	⊕OOO LOW	CRITICAL
SF36 at 2	years - Doma	ain-Energy	and vitality (Bette	er indicated by lo	wer values)							
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	115	131	-	MD 0.3 higher (5.66 lower to 6.26 higher)	⊕OOO LOW	CRITICAL
Healthca	re Utilisation(unplanne	d hospital admissi	ions for spinal in	jury, mean no. p	er patient, 3 eleme	nt MBR) (2 year) (Bette	r indicat	ed by lower values)		
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	176	173	-	MD 0.24 lower (0.32 to 0.16 lower)	⊕OOO VERY LOW	IMPORTANT
Healthcar	e Utilisation(GI	P consultati	ons, mean no. per pa	atient, 3 element l	MBR) (2 year) (B	etter indicated by I	ower value	es)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	176	173	-	MD 0.57 higher (1.29 lower to 2.43 higher)	⊕000 LOW	IMPORTANT
Healthca	re Utilisation(l	Practice N	urse consultation	s, mean no. per p	oatient, 3 elemen	t MBR) (2 year) (Be	etter indica	ated by lower	values)			
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	176	173	-	MD 0.24 higher (0.17 lower to 0.65 higher)	⊕000 LOW	IMPORTANT

Healthcar	e Utilisation(G	GP home v	risits, mean no. pe	r patient, 3 eleme	ent MBR) (2 yea	r) (Better indicated	by lower	values)						
1	randomised trials	- ,	no serious inconsistency	no serious indirectness	serious ²	none	176	173	-	MD 0.38 higher (0.07 to 0.69 higher)	⊕000 VERY LOW	IMPORTANT		
Healthcar	Healthcare Utilisation(Practise nurse home visits, mean no. per patient, 3 element MBR) (2 year) (Better indicated by lower values)													
1	randomised trials	- J .	no serious inconsistency	no serious indirectness	serious ²	none	176	173	-	MD 0.37 higher (0.02 to 0.72 higher)	⊕OOO VERY LOW	IMPORTANT		
Healthcar	Healthcare Utilisation(Prescriptions, mean no. per patient, 3 element MBR) (2 year) (Better indicated by lower values)													
1	randomised trials	- ,	no serious inconsistency	no serious indirectness	no serious imprecision	none	176	173	-	MD 0.8 higher (4.21 lower to 5.81 higher)	⊕000 LOW	IMPORTANT		

¹Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MID's.

Table 352: Clinical evidence profile: Fusion versus Different type of surgery

			Quality as	sessment			No o	f patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Spinal Fusion	Different type of surgery	Relative (95% CI)	Absolute		
Pain Seve	erity(VAS/NRS	5,0-10) ≤4	months (3 month) (Better indicate	ed by lower valu	es)						
1		very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	172	405	-	MD 0.92 higher (0.5 to 1.34 higher)	⊕OOO LOW	CRITICAL
Pain Seve	erity(VAS/NRS	5,0-10) >4	months - 1 year (1 year) (Better in	dicated by lowe	er values)						
2	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	244	485	-	MD 0.73 higher (0.32 to 1.14 higher)	⊕OOO LOW	CRITICAL
Pain Seve	Pain Severity(VAS/NRS,0-10) >4 months (2 year) (Better indicated by lower values)											
2	randomised	very	serious	no serious	no serious	none	244	485	-	MD 0.1 lower (0.89	⊕000	CRITICAL

³ Heterogeneity unexplained by subgroup analysis, random effects used

	trials	serious ¹	inconsistency ³	indirectness	imprecision					lower to 0.69 higher)	VERY LOW	
Function	(ODI,0-100) ≤₄	1 months	(3 month) (Better	indicated by lov	ver values)							
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	172	405	-	MD 8.6 higher (4.6 to 12.6 higher)	⊕OOO LOW	CRITICAL
Function	(ODI,0-100) >4	4 months	- 1 year (Better in	idicated by lowe	r values)							
2	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	244	485	ı	MD 5.9 higher (2.98 to 8.83 higher)	⊕OOO LOW	CRITICAL
Function	(ODI,0-100) >4	4 months	- 2 year (Better in	dicated by lowe	r values)							
2	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	244	485	ı	MD 4.75 higher (1.74 to 7.77 higher)	⊕OOO LOW	CRITICAL
SF36(Phy	sical Compo	nent Scor	e,PCS,0-100)≤ 4 ı	months (3 mont	h) (Better indica	ted by lower values	s)					
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	172	405	-	MD 4.5 lower (6.22 to 2.78 lower)	⊕OOO VERY LOW	CRITICAL
SF36(Phy	sical Compo	nent Scor	re,PCS,0-100)> 4 i	months - 1 year (Better indicated	by lower values)						
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	172	405	-	MD 3.1 lower (5.19 to 1.01 lower)	⊕OOO LOW	CRITICAL
SF36(Phy	sical Compo	nent Scor	e,PCS,0-100)> 4 i	months - 2 year ((Better indicated	by lower values)						
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	172	405	-	MD 3 lower (5.16 to 0.84 lower)	⊕OOO LOW	CRITICAL
SF36(Mei	ntal Compone	ent Score,	MCS,0-100)≤ 4 m	onths (3 month)	(Better indicate	d by lower values)						
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	172	405	-	MD 2.8 lower (4.91 to 0.69 lower)	⊕OOO LOW	CRITICAL
SF36(Mei	ntal Compone	ent Score,	MCS,0-100)> 4 m	onths - 1 year (B	Setter indicated b	by lower values)						
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	172	405	-	MD 2 lower (4.05 lower to 0.05 higher)	⊕OOO LOW	CRITICAL

SF36(Me	ntal Compone	ent Score.	MCS,0-100)> 4 mc	onths - 2 year (B	etter indicated h	ov lower values)						
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	172	405	-	MD 1.4 lower (3.36 lower to 0.56 higher)	⊕000 LOW	CRITICAL
EQ5D >4	months - 1 ye	ear (Bettei	r indicated by low	er values)								
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	72	80	-	MD 0.08 lower (0.17 lower to 0.01 higher)	⊕000 VERY LOW	CRITICAL
EQ5D >4	months - 2 ye	ear (Bette	r indicated by low	er values)		•						
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	72	80	-	MD 0.02 higher (0.07 lower to 0.11 higher)	⊕OOO LOW	CRITICAL
Adverse	events-Comp	lications -	- 2 year									
2	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	360/477 (75.5%)	53.2%	RR 0.97 (0.9 to 1.05)	16 fewer per 1000 (from 53 fewer to 27 more)	⊕000 LOW	IMPORTANT
Adverse	events-Comp	lications -	- 5 year			•						
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	very serious ²	none	9/72 (12.5%)	(16.3%)	RR 0.77 (0.35 to 1.69)	37 fewer per 1000 (from 106 fewer to 112 more)	⊕000 VERY LOW	IMPORTANT
Adverse	events-surge	ry at adjad	cent level at 2 year	rs								
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	6/72 (8.3%)	(1.3%)	RR 6.67 (0.82 to 54.06)	71 more per 1000 (from 2 fewer to 663 more)	⊕000 VERY LOW	IMPORTANT
Reopera	tions - 2 year											
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	very serious ²	none	7/72 (9.7%)	(10%)	RR 0.97 (0.37 to 2.55)	3 fewer per 1000 (from 63 fewer to 155 more)	⊕000 VERY LOW	IMPORTANT
Reopera	tions - 5 year											
1	randomised	very	no serious	no serious	very serious ²	none	7/72		RR 0.86 (0.34	16 fewer per 1000	⊕000	IMPORTANT

	trials	serious ¹	inconsistency	indirectness			(9.7%)	(11.3%)	to 2.2)	(from 74 fewer to 135 more)	VERY LOW	
Adverse (events-Mortal	ity (2 yea	r)									
1				no serious indirectness	very serious²	none	3/405 (0.7%)	(0.6%)	RR 1.27 (0.13 to 12.16)	2 more per 1000 (from 5 fewer to 65 more)	⊕OOO VERY LOW	IMPORTANT

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias.

² Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MID's.

³Heterogeniety unexplained by subgroup analysis, random effects used.

