REVIEW



Efficacy of Percutaneous Adhesiolysis in Managing Low Back and Lower Extremity Pain: A Systematic Review and Meta-analysis of Randomized Controlled Trials

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ABSTRACT

Introduction: Chronic refractory low back and lower extremity pain recalcitrant to conservative management and epidural injections secondary to postsurgery syndrome, spinal stenosis, and disc herniation are sometimes managed with percutaneous adhesiolysis. Consequently, this systematic review and metanalysis was undertaken to assess the efficacy of percutaneous adhesiolysis in managing low back and lower extremity pain.

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L. Manchikanti (⊠) · M. R. Sanapati Pain Management Centers of America, 67 Lakeview Drive, Paducah, KY 42001, USA e-mail: drlm@thepainmd.com

N. N. Knezevic Advocate Illinois Masonic Medical Center and College of Medicine, University of Illinois, Chicago, IL, USA e-mail: nick.knezevic@gmail.com

E. Knezevic

College of Liberal Arts and Sciences, University of Illinois at Urbana-Champaign, Champaign, IL, USA e-mail: ekneze2@illinois.edu

R. Pasupuleti University of Kentucky, Lexington, KY, USA e-mail: rpa286@g.uky.edu Methods: A systematic review and meta-analysis of randomized controlled trials (RCTs) utilizing the Preferred Reporting Items Systematic Reviews and Meta-Analyses (PRISMA) checklist was performed. A comprehensive literature search of multiple databases from 1966 to July 2022, including manual searches of the bibliography of known review articles was performed. Quality assessment of the included trials, meta-analysis, and best evidence synthesis was performed. The primary outcome measure was a significant reduction in pain (short term up to 6 months and long term more than 6 months).

Results: The search identified 26 publications, with 9 trials meeting the inclusion criteria. The results of dual-arm and single-arm analyses

A. D. Kaye

LSU Health Sciences Center, Shreveport, Ochsner Shreveport Hospital and Interventional Pain Clinic Feist-Wieller Cancer Center, Shreveport, LA, USA

L. Manchikanti · M. R. Sanapati Pain Management Centers of America, Evansville, IN, USA

M. R. Sanapati e-mail: msanapati@gmail.com

J. A. Hirsch Massachusetts General Hospital and Harvard Medical School, Boston, MA, USA e-mail: jahirsch@mgh.harvard.edu showed significant improvement in pain and function at 12 months. Opioid consumption was also significantly reduced at 6 months with dual-arm analysis, whereas single-arm analysis showed a significant decrease from baseline to treatment at the 3-, 6-, and 12-month analyses. At 1 year follow-up, seven of seven trials were positive for improvements in pain relief, function, and diminution of opioid use.

Discussion: Based on the present systematic review of nine RCTs, the evidence level is I to II, with moderate to strong recommendation for percutaneous adhesiolysis in managing low back and lower extremity pain. The limitations of the evidence include paucity of literature, lack of placebo-controlled trials, and the majority of the trials studying post lumbar surgery syndrome.

Conclusion: The evidence is level I to II or strong to moderate based on five high-quality and two moderate-quality RCTs, with 1 year follow-up that percutaneous adhesiolysis is efficacious in the treatment of chronic refractory low back and lower extremity pain.

Keywords: Chronic low back pain; Epidural scarring; Lumbar disc herniation; Lumbar spinal stenosis; Percutaneous adhesiolysis; Post lumbar surgery syndrome; Radicular pain; Spinal pain

Key Summary Points

- 1. Chronic refractory low back and lower extremity pain secondary to post lumbar surgery syndrome, spinal stenosis, and disc herniation is common.
- 2. Disc herniation, spinal stenosis, and postsurgery syndrome is managed with multiple interventional techniques, implantable therapies, or repeat surgical interventions.

- 3. The present systematic review identified seven high-quality and two moderate-quality randomized controlled trials (RCTs) evaluating the role of percutaneous adhesiolysis in managing chronic recalcitrant low back and lower extremity pain.
- 4. The evidence is level I to II with moderate to strong recommendation in managing low back and lower extremity pain after failure of conservative management and fluoroscopically directed epidural injections.
- 5. Significant paucity of the literature and heterogeneity among available trials continues to be an issue, resulting in an ongoing debate regarding efficacy, effectiveness, indications, and medical necessity.