Spinal decompression J.21

Table 353: Discectomy versus Usual Care

			Quality as:	sessment			No of patients		Effect	Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Sciatica due to herniated intervertebral disc- Discectomy	Usual Care	Relative (95% CI)	Absolute	Quanty	importance
Quality o	Quality of life, SF-36, 0-100 ≤4 months - Domain-Bodily pain (follow-up ≤4 months; Better indicated by lower values)											
		very serious¹	very serious ²	no serious indirectness	no serious imprecision	none	338	352	-	MD 8.35 higher (7.87 to 8.83 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	f life, SF-36, (0-100 ≤4 r	months - Domain-	Physical function	oning (follow-u	p ≤4 months; Bet	ter indicated by lower valu	ies)				
		very serious ¹	very serious ²	no serious indirectness	no serious imprecision	none	338	352	-	MD 9.26 higher (8.84 to 9.68 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	f life, SF-36, (0-100 ≤4 r	months - Domain-	Social function	ing (follow-up :	≤4 months; Better	· indicated by lower values	s)				
		- ,	no serious inconsistency	no serious indirectness	no serious imprecision	none	140	141	-	MD 2.3 higher (1.76 to 2.84 higher)	⊕⊕OO LOW	CRITICAL

Quality	y of life, SF-36,	0-100 ≤4	months - Domair	n-Physical role (follow-up ≤4 m	onths; Better indic	cated by lower values)		I			
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	140	141	-	MD 0.2 higher (0.54 lower to 0.94 higher)	⊕⊕OO LOW	CRITICA
Quality	y of life, SF-36,	0-100 ≤4	months - Domair	n-Emotional role	e (follow-up ≤4 ı	months; Better inc	licated by lower values)					
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	140	141	-	MD 3.1 higher (2.26 to 3.94 higher)	⊕⊕OO LOW	CRITICA
Quality	y of life, SF-36,	0-100 ≤4	months - Domair	n-Mental health	index (follow-u	p ≤4 months; Bette	er indicated by lower value	es)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	140	141	-	MD 9.1 higher (8.75 to 9.45 higher)	⊕⊕OO LOW	CRITICAL
Quality	y of life, SF-36,	0-100 ≤4 :	months - Domair	n-Vitality (follow	-up ≤4 months;	Better indicated I	oy lower values)					
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	140	141	-	MD 10.4 higher (10 to 10.8 higher)	⊕⊕OO LOW	CRITICAL
Quality	y of life, SF-36,	0-100 ≤4 :	months - Domair	n-General health	n perception (fo	llow-up ≤4 months	s; Better indicated by lowe	r values	s)			
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	140	141	-	MD 10.5 higher (10.14 to 10.86 higher)	⊕⊕OO LOW	CRITICA
Quality	y of life, SF-36,	0-100 >4	months - 1 year -	Domain-Bodily	/ pain (follow-u	p >4 months - 1 ye	ar; Better indicated by low	er valu	es)			
2	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ³	none	342	354	-	MD 3.3 higher (2.94 to 3.66 higher)	⊕OOO VERY LOW	CRITICA
Quality	y of life, SF-36,	0-100 >4	months - 1 year -	Domain-Physic	cal functioning	(follow-up >4 mon	ths - 1 year; Better indicat	ed by lo	wer values)	•		
2	randomised trials	very serious¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	342	354	-	MD 1.5 higher (1.08 to 1.92 higher)	⊕⊕OO LOW	CRITICAL
Quality	y of life, SF-36,	0-100 >4	months - 1 year -	Domain-Social	functioning (fo	ollow-up >4 month	s - 1 year; Better indicated	by low	er values)			
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	140	141	-	MD 4.5 higher (4.07 to 4.93 higher)	⊕⊕OO LOW	CRITICAL

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Quality o	of life, SF-36,	0-100 >4 :	months - 1 year -	Domain-Physic	al role (follow-	up >4 months - 1 y	year; Better indicated by	lower val	ues)			
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	140	141	-	MD 7.2 higher (6.37 to 8.03 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life, SF-36,	0-100 >4	months - 1 year -	Domain-Emotic	nal role (follow	/-up >4 months - 1	year; Better indicated b	y lower va	alues)			
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ³	none	140	141	-	MD 3.9 higher (3.23 to 4.57 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	of life, SF-36,	0-100 >4	months - 1 year -	Domain-Mental	health index (f	ollow-up 4 month	s; Better indicated by lo	wer values	s)			
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ³	none	140	141	-	MD 2.7 higher (2.37 to 3.03 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	of life, SF-36,	0-100 >4	months - 1 year -	Domain-Vitality	(follow-up >4	months - 1 year; B	Better indicated by lower	values)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	140	141	-	MD 3.2 higher (2.84 to 3.56 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life, SF-36,	0-100 >4 :	months - 1 year -	Domain-Genera	al health percep	otion (follow-up >4	1 months - 1 year; Better	indicated	by lower va	llues)		
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	140	141	-	MD 2.5 higher (2.11 to 2.89 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life, SF-36,	0-100 >4	months(2 year) -	Domain-Bodily	pain (follow-up	2 years; Better in	ndicated by lower values	s)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	very serious ³	none	186	187	-	MD 3.2 higher (2.07 lower to 8.47 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	of life, SF-36,	0-100 >4	months(2 year) -	Domain-Physic	al functioning	(follow-up 2 years	; Better indicated by low	er values				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ³	none	186	187	-	MD 0 higher (5.41 lower to 5.41 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	of life, EQ-5D,	0-1 ≤4 m	onths(3 months)	(follow-up 3 mo	onths; Better in	dicated by lower v	values)					
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	141	142	-	MD 0.06 higher (0.01 to 0.11 higher)	⊕⊕OO LOW	CRITICAL

Quality o	of life, EQ-5D,	0-1 >4 m	onths - 1 year(1 y	/ear) (follow-up	1 years; Better	indicated by lowe	er values)					
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	141	142	-	MD 0.02 higher (0.02 lower to 0.06 higher)	⊕⊕OO LOW	CRITICAL
Leg Pain	Severity(VA	S,0-10) ≤4	months (follow-	up ≤4 months; ∣	Better indicated	l by lower values)						
2	randomised trials	very serious ¹	very serious ²	no serious indirectness	serious ³	none	166	167	-	MD 1.39 lower (2.39 to 0.39 lower)	⊕OOO VERY LOW	CRITICAL
Leg Pain	Severity(VA	S,0-10) >4	l months - 1 year	(follow-up >4 n	nonths - 1 year;	Better indicated I	by lower values)					
2	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	166	167	-	MD 0.57 lower (0.87 to 0.28 lower)	⊕⊕OO LOW	CRITICAL
Leg Pain	Severity(VA	S,0-10) >4	l months(2 year)	(follow-up 2 ye	ars; Better indi	cated by lower va	lues)					
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ³	none	26	24	-	MD 0.9 lower (1.95 lower to 0.15 higher)	⊕000 VERY LOW	CRITICAL
Back Pai	in Severity(V	AS,0-10) ≤	4 months (follow	/-up ≤4 months:	Better indicate	ed by lower values	3)	- '				
2	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	166	167	-	MD 1.13 lower (1.18 to 1.08 lower)	⊕⊕OO LOW	CRITICAL
Back Pai	in Severity(V	AS,0-10) >	•4 months - 1 yea	r (follow-up >4	months - 1 yea	r; Better indicated	by lower values)					
2	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	166	166	-	MD 0.23 lower (0.28 to 0.18 lower)	⊕⊕OO LOW	CRITICAL
Back Pai	in Severity(V	AS,0-10) >	•4 months (2 yea	r) (follow-up 2	years; Better in	dicated by lower v	/alues)					
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ³	none	26	24	-	MD 1 lower (2.28 lower to 0.28 higher)	⊕OOO VERY LOW	CRITICAL
Pain Sev	verity(Back Pa	ain bothe	rsomeness, chan	ge score,0-6) ≤	4 months (follo	w-up 3 months; B	etter indicated by lower	values)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ³	none	198	211	-	MD 2.2 lower (3.46 to 0.94 lower)	⊕000 VERY	CRITICAL