INTRODUCTION

Chronic refractory low back pain with or without lower extremity pain that does not resolve after conservative therapy or even surgical treatment can present a therapeutic dilemma with limited options for proper management [1–8]. Low back and lower extremity pain recalcitrant to conservative management and epidural injections may be secondary to postsurgery syndrome, spinal stenosis, and disc herniation [1-8]. Disc herniation and spinal stenosis are often managed with surgical interventions and postsurgery syndrome may also be managed with repeat surgical interventions or implantable therapies. However, for those patients who are not responsive to or candidates for surgical interventions and/or have not adequately responded to epidural injections, percutaneous adhesiolysis may be an option [1-12]. Percutaneous adhesiolysis is also considered an option in patients not amenable to or having an inadequate response to neuromodulation therapies [1, 7–16]. Changing coverage policies have impacted utilization patterns of interventional techniques in general and percutaneous adhesiolysis in particular [1, 9–16].

National health care expenditures are an important issue, specifically since the COVID-19 pandemic, which has drastically altered health care delivery and modes of treatment [17–33]. The COVID-19 pandemic and the opioid epidemic have negatively impacted access to treatment and costs in chronic pain sufferers [17–33]. The analysis of national health care spending in the USA showed an increase of 9.7% to reach \$4.1 trillion in 2020, compared with a 4.3% increase seen in 2019 [17, 18]. The acceleration in 2020 was related to a 36% increase in federal expenditures for health care that occurred largely in response to the COVID-19 pandemic. Multiple other factors, including consolidation of providers into an employment model by health systems, which has increased substantially, has been contributing to increasing health care expenses [17, 18, 23-36]. Multiple effects due to COVID-19, with increased psychological stress and suffering, may also have a significant effect on outcomes [27, 30–32]. An analysis of the utilization patterns in the fee-for-service (FFS) Medicare population, including the impact of COVID-19, showed declining interventional techniques with an overall decrease of interventional techniques at an annual rate of 0.4% per 100,000 Medicare population from 2010 to 2019, and a decrease of 24.5% for epidural injections and adhesiolysis procedures [19]. However, the decrease from 2019 to 2020 due to the COVID-19 pandemic was 18.7% for all interventions compared with 19.0% for epidural and adhesiolysis procedures [19]. Additionally, epidural-specific utilization patterns [22] showed an overall decrease of utilization of epidural injections of 24.1% annually from 2010 to 2019, with a significant effect of the COVID-19 pandemic showing a 19.0% decrease from 2019 to 2020 [22]. Further, compared with multiple other interventions, including epidural injections, facet joint interventions, and sacroiliac joint interventions, augmentation procedures [10, 11, 14-16] and percutaneous adhesiolysis [12] have faced a substantial decline at a rapid rate. There is discordance of opinions on the efficacy and effectiveness of medical necessity and indications among various authorities [1–8].

Helm et al. [6] published a systematic review of percutaneous adhesiolysis in 2016 utilizing seven randomized controlled trials (RCTs) and three observational studies, concluding with level I or strong evidence of the efficacy of percutaneous adhesiolysis in the treatment of chronic refractory low back and lower extremity pain. In subsequent reports, Cho et al. [7] and Manchikanti et al. [2-4] have shown level II-I evidence for post lumbar surgery syndrome, spinal stenosis, and disc herniation. In fact, Cho et al. [7] have shown significant evidence for both percutaneous adhesiolysis and spinal cord stimulation (SCS) with a recommendation of level A for epidural adhesiolysis for 6--12 months of pain relief and functional improvement and level B for SCS.

In contrast, Brito-García et al. [8] in a systematic review without meta-analysis provided a rather poor methodological quality assessment of the rating of the trials, with downgrading to low quality, which have been rated as high quality in multiple other evaluations. Overall, they concluded that there was no evidence for percutaneous adhesiolysis. Thus, of the five systematic reviews, three of them, including a meta-analysis and one systematic review without meta-analysis, showed positive results compared with only one systematic review that, although it looked at similar studies, concluded very differently. Manchikanti et al. [5], in assessing systematic reviews with a systematic analysis, identified multiple issues in one of the systematic reviews. Further, Manchikanti et al. [1, 9, 37-40] and others [41, 42] have also described extensively the issues related to performance of evidence synthesis in systematic reviews and meta-analysis.