											LOW	
Pain Sev	verity(Back Pa	in bothe	someness, chan	ge score,0-6) >	4 months - 1 ye	ar (1 year) (follow	v-up 1 years; Better indica	ted by lo	ower values)			
1	randomised trials	very serious¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	202	211	-	MD 1.6 lower (2.86 to 0.34 lower)	⊕⊕OO LOW	CRITICAL
Pain Sev	verity(Back Pa	in bothe	someness, chan	ge score,0-6) >	4 months (2 ye	ar) (follow-up 2 ye	ears; Better indicated by l	ower val	ues)			
1	randomised trials	very serious¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	186	187	-	MD 1.6 lower (2.92 to 0.28 lower)	⊕⊕OO LOW	CRITICAL
Function	n(RMDQ, final	score) ≤₄	l months (follow-	-up ≤4 months;	Better indicated	d by lower values)						
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	140	141	-	MD 3.1 lower (3.22 to 2.98 lower)	⊕⊕OO LOW	CRITICAL
Function	n(RMDQ final	score) >4	months - 1 year	(follow-up >4 m	onths - 1 year;	Better indicated b	by lower values)					
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	140	141	-	MD 0.8 lower (0.92 to 0.68 lower)	⊕⊕OO LOW	CRITICAL
Function	n(,ODI change	score) ≤	4 months (follow	-up ≤4 months;	Better indicate	d by lower values)					
2	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	224	237	-	MD 5.1 lower (8.91 to 1.3 lower)	⊕⊕OO LOW	CRITICAL
Function	n(,ODI change	score) >	4 months - 1 year	r (follow-up >4 ı	months - 1 year	; Better indicated	by lower values)					
2	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	228	239	-	MD 2.58 lower (6.47 lower to 1.3 higher)	⊕⊕OO LOW	
Function	n(,ODI change	score) >	4 months (2 year) (follow-up 2 ye	ears; Better ind	icated by lower va	alues)					
2	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	212	211	-	MD 3.38 lower (7.33 lower to 0.58 higher)	⊕⊕OO LOW	CRITICAL
Respond	der criteria (co	omplete o	r nearly complete	e disappearanc	e of symptoms) ≤ 4 months(8 we	eeks) (follow-up 8 weeks)	_		, <i>,</i>		
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	86/140 (61.4%)	31.2%	RR 1.97 (1.49 to 2.6)	303 more per 1000 (from 153 more to 499 more)	⊕⊕OO LOW	IMPORTANT

Respond	ler criteria (co	omplete o	r nearly complete	e disappearance	e of symptoms)	> 4 months(26 w	veeks) (follow-up 26 weeks	s)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ³	none	127/140 (90.7%)	66%	RR 1.38 (1.21 to 1.57)	251 more per 1000 (from 139 more to 376 more)	⊕OOO VERY LOW	IMPORTANT
Reopera	tions (1 year)	(follow-u	p 1 years)									
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious³	none	2/26 (7.7%)	0%	OR 7.12 (0.43 to 117.44)	-	⊕OOO VERY LOW	IMPORTANT
Reopera	tions (2 years	5)										
2	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	27/269 (10%)	0%	OR 8.33 (3.85 to 18.04)	-	⊕⊕OO LOW	IMPORTANT
Adverse	events(intrac	perative	complications) ≤	4 months (follo	w-up ≤4 month	s)						
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	13/243 (5.3%)	0%	OR 8.27 (2.75 to 24.86)	-	⊕⊕OO LOW	
Adverse	events(posto	perative	complications/ev	ents) ≤ 4 month	s(8 weeks) (fo	llow-up 8 weeks)						
1	randomised trials	· ,	no serious inconsistency	no serious indirectness	no serious imprecision	none	13/243 (5.3%)	0%	OR 8 (2.66 to 24.05)	-	⊕⊕OO LOW	IMPORTANT
Healthca	re Utilisation	(Number	of patients with a	dditional physi	cal therapy visi	ts)> 4 months (2 y	year) (follow-up 2 years)					
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	Serious ³	none	8/26 (30.8%)	62.5%	RR 0.49 (0.26 to 0.95)	319 fewer per 1000 (from 31 fewer to 463 fewer)	⊕OOO VERY LOW	IMPORTANT

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Downgraded by 1 or 2 increments because of Heterogeneity, I2=50%, p=0.04, unexplained by subgroup analysis.

³ Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

Table 354: Discectomy versus usual care (cohort and RCT+cohort)

servational	Risk of bias 00 ≤4 mo	Inconsistency	Indirectness	Imprecision	Other	No of patients Ef					Importance
servational	00 ≤4 mo	nthe(3 month)			considerations	herniated disc- Discectomy	Usual care	Relative (95% CI)	Absolute		
		iiiiis(3 iii0iiiii) - i	ا Domain-Bodily إ	pain (follow-up	3 months; Better in	ndicated by lower va	lues)				
udies	,	no serious inconsistency	no serious indirectness	serious ²	none	466	190	-	MD 14.9 higher (10.77 to 19.03 higher)	⊕OOO VERY LOW	CRITICAL
fe, SF-36, 0-1	00 ≤4 mo	nths(3 month) - I	Domain-Physica	al functioning (f	ollow-up 3 months	; Better indicated by	lower va	lues)			
		no serious inconsistency	no serious indirectness	serious ²	none	466	190	-	MD 15.4 higher (11.53 to 19.27 higher)	⊕OOO VERY LOW	CRITICAL
fe, SF-36, 0-1	00 >4 mo	nths - 1 year(1 ye	ear) - Domain-B	odily pain (follo	w-up 1 years; Bett	er indicated by lowe	r values)				
		no serious inconsistency	no serious indirectness	serious ²	none	460	171	-	MD 10.8 higher (6.5 to 15.1 higher)	⊕OOO VERY LOW	CRITICAL
fe, SF-36, 0-1	00 >4 mo	nths - 1 year(1 ye	ear) - Domain-P	hysical function	ning (follow-up 1 ye	ears; Better indicated	d by lowe	r values)	l		
	,	no serious inconsistency	no serious indirectness	serious ²	none	460	171	-	MD 15.1 higher (10.9 to 19.3 higher)	⊕OOO VERY LOW	CRITICAL
s ic	servational dies e, SF-36, 0-1 servational dies e, SF-36, 0-1 servational dies	servational very serious¹ e, SF-36, 0-100 >4 mo servational very serious¹ e, SF-36, 0-100 >4 mo servational very serious¹ servational very serious¹	servational dies serious no serious inconsistency serious no serious inconsistency serious no serious inconsistency servational dies serious no serious inconsistency servational very servational very serious no serious inconsistency serious no serious inconsistency serious inconsistency	servational dies serious inconsistency indirectness inconsistency indirectness inconsistency indirectness inconsistency indirectness inconsistency inconsistency inconsistency inconsistency inconsistency indirectness inconsistency indirectness inconsistency indirectness inconsistency indirectness inconsistency indirectness	servational very serious inconsistency inconsistency indirectness serious² a, SF-36, 0-100 >4 months - 1 year(1 year) - Domain-Bodily pain (follower vational dies serious¹ inconsistency indirectness serious² a, SF-36, 0-100 >4 months - 1 year(1 year) - Domain-Physical function servational very serious¹ inconsistency inconsistency inconsistency serious² beginning the property of the property	servational very serious inconsistency indirectness serious serious indirectness serious none indirectness serious none indirectness serious none indirectness serious none serious serious none indirectness serious none serious indirectness serious none indirectness serious none indirectness serious none serious inconsistency inconsistency indirectness serious none indirectness serious none serious inconsistency indirectness serious none indirectness serious none serious inconsistency indirectness serious none serious inconsistency indirectness serious none serious indirectness serious none serious indirectness serious none serious indirectness serious none serious serious none serious indirectness serious none serious indirectness serious none serious serious serious none serious indirectness serious none serious indirectness serious none serious serious serious none serious indirectness serious none serious serious serious none serious serious serious none serious serious serious none serious serious serious serious serious none serious serious serious serious serious serious none serious	servational dies serious no serious indirectness serious none 466 e, SF-36, 0-100 >4 months - 1 year(1 year) - Domain-Bodily pain (follow-up 1 years; Better indicated by lowe servational very serious inconsistency indirectness serious none 460 e, SF-36, 0-100 >4 months - 1 year(1 year) - Domain-Physical functioning (follow-up 1 years; Better indicated servational very no serious no serious serious none 460	servational very serious¹ inconsistency indirectness serious² none 466 190 a, SF-36, 0-100 >4 months - 1 year(1 year) - Domain-Bodily pain (follow-up 1 years; Better indicated by lower values) servational very serious¹ no serious indirectness serious² none 460 171 a, SF-36, 0-100 >4 months - 1 year(1 year) - Domain-Physical functioning (follow-up 1 years; Better indicated by lower values) a, SF-36, 0-100 >4 months - 1 year(1 year) - Domain-Physical functioning (follow-up 1 years; Better indicated by lower values) servational very no serious inconsistency indirectness serious² none 460 171	dies serious¹ inconsistency indirectness e, SF-36, 0-100 >4 months - 1 year(1 year) - Domain-Bodily pain (follow-up 1 years; Better indicated by lower values) servational very serious¹ inconsistency indirectness serious² none 460 171 - e, SF-36, 0-100 >4 months - 1 year(1 year) - Domain-Physical functioning (follow-up 1 years; Better indicated by lower values) servational very serious¹ no serious inconsistency indirectness serious² none 460 171 - servational very serious¹ no serious indirectness indirectness serious² none 460 171 -	rervational very serious no serious indirectness serious none serious none were values none were values. Servational very serious no serious indirectness serious none were values. Servational very serious no serious indirectness none were values. Servational very serious no serious indirectness none were values. Servational very serious no serious indirectness none were values. Servational very serious no serious indirectness none were values. Servational very serious no serious indirectness none were values. Servational very serious no serious indirectness none were values. Servational very serious no serious indirectness none were values. Servational very serious no serious indirectness none were values. Servational very serious no serious indirectness none were values. Servational very serious no serious indirectness none were values. Servational very serious no serious indirectness none were values. Servational very serious no serious indirectness none were values. Servational very serious no serious indirectness none were values.	Pervational dies serious no serious no serious no serious no serious none 466 190 - MD 15.4 higher (11.53 to 19.27 higher) MD 2.7 higher) None 466 190 - MD 15.4 higher (11.53 to 19.27 higher) None 460 171 - MD 10.8 higher (6.5 ⊕OOO to 15.1 higher) None 460 171 - MD 10.8 higher (6.5 ⊕OOO to 15.1 higher) None 460 171 - MD 10.8 higher None 460 171 - NONE NONE