Consequently, to assess the efficacy of percutaneous adhesiolysis in managing low back and lower extremity pain this systematic review and meta-analysis of RCTs was undertaken..

METHODS

A systematic review and meta-analysis were performed based on the methodological and reporting quality of systematic reviews, as described by the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) [43]. Methodology from other reviews was also utilized [2–5, 37–39, 44].

Eligibility Criteria

Randomized trials of interest included patients suffering from chronic low back and lower extremity pain due to postsurgery syndrome, spinal stenosis, and disc herniation and treated with percutaneous epidural neurolysis or adhesiolysis. Trials of patients with fractures, malignancies, acute trauma, and inflammatory diseases were excluded. All RCTs were included.

This review focused on lumbar percutaneous adhesiolysis/neurolysis for postsurgery syndrome, central spinal stenosis, and disc herniation with multiple approaches. All trials that provided appropriate outcome data and analysis for 6 months were reviewed. Book chapters, case reports, and reports without an appropriate diagnosis were not considered.

Information Sources

All available studies in the English language, or with available translation, with appropriate reporting of outcomes data for 6 months were included. Searches were performed using multiple databases, including PubMed, www.ncbi.nlm.nih.gov/pubmed; Cochrane Library, www.thecochranelibrary.com; US National Guideline Clearinghouse (NGC), www.guideline.gov/; clinical trials, www.clinicaltrials.gov/; and Google Scholar, https://scholar.google.com; from 1966 to July 2022 [4].

Search Strategy

The search terminology was as follows:

(chronic low back pain OR nerve root compression OR lumbosciatic pain OR radicular pain OR radiculitis OR sciatica OR disc herniation, postsurgery syndrome, failed surgery syndrome, spinal stenosis) AND (epidural injection OR epidural adhesiolysis OR neurolysis OR epidural neuroplasty OR epidural lysis of

adhesions OR percutaneous adhesiolysis OR neurolysis OR transforaminal injection OR corticosteroid OR methylprednisolone OR bupivacaine OR lidocaine) AND (meta-analysis [pt] OR randomized controlled trial [pt] OR controlled clinical trial [pt] OR systematic review OR randomized controlled trials [mh] OR nonrandomized studies OR observational studies OR random allocation [mh] OR double-blind method [mh] OR single-blind method [mh] OR clinical trial [pt] OR clinical trials [mh] OR ("clinical trial" [tw]) OR ((singl* [tw] OR doubl* [tw] OR trebl* [tw] OR tripl* [tw]) AND (mask* [tw] OR blind* [tw])) OR (placebos [mh] OR placebo* [tw] OR random* [tw] OR research design [mh:noexp]).

Data Selection

In the identification of the relevant literature, the article selection and extraction of the data from the included studies was conducted independently, by two review authors (N.N.K. and M.R.S.). Any disagreement among the reviewer authors were resolved by the third author (A.D.K.). All conflicts of interest of the reviewers with authorship of the article was resolved by assigning them to other reviewers.

Study Risk of Bias Assessment

Two authors completed the quality assessment of each individual article. Three authors completed evidence synthesis. All conflicts were resolved as stated above by a fourth author.

The quality of each RCT was assessed using the Cochrane Review rating system (see Table S1 in the electronic supplementary material for details) [45] and the Interventional Pain Management Techniques—Quality Appraisal of Reliability and Risk of Bias Assessment Tool (IPM–QRB) for RCTs (aee Table S2 in the electronic supplementary material for details) [46].

Randomized trials meeting at least 9 of the 13 inclusion criteria of the Cochrane Review were considered high quality. The trials meeting 5–8 criteria were considered moderate quality,

and those meeting fewer than 5 criteria were considered low quality and were excluded.

Based on the IPM–QRB criteria, randomized trials with scores of 32–48 were considered high quality, studies scoring 16–31 were considered moderate quality, and studies scoring less than 16 were considered low quality and were excluded.

Analysis of the Evidence

Analysis of the evidence was performed by two authors, N.N.K. and E.K., with consultation from A.D.K., M.R.S., and J.A.H. Any disagreements among the authors was resolved by consensus or by A.D.K. and J.A.H.

Outcome of the Studies

Clinically important outcome measures were 50% significant improvement from the baseline pain score or a change of at least 3 points on an 11-point pain scale of 0 to 10 and a change of 30% or more on disability scores [4].