1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	456	165	-	MD 10.2 higher (5.9 to 14.5 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	f life, SF-36, 0-1	100 >4 mo	onths(2 year) - Do	omain-Physical	functioning (fo	llow-up 2 years; Be	tter indicated by low	er values	s)			
1		very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	456	165	-	MD 12 higher (7.8 to 16.2 higher)	⊕OOO VERY LOW	CRITICAL
Pain Sev	erity(Sciatica b	othersom	eness index, cha	inge score,0-24) ≤4 months (3	months) (follow-up	3 months; Better inc	licated by	y lower valu	es)		
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	466	190	-	MD 3.9 lower (4.93 to 2.87 lower)	⊕OOO VERY LOW	CRITICAL
Pain Sev	erity(Sciatica be	othersom	eness index, cha	inge score,0-24) >4 months - 1	year (1 year) (follo	w-up 1 years; Better i	ndicated	by lower va	llues)		
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	460	171	-	MD 2.6 lower (3.67 to 1.53 lower)	⊕OOO VERY LOW	CRITICAL
Pain Sev	erity(Sciatica b	othersom	eness index, cha	nge score,0-24) >4 months (2 y	/ear) (follow-up 2 y	ears; Better indicated	by lowe	er values)			
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	serious ¹	none	456	165	-	MD 2.1 lower (3.17 to 1.03 lower)	⊕OOO VERY LOW	CRITICAL
Function	(ODI change so	ore) ≤4 m	onths (follow-up	3 months; Bett	ter indicated by	lower values)						
1	observational studies	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	466	190	-	MD 15.2 lower (18.6 to 11.8 lower)	⊕OOO VERY LOW	CRITICAL
Function	(,ODI change so	core) 4 m	onths (1 year) (fo	ollow-up 1 years	s; Better indicat	ed by lower values)					
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	460	171	-	MD 15.3 lower (19.03 to 11.57	⊕OOO VERY	CRITICAL

										lower)	LOW	
Function	(,ODI change so	ore) ≤4 n	nonths (2 year) (follow-up 2 yea	rs; Better indica	ated by lower value	es)	1				
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	456	165	-	MD 13.4 lower (17.13 to 9.67 lower)	⊕OOO VERY LOW	CRITICAL
Pain Seve	erity(Back Pain	botherso	meness,0-6) ≤4 n	nonths (follow-	up 3 months; B	etter indicated by	ower values)					
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	775	416	-	MD 0.9 lower (0.91 to 0.89 lower)	⊕OOO VERY LOW	CRITICAL
Pain Seve	erity(Back Pain	botherso	meness,0-6) >4 r	nonths - 1 year	(1 year) (follow	v-up 1 years; Bette	r indicated by lower v	alues)				
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	775	416	-	MD 0.7 lower (0.71 to 0.69 lower)	⊕OOO VERY LOW	CRITICAL
Pain Seve	erity(Back Pain	botherso	meness,0-6) >4 r	nonths (2 year)	(follow-up 2 ye	ears; Better indicat	ed by lower values)					
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	775	416	-	MD 0.5 lower (0.51 to 0.49 lower)	⊕OOO VERY LOW	CRITICAL
Healthca	re Utilisation(N	umber of	patients with mo	re reported dia	gnostic test us	e)> 4 months (2 ye	ar) (follow-up 2 years)					
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	410/775 (52.9%)	33.9%	RR 1.56 (1.34 to 1.81)	190 more per 1000 (from 115 more to 275 more)	⊕OOO VERY LOW	IMPORTANT
Healthca	re Utilisation(N	umber of	patients with add	litional physica	I therapy visits)	> 4 months (2 year	r) (follow-up 2 years)					
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	383/775 (49.4%)	44%	RR 1.12 (0.99 to 1.28)	53 more per 1000 (from 4 fewer to 123 more)	⊕OOO VERY LOW	IMPORTANT

Low back pain and sciatica in over 16s Quality assessment

Healthcar	e Utilisation(N	umber of	patients with rep	orted healthcar	e visits)> 4 mor	nths (2 year) (follo	w-up 2 years)							
	observational studies		no serious inconsistency	no serious indirectness	no serious imprecision ²	none	698/775 (90.1%)	88%	RR 1.02 (0.98 to 1.07)	18 more per 1000 (from 18 fewer to 62 more)	⊕OOO VERY LOW	IMPORTANT		
Healthcar	Healthcare Utilisation(Medication use)> 4 months (2 year) (follow-up 2 years)													
	observational studies		no serious inconsistency	no serious indirectness	no serious imprecision	none	744/775 (96%)	88.9%	RR 1.08 (1.04 to 1.12)	71 more per 1000 (from 36 more to 107 more)		IMPORTANT		

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 355: Discectomy versus combination treatment (manual therapy+ biomechanical exercise + self-management)

			Quality ass	essment			No of	patients		Effect	Quality	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Sciatica due to herniated intervertebral disc- Discectomy	Manual therapy+ biomechanical exercise + self-management	Relative (95% CI)	Absolute	Quality	Importance
Quality o	f life, SF-36,	0-100 ≤4	months(12 wee	ks) - Domain-B	odily pain (fo	ollow-up 12 week	s; Better indicated by l	ower values)				
	randomised trials		no serious inconsistency	no serious indirectness	serious ²	none	20	20	-	MD 10.3 higher (2.37 lower to 22.97 higher)	⊕⊕OO LOW	CRITICAL
Quality o	f life, SF-36,	0-100 ≤4	months(12 wee	ks) - Domain-P	hysical role	(follow-up 12 wee	ks; Better indicated by	lower values)				
	randomised trials			no serious indirectness	very serious²	none	20	20	-	MD 3.7 lower (27.1 lower to 19.7 higher)	#000 VERY LOW	CRITICAL
Quality o	f life, SF-36,	0-100 ≤4	months(12 wee	ks) - Domain-E	motional role	e (follow-up 12 we	eeks; Better indicated I	by lower values)	ı			

² Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	very serious ²	none	20	20	-	MD 9.5 lower (34.49 lower to 15.49 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	of life, SF-36,	0-100 ≤4	months(12 wee	eks) - Domain-V	itality (follow	w-up 12 weeks; I	Better indicated by lower	values)				
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	very serious ²	none	20	20	-	MD 8.20 higher (3.37 lower to 19.77 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	of life, SF-36,	0-100 ≤4	months(12 wee	eks) - Domain-F	hysical fund	tion (follow-up	12 weeks; Better indicated	d by lower values)				
1	randomised trials		no serious inconsistency	no serious indirectness	very serious ²	none	20	20	-	MD 6.80 higher (9.64 lower to 23.24 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	of life, SF-36,	0-100 ≤4	months(12 wee	eks) - Domain-S	Social function	on (follow-up 12	weeks; Better indicated b	y lower values)				
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	very serious ²	none	20	20	-	MD 6.30 lower (23.79 lower to 11.19 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	of life, SF-36,	0-100 ≤4	months(12 wee	eks) - Domain-N	lental health	(follow-up 12 w	eeks; Better indicated by	lower values)				
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	very serious ²	none	20	20	-	MD 0.40 higher (5.61 lower to 6.41 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	of life, SF-36,	0-100 ≤4	months(12 wee	eks) - Domain-C	Seneral healt	h (follow-up 12 v	weeks; Better indicated b	y lower values)		•		
1	randomised trials	,	no serious inconsistency	no serious indirectness	very serious ²	none	20	20	-	MD 5.40 higher (5.61 lower to 6.41 higher)	⊕OOO VERY LOW	CRITICAL
Pain Sev	verity(McGill,	0-78) ≤ 4	months(12 wee	eks) (follow-up	12 weeks; B	etter indicated b	y lower values)			,		
1	randomised trials		no serious inconsistency	no serious indirectness	serious ²	none	20	20	-	MD 6.4 lower (3.40 lower to	⊕⊕OO LOW	CRITICAL