Based on the relevance and effectiveness of the adhesiolysis, either compared with a control group or from baseline to follow-up, a trial was categorized as positive or negative neutral. Reference point measurements were considered at 3 months, 6 months, and 1 year [4].

The best-evidence synthesis developed by American Society of Interventional Pain Physicians (ASIPP), modified, and collated using multiple criteria, was used for qualitative analysis (Table 1) [47]. The evidence synthesis varied from strong to opinion- or consensus-based using five levels of evidence.

Table 2 presents guidance for the strength of recommendations from weak to strong [48].

The results of best evidence as per grading were utilized and the Grading of Recommendations, Assessment, Development and Evaluations (GRADE) system of appraisal was used for determining the body of evidence [49]. Clinical relevance and pragmatism of all studies were assessed [50].

Table 1 Qualitative modified approach to grading of evidence. Adapted from Manchikanti et al. [47]

Level I	Strong	Evidence obtained from multiple relevant high-quality randomized controlled trials
Level II	Moderate	Evidence obtained from at least one relevant high-quality randomized controlled trial or multiple relevant moderate- or low-quality randomized controlled trials
Level III	Fair	Evidence obtained from at least one relevant moderate- or low-quality randomized trial
		or
		Evidence obtained from at least one relevant high-quality non-randomized trial or observational study with multiple moderate- or low-quality observational studies
Level IV	Limited	Evidence obtained from multiple moderate- or low-quality relevant observational studies
Level V	Consensus based	Opinion or consensus of a large group of clinicians and/or scientists

Meta-analysis

Dual-Arm Meta-analysis

For the dual-arm meta-analysis, Review Manager version 5.4 (The Cochrane Collaboration) 2020, software was used. For pain and functionality improvement data, the studies were reported as the standardized mean differences (SMD) with 95% confidence intervals (CI). Data were plotted using forest plots to evaluate treatment effects using a random effects model. Heterogeneity was interpreted through I^2 statistics [40].

Single-Arm Meta-analysis

For the single-arm meta-analysis, Comprehensive Meta-Analysis version 3.0 (Biostat Inc.,

Table 2 Guide for strength of recommendations Source: National Guideline Clearinghouse Extent Adherence to Trustworthy Standards (NEATS) instrument [48]

Strong There is high confidence that the recommendation reflects best practice. This is based on: (a) strong evidence for a true net effect (e.g., benefits exceed harms);
(b) consistent results, with no or minor exceptions; (c) minor or no concerns about

study quality; and/or (d) the extent the panelists' agreement. Other compelling considerations (discussed in the guideline's literature review and analyses) may also warrant a strong recommendation

Moderate There is moderate confidence that the recommendation reflects best practice. This is based on: (a) good evidence for a true net effect (e.g., benefits exceed harms);
(b) consistent results, with minor and/or few exceptions; (c) minor and/or few concerns about study quality; and/or (d) the extent of panelists' agreement. Other compelling considerations (discussed in the guideline's literature review and analyses) may also warrant a moderate recommendation

Weak There is some confidence that the recommendation offers the best current guidance for practice. This is based on:

(a) limited evidence for a true net effect (e.g., benefits exceed harms); (b) consistent results, but with important exceptions; (c) concerns about study quality; and/or (d) the extent of panelists' agreement. Other considerations (discussed in the guideline's literature review and analyses) may also warrant a weak recommendation

Englewood, NJ) software was used. For pain and functionality improvement data, the studies were reported as the mean differences with 95% confidence intervals. Data were plotted using forest plots to evaluate treatment effects.

Heterogeneity was interpreted through I^2 statistics [40].

Compliance with Ethics Guidelines

This article is based on previously conducted studies and does not contain any new studies with human participants or animals performed by any of the authors.

RESULTS

Study Selection

Figure 1 shows a flow diagram of the study selection using the PRISMA study selection process.

Based on the search criteria, 26 publications were identified and considered for inclusion [51–76]. A total of 12 trials [51–59, 61, 62, 76] met the inclusion criteria and 9 trials were included after exclusion of duplicates, follow-up evaluations, and qualifications [51-54, 56, 58, 61, 62, 76]. Three trials reported follow-up results, consequently, these were not considered as separate studies [54-59]. Of the nine trials included, six of them studied postsurgery syndrome [51–54, 62, 76], two trials studied spinal stenosis [56, 61], and one trial studied disc herniation [58]. Of the included trials, only one trial was placebo controlled [58], and eight were active controlled trials [51–54, 56, 61, 62, 76].