Quality ass	_ow back
assessm	pain
ent	and
	nd sciatica
	\exists
	over
	16

										14.20 higher)		
Function	(RMDQ,0-24)	≤4 mont	hs (follow-up 12	weeks; Better	indicated by	lower values)						
	randomised trials			no serious indirectness	serious ²	none	20	20	1	MD 1.8 lower (5.87 lower to 2.27 higher)	⊕⊕OO LOW	CRITICAL

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 356: Percutaneous decompression versus usual care

			Quality as	sessment			No of patients	Effect		Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Sciatica due to herniated intervertebral disc- Percutaneous disc decompression	Usual Care	Relative (95% CI)	Absolute		
Pain Sev	erity(Leg Pai	n NVS,0-	l 10) ≤4 months(3 r	nonths) (follow	up 3 months; E	Better indicated b	y lower values)		<u> </u>		<u>l</u>	
1				no serious indirectness	serious ²	none	31	31	-	MD 1.6 lower (2.95 to 0.25 lower)	⊕OOO VERY LOW	CRITICAL
Pain Sev	erity(Leg Pai	n NVS,0-	10) >4 months - 1	year(1 year) (fo	bllow-up 1 years	s; Better indicated	d by lower values)		1		1	
1				no serious indirectness	no serious imprecision	none	31	31	-	MD 2.8 lower (4.02 to 1.58 lower)	⊕⊕OO LOW	CRITICAL
Pain Sev	erity(Leg Pai	n NVS,0-	10) >4 months(2 y	/ears) (follow-u	p 2 years; Bette	r indicated by lov	wer values)					
1				no serious indirectness	no serious imprecision	none	31	31	-	MD 3.10 lower (4.45 to 1.75 lower)	⊕⊕OO LOW	CRITICAL

² Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

Table 357: Plasma disc decompression versus other treatment (epidural steroid)

			Quality as	sessment			No of pa	atients	E	Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Sciatica due to herniated intervertebral disc- Plasma disc decompression	Other treatment (Transforaminal epidural steroid injections)	Relative (95% CI)	Absolute	Quality	Importance
Pain Sev	erity(Leg P	ain VAS	0-10) ≤4 months	s(3 months) (fo	ollow-up 3 mo	nths; Better indi	cated by lower values)					
	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	45	40	-	MD 1.8 lower (3.05 to 0.55 lower)	⊕⊕OO LOW	CRITICAL
Pain Sev	verity(Leg P	ain VAS	0-10) >4 months	s - 1 year(6 mg	onths) (follow	-up 6 months; Bo	etter indicated by lower	values)				
	randomised trials	serious ¹		no serious indirectness	serious ²	none	45	40	-	MD 1.8 lower (3.05 to 0.55 lower)	⊕⊕OO LOW	CRITICAL
Pain Sev	erity(Back	Pain VA	5,0-10) ≤4 montl	ns(3 months) (follow-up 3 m	onths; Better inc	dicated by lower values					
	randomised trials	serious ¹	no serious inconsistency		no serious imprecision	none	45	40	-	MD 2.2 lower (3.18 to 1.22 lower)	⊕⊕⊕O MODERATE	CRITICAL
Pain Sev	erity(Back	Pain VA	S,0-10) >4 mont	hs - 1 year(6 m	onths) (follow	v-up 6 months; B	Setter indicated by lower	r values)				
	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	45	40	-	MD 1.62 lower (2.73 to 0.51 lower)	⊕⊕OO LOW	CRITICAL

	randomised trials		no serious inconsistency	no serious indirectness	serious ²	none	45	40	-	MD 1.2 lower (1.91 to 0.49 lower)	⊕⊕OO LOW	CRITICAL
nctio	n(ODI,0-100)	>4 mont	ths - 1 year (6 n	nonths) (follow	-up 6 months;	Better indicate	d by lower values)					
	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	45	40	-	MD 1.6 lower (2.31 to 0.89 lower)	⊕⊕⊕O MODERATE	CRITICAL
roced	ure related a	dverse e	vents> 4 month	ns (6 months) (follow-up 6 m	onths)						
	randomised trials		no serious inconsistency	no serious indirectness	very serious ²	none	5/45 (11.1%)	17.5%	RR 0.63 (0.22 to 1.84)	65 fewer per 1000 (from 137 fewer to 147 more)	⊕000 VERY LOW	IMPORTAN

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

Table 358: Discectomy versus fusion

			Quality asses	ssment			No of patients Effect				Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Sciatica due to herniated disc- Discectomy	Fusion	Relative (95% CI)	Absolute			
Function	(ODI 0-100) >4 r	nonths - 1	year (follow-up >	4 months - 1 ye	ar; Better inc	licated by lower v	alues)						
	observational studies		no serious inconsistency	no serious indirectness	serious ²	none	25	30		MD 1.52 lower (8.76 lower to 5.72 higher)	⊕OOO VERY LOW	CRITICAL	
Revision	evision surgery >4 months - 1 year (follow-up >4 months - 1 year)												

1	observational	very	no serious	no serious	serious ²	none	3/25	0%	OR 9.82	-	\oplus OOO	IMPORTANT
	studies	serious ¹	inconsistency	indirectness			(12%)		(0.97 to		VERY	
									99.53)		LOW	

Table 359: Laminectomy versus usual care

			Quality as	sessment			No of patients			Effect		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Sciatica due to stenosis (foraminal and/or canal)- Laminectomy versus Usual Care	Control	Relative (95% CI)	Absolute	Quality	Importance
Quality o	f life, SF-36,	l 0-100 ≤4 r	months - Domain	-Bodily pain (fo	llow-up 3 mont	hs; Better indicate	ed by lower values)	<u> </u>				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	116	135	-	MD 2.5 higher (4.16 lower to 9.16 higher)	⊕⊕OO LOW	CRITICAL
Quality o	f life, SF-36,	0-100 ≤4 r	months - Domain	-Physical functi	oning (follow-u	p 3 months; Bette	er indicated by lower values)		<u> </u>			
1	randomised trials	· ,	no serious inconsistency	no serious indirectness	no serious imprecision	none	116	135	-	MD 4.2 lower (10.86 lower to 2.46 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life, SF-36,	0-100 >4 ı	months - 1 year (1 year) - Domai	n-Bodily pain (follow-up 1 years;	Better indicated by lower value	s)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	120	126	-	MD 5.5 higher (0.74 lower to 11.74 higher)	⊕⊕OO LOW	CRITICAL
Quality o	of life, SF-36,	0-100 >4 ı	months - 1 year (1 year) - Domai	n-Physical fund	ctioning (follow-up	o 1 years; Better indicated by lov	ver value	es)			

¹ Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

² Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

1	randomised	very	no serious	no serious	no serious	none	120	126	-	MD 1.6 higher	⊕⊕OO	CRITICAL
	trials	serious ¹	inconsistency	indirectness	imprecision					(4.64 lower to 7.84 higher)	LOW	
Quality	of life, SF-36,	0-100 >4	months (2 year)	- Domain-Bodil	y pain (follow-ι	up 2 years; Better	indicated by lower values)					
1		very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	108	113	-	MD 7.8 higher (1.56 to 14.04 higher)	⊕OOO VERY LOW	CRITICAL
Quality	of life, SF-36,	0-100 >4	months (2 year)	- Domain-Physi	cal functioning	g (follow-up 2 year	s; Better indicated by lower valu	es)				
1		very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	108	113	-	MD 0 higher (6.52 lower to 6.52 higher)	⊕⊕OO LOW	CRITICAL
Pain Sev	verity(Low Ba	ck Pain b	othersomeness,	change score,0	0-24) ≤4 months	s (follow-up 3 mon	ths; Better indicated by lower va	lues)				
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	116	135	-	MD 0.4 higher (0.15 lower to 0.95 higher)	⊕⊕OO LOW	CRITICAL
Pain Sev	verity(Low Ba	ck Pain b	othersomeness,	change score,0	0-24) >4 months	s - 1 year (follow-u	p 1 years; Better indicated by lov	wer valu	es)	'		
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	120	126	-	MD 0 higher (0.55 lower to 0.55 higher)	⊕⊕OO LOW	CRITICAL
Pain Sev	verity(Low Ba	ck Pain b	othersomeness,	change score,0	0-24) >4 months	s (2 year) (follow-	up 2 years; Better indicated by lo	wer valu	ies)			
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	108	113	-	MD 0.3 higher (0.26 lower to 0.86 higher)	⊕⊕OO LOW	CRITICAL
Pain Sev	verity(Sciatica	Pain bot	hersomeness, ch	nange score,0-2	24) ≤4 months (follow-up 3 month	s; Better indicated by lower valu	es)		<u> </u>		
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	116	135	-	MD 0.3 lower (1.01 lower to 0.41 higher)	⊕⊕OO LOW	CRITICAL