Methodological Quality and Risk of Bias Assessment

Tables 3 and 4 present the methodological quality assessment and risk of bias of the nine RCTs utilizing the Cochrane review criteria and the IPM–QRB criteria, respectively [51–54, 56, 58, 61, 62, 76]. Assessment by the Cochrane review criteria showed all of them as high-quality trials, scoring at least 9 of 13. However, based on the IPM–QRB instrument, seven of the nine trials [52, 54, 56, 58, 61, 62, 76] scored as high, with scores of over 32 of 48. The

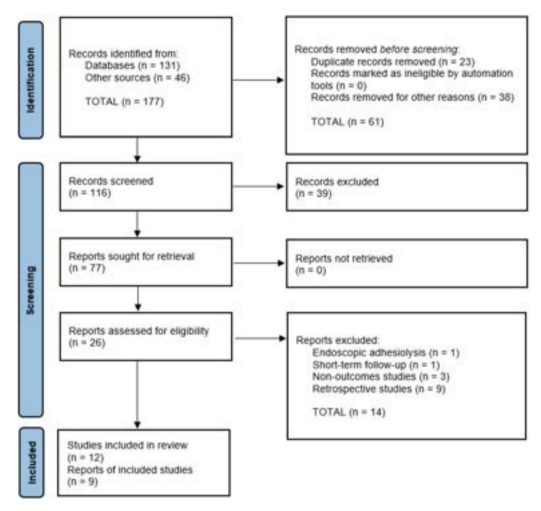


Fig. 1 Schematic presentation of the study selection of percutaneous adhesiolysis based on the PRISMA 2020 flow diagram for new systematic reviews

remaining two studies [51, 53] showed moderate quality, with scores above 16.

Study Characteristics

Table 5 presents the characteristics and outcomes of the studies meeting the inclusion criteria for receiving percutaneous adhesiolysis/neurolysis for lumbar disc herniation.

Results of Individual Studies

Qualitative Analysis

Qualitative analysis at 6-months follow-up, showed that one of the nine trials had negative

results [61]; however, at 1-year follow-up, seven trials showed positive results [51–54, 56, 58, 62].

Qualitative analysis was also performed, utilizing a modified approach of grading of evidence with moderate (level II) evidence from seven relevant high-quality RCTs [52, 54, 56, 58, 61, 62, 76] and two relevant moderate-quality RCTs [51, 53].

Utilizing the GRADE criteria [49], there was no change in the evidence level. All the trials were considered to meet clinical relevance and pragmatism [50]. All the included trials met pragmatic criteria for clinical relevance and pragmatism [50]. In addition, the evidence was assessed by qualitative and quantitative evidence synthesis utilizing conventional dual-arm

	Heavner et al. [51]	Manchikanti et al. [52]	Veihelmann et al. [53]	Manchikanti et al. [54, 55]	Manchikanti et al. [56, 57]	Chunjing et al. [76]	Gerdesmeyer et al. [58]	Karm et al. [61]	Akbas et al. [62]
Randomization adequate	U	Y	Y	Y	Y	Y	Y	Y	Y
Concealed treatment allocation	Ω	Y	Y	Y	Y	Y	Y	Y	Y
Patient blinded	Y	Y	Z	Y	Y	Y	Y	Y	Z
Care provider blinded	Z	Z	Z	Z	Z	z	Y	Y	Z
Outcome assessor blinded	10	Y	Z	Ω	Z	Y	Y	Y	Z
Dropout rate described	Y	Y	Y	Y	Y	Y	Y	Z	NA
All randomized participants analyzed in the group	Y	Y	¥	¥	Y	Y	Y	Z	\succ
Reports of the study free of suggestion of selective outcome reporting	¥	¥	¥	X	Y	X	¥	>	×
Groups similar at baseline regarding most important prognostic indicators	>	Y	Y	X	Y	X	¥	×	> -
Co-intervention avoided or similar in all groups	¥	¥	×	¥	¥	×	¥	X	X
Compliance acceptable in all groups	¥	¥	¥	¥	¥	¥	¥	X	>
Time of outcome assessment in all groups similar	¥	Y	¥	¥	¥	¥	¥	¥	¥
Are other sources of potential bias not likely	¥	Y	>	¥	¥	×	Y	X	¥
Score	10/13	12/13	10/13	11/13	11/13	12/13	13/13	11/13	9/13