	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	120	126	-	MD 0.6 lower (1.15 to 0.05	⊕000 VERY	CRITICAL
		00000								lower)	LOW	
ain Se	verity(Sciatica	Pain bot	hersomeness, c	hange score,0-2	24) >4 months	(2 year) (follow-	up 2 years; Better indicated by	lower values	s)			
	randomised	very	no serious	no serious	no serious	none	108	113	-	MD 0.4 lower	⊕⊕00	CRITICAL
	trials	serious ¹	inconsistency	indirectness	imprecision					(0.96 lower to	LOW	
										0.16 higher)		
inctio	n(,ODI change	score) ≤	4 months (follow	-up 3 months;	Better indicate	d by lower value	es)					
									1	1		
	randomised	very	no serious	no serious	no serious	none	116	135	-	MD 0.5 higher	$\oplus \oplus OO$	CRITICAL
	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	116	135	-	MD 0.5 higher (5.05 lower to	⊕⊕OO LOW	CRITICAL
						none	116	135	-			CRITICAL
unctio	trials	serious ¹		indirectness	imprecision			135	-	(5.05 lower to		CRITICAL
ınctic	trials	serious ¹	inconsistency	indirectness	imprecision			135	-	(5.05 lower to		CRITICAL
ınctic	trials on(,ODI change	serious ¹ score) >	inconsistency 4 months - 1 yea	indirectness	imprecision ears; Better in	dicated by lower	r values)		-	(5.05 lower to 6.05 higher)	LOW	
inctic	trials n(,ODI change	serious¹ score) >	inconsistency 4 months - 1 yea no serious	indirectness Ir (follow-up 1 y	imprecision ears; Better in no serious	dicated by lower	r values)		-	(5.05 lower to 6.05 higher) MD 2.2 lower	LOW	
	randomised trials	serious¹ score) > very serious¹	inconsistency 4 months - 1 yea no serious	indirectness Ir (follow-up 1 y no serious indirectness	ears; Better in no serious imprecision	dicated by lower	r values)		-	(5.05 lower to 6.05 higher) MD 2.2 lower (7.33 lower to	LOW	
	randomised trials	very serious ¹	inconsistency 4 months - 1 yea no serious inconsistency	indirectness Ir (follow-up 1 y no serious indirectness	ears; Better in no serious imprecision	dicated by lower	r values)		-	(5.05 lower to 6.05 higher) MD 2.2 lower (7.33 lower to 2.93 higher)	LOW	
	randomised trials	serious¹ score) > very serious¹	inconsistency 4 months - 1 yea no serious inconsistency 4 months (2 year	indirectness Ir (follow-up 1 y no serious indirectness r) (follow-up 2 y	imprecision ears; Better in no serious imprecision ears; Better in	none dicated by lower	r values)	126	-	(5.05 lower to 6.05 higher) MD 2.2 lower (7.33 lower to	LOW ⊕⊕OO LOW	CRITICA

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias ² Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

Table 360: Laminectomy versus usual care (cohort and RCT+ Cohort)

Quality assessment	No of patients	Effect	Quality Impor	ortance

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Sciatica due to stenosis (foraminal and/or canal)- Laminectomy versus Usual Care	Control	Relative (95% CI)	Absolute		
Quality o	of life, SF-36, 0-	100 ≤4 mo	onths - Domain-B	odily pain (follo	ow-up 3 months	s; Better indicated	by lower values)					
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	very serious ²	none	378	313	-	MD 16.1 higher (12.91 to 19.29 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	f life, SF-36, 0-	100 ≤4 mo	onths - Domain-P	hysical function	ning (follow-up	3 months; Better	indicated by lower values)					
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	378	313	-	MD 14.8 higher (11.48 to 18.12 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	f life, SF-36, 0-	100 >4 mo	onths - 1 year (1	year) - Domain-	Bodily pain (fo	llow-up 1 years; E	Better indicated by lower values)				
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	302	230	-	MD 14.5 higher (10.89 to 18.11 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	of life, SF-36, 0-	100 >4 mo	onths - 1 year (1	year) - Domain-	Physical funct	oning (follow-up	1 years; Better indicated by low	er values	5)			
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	302	230	-	MD 16 higher (12.39 to 19.61 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	of life, SF-36, 0-	100 >4 mo	onths (2 year) - D	l Domain-Bodily p	pain (follow-up	2 years; Better inc	dicated by lower values)					
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	335	198	-	MD 13.6 higher (9.99 to 17.21 higher)	⊕OOO VERY LOW	CRITICAL
Quality o	of life, SF-36, 0-	100 >4 mo	onths (2 year) - D	l Iomain-Physica	I functioning (f	ollow-up 2 years;	Better indicated by lower value	s)				
1	observational	very	no serious	no serious	serious ²	none	335	113	-	MD 11.2 higher (6.76 to 15.64	⊕000 VERY	CRITICAL

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	studies	serious ¹	inconsistency	indirectness						higher)	LOW	
ain Sev	 verity(Low Back	Pain bot	:hersomeness, c	hange score,0-2	24) ≤4 months (follow-up 3 month	l ns; Better indicated by lower values	ues)				
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	378	313	-	MD 1.2 lower (1.48 to 0.92 lower)	#000 VERY LOW	CRITICAL
Pain Sev	 verity(Low Back	Pain bot	 :hersomeness, c	hange score,0-2	24) >4 months -	1 year (follow-up	1 years; Better indicated by low	er values	5)			
1	randomised trials	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	302	230	-	SMD 3.00 lower (3.28 to 2.72 lower)	⊕⊕OO LOW	CRITICAL
Pain Sev	verity(Low Back	Pain bot	thersomeness, c	hange score,0-2	24) >4 months (2 year) (follow-up	o 2 years; Better indicated by lov	ver value	s)		1	
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	335	198	-	MD 0.9 lower (1.18 to 0.62 lower)	⊕000 VERY LOW	CRITICAL
Pain Sev	verity(Sciatica P	Pain both	ersomeness, cha	ange score,0-24) ≤4 months (fo	llow-up 3 months	 ; Better indicated by lower value	s)				
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	378	313	-	MD 1.8 lower (2.08 to 1.52 lower)	#000 VERY LOW	CRITICAL
Pain Sev	verity(Sciatica P	Pain both	ersomeness, cha	ange score,0-24) >4 months - 1	year (1 year) (fol	low-up 1 years; Better indicated	by lower	values)			
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	302	230	-	MD 1.2 lower (1.48 to 0.92 lower)	#000 VERY LOW	CRITICAL
Pain Sev	verity(Sciatica P	Pain both	ersomeness, cha	ange score,0-24) >4 months (2	year) (follow-up 2	! ! years; Better indicated by lowe	r values)		l	<u> </u>	l
1	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	335	198	-	MD 1.1 lower (1.38 to 0.82 lower)	#000 VERY LOW	CRITICAL

	observational studies		no serious inconsistency	no serious indirectness	no serious imprecision	none	378	313	-	MD 13.8 lower (16.44 to 11.16 lower)	⊕OOO VERY LOW	CRITICA
ınctior	n(,ODI change s	score) >4	months - 1 year (follow-up 1 yea	rs; Better indic	ated by lower valu	ues)					
	observational studies	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	302	230	-	MD 12.5 lower (15.41 to 9.59 lower)	⊕OOO VERY LOW	CRITICA
unction	n(,ODI change s	score) >4	months (2 year) (follow-up 2 yea	rs; Better indic	ated by lower valu	ues)					
	observational	very	no serious	no serious	serious ²	none	335	198	-	MD 11.2 lower	⊕OOO	CRITICA

¹ Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 361: Discectomy versus fusion

	<u> </u>		Quality asso	essment	No of patients				_	Importance		
No of studies	Design Inconsistency Indirect		Indirectness	Imprecision	Other considerations	Sciatica due to stenosis- Discectomy	Fusion	Relative (95% CI)	Absolute			
Adverse ev	vents (complicati	ions) >4 m	onths - 1 year (folio	ow-up >4 months	- 1 year)							
					no serious imprecision	none	0/47 (0%)	0%	not pooled	not pooled	⊕000 VERY LOW	IMPORTANT

² Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

References

- Anema JA, Steenstra IA, Bongers PM, de Vet HC, Knol DL, Loisel P et al. Multidisciplinary rehabilitation for subacute low back pain: graded activity or workplace intervention or both? a randomized controlled trial. Spine. 2007; 32(3):291-298
- Apeldoorn AT, Bosmans JE, Ostelo RW, de Vet HC, van Tulder MW. Cost-effectiveness of a classification-based system for sub-acute and chronic low back pain. European Spine Journal. 2012; 21(7):1290-1300
- Apeldoorn AT, Ostelo RW, van Helvoirt H, Fritz JM, de Vet HCW, van Tulder MW. The costeffectiveness of a treatment-based classification system for low back pain: design of a randomised controlled trial and economic evaluation. BMC Musculoskeletal Disorders. 2010; 11:58
- 4 Apeldoorn AT, Ostelo RW, van Helvoirt H, Fritz JM, Knol DL, van Tulder MW et al. A randomized controlled trial on the effectiveness of a classification-based system for subacute and chronic low back pain. Spine. 2012; 37(16):1347-1356
- 5 Berg S. On total disc replacement. Acta Orthopaedica. Sweden 2011; 82(Suppl.343):1-34
- Berg S, Tullberg T, Branth B, Olerud C, Tropp H. Total disc replacement compared to lumbar fusion: A randomised controlled trial with 2-year follow-up. European Spine Journal. 2009; 18(10):1512-1519
- 7 Chuang LH. Mapping the roland morris disability questionnaire for back pain into utility index. Value in Health. 2009; 12(7):A383
- 8 Chuang LH, Soares MO, Tilbrook H, Cox H, Hewitt CE, Aplin J et al. A pragmatic multicentered randomized controlled trial of yoga for chronic low back pain: economic evaluation. Spine. 2012; 37(18):1593-1601
- 9 Critchley DJ, Ratcliffe J, Noonan S, Jones RH, Hurley M, V. Effectiveness and cost-effectiveness of three types of physiotherapy used to reduce chronic low back pain disability: a pragmatic randomized trial with economic evaluation. Spine. 2007; 32(14):1474-1481
- Fairbank J, Frost H, Wilson-MacDonald J, Yu LM, Barker K, Collins R. Randomised controlled trial to compare surgical stabilisation of the lumbar spine with an intensive rehabilitation programme for patients with chronic low back pain: the MRC spine stabilisation trial. BMJ. 2005; 330(7502):1233
- Foster L, Clapp L, Erickson M, Jabbari B. Botulinum toxin A and chronic low back pain: a randomized, double-blind study. Neurology. 2001; 56(10):1290-1293
- Foster NE, Mullis R, Hill JC, Lewis M, Whitehurst DGT, Doyle C et al. Effect of stratified care for low back pain in family practice (IMPaCT Back): a prospective population-based sequential comparison. Annals of Family Medicine. 2014; 12(2):102-111
- Fritzell P, Berg S, Borgstrom F, Tullberg T, Tropp H. Cost effectiveness of disc prosthesis versus lumbar fusion in patients with chronic low back pain: randomized controlled trial with 2-year follow-up. European Spine Journal. 2011; 20(7):1001-1011

- Gilbert FJ, Grant AM, Gillan MG, Vale LD, Campbell MK, Scott NW et al. Low back pain: influence of early MR imaging or CT on treatment and outcome. Multicenter randomized trial. Radiology. 2004; 231(2):343-351
- Gilbert FJ, Grant AM, Gillan MGC, Vale L, Scott NW, Campbell MK. Does early magnetic resonance imaging influence management or improve outcome in patients referred to secondary care with low back pain? A pragmatic randomised controlled trial. Health Technology Assessment. England 2004; 8(17)
- Haas M, Vavrek D, Peterson D, Polissar N, Neradilek MB. Dose-response and efficacy of spinal manipulation for care of chronic low back pain: a randomized controlled trial. Spine Journal. 2014; 14(7):1106-1116
- Hellum C, Berg L, Gjertsen O, Johnsen LG, Neckelmann G, Storheim K et al. Adjacent level degeneration and facet arthropathy after disc prosthesis surgery or rehabilitation in patients with chronic low back pain and degenerative disc: second report of a randomized study. Spine. 2012; 37(25):2063-2073
- Hellum C, Johnsen LG, Gjertsen O, Berg L, Neckelmann G, Grundnes O et al. Predictors of outcome after surgery with disc prosthesis and rehabilitation in patients with chronic low back pain and degenerative disc: 2-year follow-up. European Spine Journal. 2012; 21(4):681-690
- Hellum C, Johnsen LG, Storheim K, Nygaard OP, Brox JI, Rossvoll I et al. Surgery with disc prosthesis versus rehabilitation in patients with low back pain and degenerative disc: two year follow-up of randomised study. BMJ. 2011; 342:d2786
- Hill JC, Dunn KM, Lewis M, Mullis R, Main CJ, Foster NE et al. A primary care back pain screening tool: identifying patient subgroups for initial treatment. Arthritis and Rheumatism. 2008; 59(5):632-641
- 21 Hill JC, Whitehurst DG, Lewis M, Bryan S, Dunn KM, Foster NE et al. Comparison of stratified primary care management for low back pain with current best practice (STarT Back): a randomised controlled trial. Lancet. 2011; 378(9802):1560-1571
- Hlobil H, Uegaki K, Staal JB, Bruyne M, Smid T, Mechelen W. Substantial sick-leave costs savings due to a graded activity intervention for workers with non-specific sub-acute low back pain. European Spine Journal. 2007; 16(7):919-924
- Hollinghurst S, Sharp D, Ballard K, Barnett J, Beattie A, Evans M et al. Randomised controlled trial of Alexander technique lessons, exercise, and massage (ATEAM) for chronic and recurrent back pain: economic evaluation. BMJ. 2008; 337:a2656
- Jellema P, van der Roer N, Van Der Windt DAWM, van Tulder MW, Van Der Horst HE, Stalman WAB et al. Low back pain in general practice: Cost-effectiveness of a minimal psychosocial intervention versus usual care. European Spine Journal. 2007; 16(11):1812-1821
- Jellema P, van der Windt DA, Van Der Horst HE, Twisk JW, Stalman WA, Bouter LM. Should treatment of (sub)acute low back pain be aimed at psychosocial prognostic factors? Cluster randomised clinical trial in general practice. BMJ. 2005; 331(7508):84
- Jellema P, van Tulder MW, Van Poppel MN, Nachemson AL, Bouter LM. Lumbar supports for prevention and treatment of low back pain: a systematic review within the framework of the cochrane back review group. Spine. 2001; 26(4):377-386

- Johnsen LG, Brinckmann P, Hellum C, Rossvoll I, Leivseth G. Segmental mobility, disc height and patient-reported outcomes after surgery for degenerative disc disease: a prospective randomised trial comparing disc replacement and multidisciplinary rehabilitation. Bone and Joint Journal. 2013; 95-B(1):81-89
- Johnsen LG, Hellum C, Storheim K, Nygaard OP, Brox JI, Rossvoll I et al. Cost-effectiveness of total disc replacement versus multidisciplinary rehabilitation in patients with chronic low back pain: A norwegian multicenter RCT. Spine. 2014; 39(1):23-32
- 29 Lamb SE, Hansen Z, Lall R, Castelnuovo E, Withers EJ, Nichols V et al. Group cognitive behavioural treatment for low-back pain in primary care: a randomised controlled trial and cost-effectiveness analysis. Lancet. United Kingdom 2010; 375(9718):916-923
- Lamb SE, Lall R, Hansen Z, Castelnuovo E, Withers EJ, Nichols V et al. A multicentred randomised controlled trial of a primary care-based cognitive behavioural programme for low back pain. the back skills training (BeST) trial. Health Technology Assessment. 2010; 14(41)
- Lamb SE, Lall R, Hansen Z, Withers EJ, Griffiths FE, Szczepura A et al. Design considerations in a clinical trial of a cognitive behavioural intervention for the management of low back pain in primary care: Back Skills Training Trial. BMC Musculoskeletal Disorders. 2007; 8:14
- Lamb SE, Mistry D, Lall R, Hansen Z, Evans D, Withers EJ et al. Group cognitive behavioural interventions for low back pain in primary care: extended follow-up of the Back Skills Training Trial (ISRCTN54717854). Pain. 2012; 153(2):494-501
- Lambeek LC, Bosmans JE, van Royen BJ, van Tulder MW, van MW, Anema JR. Effect of integrated care for sick listed patients with chronic low back pain: economic evaluation alongside a randomised controlled trial. BMJ. 2010; 341:c6414
- 34 Lambeek LC, van Mechelen W, Knol DL, Loisel P, Anema JR. Randomised controlled trial of integrated care to reduce disability from chronic low back pain in working and private life. BMJ. 2010; 340:c1035
- Little P, Roberts L, Blowers H, Garwood J, Cantrell T, Langridge J et al. Should we give detailed advice and information booklets to patients with back pain? A randomized controlled factorial trial of a self-management booklet and doctor advice to take exercise for back pain. Spine. 2001; 26(19):2065-2072
- Little P, Lewith G, Webley F, Evans M, Beattie A, Middleton K et al. Randomised controlled trial of Alexander technique lessons, exercise, and massage (ATEAM) for chronic and recurrent back pain. BMJ. 2008; 337:a884
- Lloyd A, Scott DA, Akehurst RL, Lurie-Luke E, Jessen G. Cost-effectiveness of low-level heat wrap therapy for low back pain. Value in Health. 2004; 7(4):413-422
- 38 Morera-Dominguez C, Ceberio-Balda F, Florez G, Masramon X, Lopez-Gomez V. A costconsequence analysis of pregabalin versus usual care in the symptomatic treatment of refractory low back pain: sub-analysis of observational trial data from orthopaedic surgery and rehabilitation clinics. Clinical Drug Investigation. 2010; 30(8):517-531
- Nadler SF, Steiner DJ, Erasala GN, Hengehold DA, Hinkle RT, Beth Goodale M et al. Continuous low-level heat wrap therapy provides more efficacy than Ibuprofen and acetaminophen for acute low back pain. Spine. 2002; 27(10):1012-1017