Y yes, N no, U unclear

 Table 4
 Methodological quality assessment of randomized trials of percutaneous adhesiolysis utilizing the IPM-QRB criteria. Source: Manchikanti et al. [48]

Trial design and guidance reporting Trial design and guidance Feprotring			Heavner et al. [51]	Manchikanti et al. [52]	Veihelmann et al. [53]	Manchikanti et al. [54, 55]	Manchikanti et al. [56, 57]	Chunjing et al. [76]	Gerdesmeyer et al. [58]	Karm et al. [61]	Akbas et al. [62]
CoNSORT or SPIRIT 0 3 2 0 3 2 Design factors Type and design of trial 2 2 2 3 2 3 2 3 2 3 2 3 3 2 3 3 2 2 3 2 2 3 2 2 3		Trial design and guidance									
Design factors Type and design of trial Scruting/physician Limaging Sample size Sample size	_	CONSORT or SPIRIT	0	3	1	3	2	0	8	2	2
Type and design of trial 2 3 2 2 3 2 3 2 3 2 3 2 3 3 2 3 <td>Η</td> <td>Design factors</td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td>	Η	Design factors									
Setting/physician 2 2 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 3	7	Type and design of trial	2	2	3	2	2	2	3	2	2
Imaging 3 </td <td>3</td> <td>Setting/physician</td> <td>2</td> <td>2</td> <td>1</td> <td>2</td> <td>2</td> <td>1</td> <td>2</td> <td>2</td> <td>2</td>	3	Setting/physician	2	2	1	2	2	1	2	2	2
Sample size 0 2 2 3 2 2 2 1 Patient factors 1 1 1 1 1 1 1 1 Inclusiveness of population 1 2 <	4	Imaging	3	3	3	3	3	3	3	3	8
Patient factors 1 2	\sim	Sample size	0	2	2	3	2	2	2	_	2
Patient factors Inclusiveness of population 1 2	9	Statistical methodology	1	1	1	1	1	1	1	_	1
Inclusiveness of population 1 1 2<	II	Patient factors									
Duration of pain 2	_	Inclusiveness of population	1	1	2	2	2	2	2	2	2
Previous treatments 2 3 2 3 4 3 4 3 4 3 4 3 4 4 4 4 4 4	∞	Duration of pain	2	2	0	2	2	1	2	2	2
Outcomes A malysis of all randomized participants in the groups 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 4 4 4 2	6	Previous treatments	2	2	2	2	2	2	2	2	7
Outcomes 1 Outcomes assessment criteria 0 4 4 2 4 2 for significant improvement 1 2 1 2 1 2 1 2 Analysis of all randomized 1 2 1 2 2 1 participants in the groups 1 0 1 1 1 1 1	10	Duration of follow-up with appropriate interventions	2	2	2	2	2	1	7	2	П
Outcomes assessment criteria 0 4 4 4 5 4 2 4 2 4 2 4 2 2 4 2 4 2 2 2 2 2 2 1 3 3 1 3 1 3 1	>	Outcomes									
Analysis of all randomized 1 2 1 2 2 1 participants in the groups 1 0 1 <	11	Outcomes assessment criteria for significant improvement	0	2	0	4	4	2	4	2	7
Description of dropout rate 0 1 1 1 1 1 1 1 1	12	Analysis of all randomized participants in the groups	1	2	_	2	1	2	7	1	7
	13	Description of dropout rate	0	1	0	1	1	1	1	Т	0

Lable	lable 4 continued									
		Heavner et al. [51]	Manchikanti et al. [52]	Veihelmann et al. [53]	Manchikanti et al. [54, 55]	Manchikanti et al. [56, 57]	Chunjing et al. [76]	Gerdesmeyer et al. [58]	Karm et al. [61]	Akbas et al. [62]
41	Similarity of groups at baseline for important prognostic indicators	7	7		7		2	7	1	2
15	Role of co-interventions	1	1	1	1	1	1	-	1	1
>	Randomization									
16	Method of randomization	0	2	1	2	2	2	2	2	2
M	Allocation of concealment									
17	Concealed treatment allocation	1	1	1	7	_	2	2	2	2
VII	Blinding									
18	Patient blinding	1	1	1	1	1	1	1	П	0
19	Care provider blinding	0	0	0	0	0	0	1	0	0
20	Outcome assessor blinding	0	1	0	0	0	1	1	0	0
VIII	VIII Conflicts of interest									
21	Funding and sponsorship	2	2	2	2	2	2	2	2	7
22	Conflicts of interest	0	3	1	3	2	3	3	2	3
Total		23/48	38/48	27/48	42/48	36/48	34/48	44/48	34/48	35/48

Table 5 Study characteristics of randomized trials assessing percutaneous adhesiolysis

Study	Number of	Interventions	Outcome	Pain relief and function	function		Results			Comments/conclusions
Study characteristic Methodological quality scoring	patients and selection criteria		measures	3 months	6 months	l year	3 months	3 months 6 months 1 year	l year	
Lumbar postsurgery syndrome Chun-jing et al. 92 patient (2012) [76] pain ar Randomized, 6 mon active control surgery Quality scores: herniat Cochrane = 12/ patient 13 evaluat 13	mbar postsurgery syndrome Chun-jing et al. 92 patients with (2012) [76] pain and andomized, formuchs after active control surgery for disc Quality scores: herniation; 76 cochrane = 12/ patients were 13 cvaluated AA-QRB = 34/	Catheter with guidewire was passed to ventral epidural space: 50–90 ml and 10 mg of dexamethasone were injected. Control received 10 mg dexamethasone only	VAS, opioid use; Macnab criteria	¥ Z	> 3 points relief on VAS, with 46% relief	N N	N.	<u>e</u>	V. ∀	Adhesiolysis is effective using the vascular catheter with ventral placement of the catheter. Improvements in dye flow are necessary for good clinical outcomes
Manchikanti et al. (2009) [54, 55]. Randomized, active control Quality scores: Cochrane = 11/ 13 IPM-QRB = 42/ 48	120 Post lumbar surgery syndrome	Adhesiolysis with 10% saline versus S3 caudal injection with 0.9% saline	NRS, ODI, employment status, opioid use	s8% of adhesiolysis had > 50% relief versus 38% of comparator	54% of adhesiolysis had > 50% relief versus 27% of comparator	s1% of adhesiolysis had > 50% relief versus 23% of comparator	ď	ď	ط	90% of adhesiolysis group had > 50% relief at 3 months and 73% did at 12 months; 35% of caudal group had > 50% relief at 3 months and 12% did at 12 months; 77% of adhesiolysis group had > 40% improvement in ODI at 12 months compared with 13% of the caudal group. Average of 3.5 adhesiolysis procedures/year with an average relief/year of 4.5 weeks

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-	9	
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Study	Number of patients and	Interventions	Outcome	Pain relief and function	nction		Results			Comments/conclusions
Study characteristic Methodological quality scoring	selection criteria		measures	3 months	6 months	l year	3 months	3 months 6 months 1 year	l year	
Heavner et al. (1999) [51] Randomized, active control Quality scores: Cochrane = 10/ 13 1PM-QRB = 23/ 48	83	3-day adhesiolysis in four groups with either 0.9% or 10% saline and with or without hyaluronidase	VAS, McGill pain questionnaire	About 50% of subjects had more than 10/100 improvement in VAS	About 50% of subjects had more than 10/100 improvement in VAS	About 50% of subjects had more than 10/100 improvement in VAS	<u>a</u>	<u>a</u>	۵	Moderate-quality study comparing four treatment options. Reduced additional procedures First study to compare various groups with positive results
Manchikanti et al. (2004) [52] Randomized, active control Quality scores: Cochrane = 12/ 13 13 1PM—QRB = 38/	Low back pain without response to epidural injection and no facet disease. Between 64% and 72% of patients had prior lumbar surgery; between 4% and 20% had spinal stenosis	One day adhesiolysis with 0.9% and 10% saline versus epidural injection	VAS, ODI, work status, opioid intake, range of motion measurement, and P-3 \geq 50% pain relief and functional status	72% of the 10% saline group. 64% of the 0.9% group and 0% of the caudal group had > 50% relief	saline group, 60% of the 0.9% group and 0% of the caudal group had > 50% relief	72% of the 10% saline group, 60% of the 0.9% group and 0% of the caudal had > 50% relief	<u>د</u>	۵.	٩	High-quality RCT showing that adhesiolysis provides significant relief regardless of whether normal saline or hypertonic saline is used