- 40 Niemisto L, Rissanen P, Sarna S, Lahtinen-Suopanki T, Lindgren K-A, Hurri H. Cost-effectiveness of combined manipulation, stabilizing exercises, and physician consultation compared to physician consultation alone for chronic low back pain: a prospective randomized trial with 2-year follow-up. Spine. 2005; 30(10):1109-1115
- 41 Niemisto L, Lahtinen-Suopanki T, Rissanen P, Lindgren KA, Sarna S, Hurri H. A randomized trial of combined manipulation, stabilizing exercises, and physician consultation compared to physician consultation alone for chronic low back pain. Spine. 2003; 28(19):2185-2191
- Organisation for Economic Co-operation and Development (OECD). Purchasing power parities (PPP). 2012. Available from: http://www.oecd.org/std/ppp [Last accessed: 1 November 2015]
- Peul WC, van den Hout WB, Brand R, Thomeer RTWM, Koes BW, Leiden-The Hague Spine Intervention Prognostic Study Group. Prolonged conservative care versus early surgery in patients with sciatica caused by lumbar disc herniation: two year results of a randomised controlled trial. BMJ. 2008; 336(7657):1355-1358
- Price C, Arden N, Coglan L, Rogers P. Cost-effectiveness and safety of epidural steroids in the management of sciatica. Health Technology Assessment. 2005; 9(33)
- Ratcliffe J, Thomas KJ, MacPherson H, Brazier J. A randomised controlled trial of acupuncture care for persistent low back pain: cost effectiveness analysis. BMJ. 2006; 333:626-628
- 46 Rivero-Arias O, Campbell H, Gray A, Fairbank J, Frost H, Wilson-MacDonald J. Surgical stabilisation of the spine compared with a programme of intensive rehabilitation for the management of patients with chronic low back pain: cost utility analysis based on a randomised controlled trial. BMJ. 2005; 330:1239-1243
- 47 Smeets RJEM. Do lumbar stabilising exercises reduce pain and disability in patients with recurrent low back pain? Australian Journal of Physiotherapy. 2009; 55(2):138
- Smeets RJEM, Vlaeyen JWS, Hidding A, Kester ADM, van der Heijden GJMG, Knottnerus JA. Chronic low back pain: physical training, graded activity with problem solving training, or both? The one-year post-treatment results of a randomized controlled trial. Spine. 2008; 134(3):263-276
- 49 Smeets RJEM, Vlaeyen JWS, Hidding A, Kester ADM, van der Heijden GJMG, van Geel ACM et al. Active rehabilitation for chronic low back pain: cognitive-behavioral, physical, or both? First direct post-treatment results from a randomized controlled trial [ISRCTN22714229]. BMC Musculoskeletal Disorders. 2006; 7:5
- Spijker-Huiges A, Vermeulen K, Winters JC, van WM, van der Meer K. Costs and costeffectiveness of epidural steroids for acute lumbosacral radicular syndrome in general practice: an economic evaluation alongside a pragmatic randomized control trial. Spine. 2014; 39(24):2007-2012
- 51 Staal JB, Hlobil H, Twisk JWR, Smid T, Koke AJA, van Mechelen W. Graded activity for low back pain in occupational health care: a randomized, controlled trial. Annals of Internal Medicine. 2004; 140(2):77-84
- Steenstra IA, Anema JR, van Tulder MW, Bongers PM, de Vet HC, van MW. Economic evaluation of a multi-stage return to work program for workers on sick-leave due to low back pain. Journal of Occupational Rehabilitation. 2006; 16(4):557-578

- 53 Steenstra IA, Anema JR, Bongers PM, de Vet HCW, van Mechelen W. Cost effectiveness of a multi-stage return to work program for workers on sick leave due to low back pain, design of a population based controlled trial [ISRCTN60233560]. BMC Musculoskeletal Disorders. 2003; 4:26
- Thomas KJ, MacPherson H, Ratcliffe J, Thorpe L, Brazier J, Campbell Mea. Longer term clinical and economic benefits of offering acupuncture care to patients with chronic low back pain. Health Technology Assessment. 2005; 9(32)
- Thomas KJ, MacPherson H, Thorpe L, Brazier J, Fitter M, Campbell MJ et al. Randomised controlled trial of a short course of traditional acupuncture compared with usual care for persistent non-specific low back pain. BMJ. 2006; 333(7569):623
- Thomas M, Lundberg T. Importance of modes of acupuncture in the treatment of chronic nociceptive low back pain. Acta Anaesthesiologica Scandinavica. 1994; 38(1):63-69
- 57 Tilbrook HE, Cox H, Hewitt CE, Kang'ombe AR, Chuang LH, Jayakody S et al. Yoga for chronic low back pain. Annals of Internal Medicine. 2011; 155(9):569-578
- Tosteson AN, Lurie JD, Tosteson TD, Skinner JS, Herkowitz H, Albert T et al. Surgical treatment of spinal stenosis with and without degenerative spondylolisthesis: cost-effectiveness after 2 years. Annals of Internal Medicine. 2008; 149(12):845-853
- Tosteson ANA, Skinner JS, Tosteson TD, Lurie JD, Andersson GB, Berven S et al. The cost effectiveness of surgical versus nonoperative treatment for lumbar disc herniation over two years: Evidence from the Spine Patient Outcomes Research Trial (SPORT). Spine. 2008; 33(19):2108-2115
- 60 UK BEAM Trial Team. United Kingdom back pain exercise and manipulation (UK BEAM) randomised trial: Cost effectiveness of physical treatments for back pain in primary care. BMJ. 2004; 329(7479):1381-1385
- 61 Underwood M. United Kingdom back pain exercise and manipulation (UK BEAM) randomised trial: Effectiveness of physical treatments for back pain in primary care. BMJ. 2004; 329(7479):1377-1381
- van den Hout WB, Peul WC, Koes BW, Brand R, Kievit J, Thomeer RT. Prolonged conservative care versus early surgery in patients with sciatica from lumbar disc herniation: cost utility analysis alongside a randomised controlled trial. BMJ. Netherlands 2008; 336(7657):1351-1354
- van Wijk RMAW, Geurts JWM, Wynne HJ, Hammink E, Buskens E, Lousberg R et al.
 Radiofrequency denervation of lumbar facet joints in the treatment of chronic low back pain: a randomized, double-blind, sham lesion-controlled trial. Clinical Journal of Pain. 2005; 21(4):335-344
- Vavrek D, Sharma R, Haas M. Cost-analysis related to dose-response for spinal manipulative therapy for chronic low back pain: outcomes from a randomized controlled trial. Journal of Alternative and Complementary Medicine. 2014; 20(5):A18
- Whitehurst DG, Bryan S, Lewis M, Hay EM, Mullis R, Foster NE. Implementing stratified primary care management for low back pain: cost-utility analysis alongside a prospective, population-based, sequential comparison study. Spine. 2015; 40(6):405-414

- Whitehurst DGT, Bryan S, Lewis M, Hill J, Hay EM. Exploring the cost-utility of stratified primary care management for low back pain compared with current best practice within risk-defined subgroups. Annals of Rheumatic Diseases. 2012; 71(11):1796-1802
- Whitehurst DGT, Lewis M, Yao GL, Bryan S, Raftery JP, Mullis R et al. A brief pain management program compared with physical therapy for low back pain: results from an economic analysis alongside a randomized clinical trial. Arthritis and Rheumatism. 2007; 57(3):466-473
- Wielage RC, Bansal M, Andrews JS, Wohlreich MM, Klein RW, Happich M. The costeffectiveness of duloxetine in chronic low back pain: a US private payer perspective. Value in Health. 2013; 16(2):334-344