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Study	Number of patients	Interventions	Outcome	Pain relief and function	on		Results			Comments/conclusions
Study characteristic Methodological quality scoring	and selection criteria		measures	3 months	6 months	1 year	3 months	6 months	1 year	
Veihelmann et al. (2006) [53] Randomized, active control Quality scores: Cochrane = 10/ 13 IPM-QRB = 27/ 48	99 patients with radicular pain with concordant imaging findings	One day adhesiolysis with 10% saline versus physical therapy	VAS, ODI, GHS	Mean improvement of the treated group was > 50% in VAS and > 40% in ODI. Treatment group had ~ 10% relief	Mean improvement of the treated group was > 50% in VAS and > 40% in ODI. Treatment group had ~ 10% relief	Mean improvement of the treated group was > 50% in VAS and > 40% in ODI. Treatment group had ~ 10% relief	۵	۵.	<u>a</u>	Adhesiolysis is superior to physical therapy in treating persistent back and leg pain with concordant imaging findings
Akbas et al. (2018) [62] Randomized, active control Quality scores: Cochrane = 9/13 IPM-QRB = 35/ 48	60 patients Post lumbar surgery syndrome Three groups: Caudal = 20 SI foraminal = 20 L5 transforaminal = 20	All patients underwent placement of 16-gauge RX Coude needle in the Racz catheter with three approaches along with adhesiolysis. They also received exercises with neural flossing 3-4 times daily for 3 months With the caudal approach, L5 and S1 transforaminal approach, L5 and S1 transforaminal approach, local anesthetic, hyaluronidase, and steroids were administered	VAS, ODI	Significant improvement was seen in pain and functional status, with a reduction in scores with all three approaches, and with no significant differences between the approaches	Significant improvement was seen in pain and functional status, with a reduction in scores with all three approaches, and with no significant differences between the approaches	Significant improvement was seen in pain and functional sratus, with a reduction in scores with all three approaches, and with no significant differences between the approaches	۵.	۵-	۵.	The three approaches result in the same outcome with regard to pain relief and complication rate Adhesiolysis is an effective technique in managing post lumbar surgery syndrome pain

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Study	Number of	Interventions	Outcome measures	Pain relief and function	function		Results			Comments/conclusions
Study characteristic Methodological quality scoring	patients and selection criteria			3 months	6 months	1 year	3 months	3 months 6 months 1 year	l year	
Lumbar spinal stenosis	osis									
Manchikanti	50 patients	Adhesiolysis with Racz catheter, NRS, ODI,	NRS, ODI,	90 %08	Jo %08	Jo %92	Ъ	Ь	Ь	This is the first RCT
et al. (2009)	Percutaneous	followed by injection of 5 ml	employment status,	adhesiolysis	adhesiolysis	adhesiolysis				conducted in
[56, 57]	adhesiolysis = 25	of 2% preservative free	opioid use	had > 50%	had > 50%	had > 50%				managing chronic low
Randomized	patients	lidocaine and subsequent	Significant	relief versus	relief versus	relief at				back pain secondary
active control	Carron	injection of 6 ml of 10%	improvement = 50%	26% for	12% for	12 months				to lumbar central
	Caudal epidural	hypertonic sodium chloride	or more pain relief	caudal	caudal	after 3.5				spinal canal stenosis
Quality scores:	injections = 25	solution and 6 mg of	and improvement in			average				with percutaneous
Cochrane = 11/	patients	nonparticulate	functional status			injections				adhesiolysis
13		betamethasone								The treatment was
IPM-QRB = 36/		Control group: Catheter passed								offered based on the
48		to S2 with injection of 5 ml								return of pain
		of 2% preservative free								Robust outcome criteria
		lidocaine with subsequent								were utilized
		sodium chloride solution and								High-quality study with
		6 mg of nonparticulate betamethasone								positive results
		There was reduction of opioid								
		intake in the assessment								

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Study	Number of	Interventions	Outcome	Pain relief and function	ion		Results			Comments/conclusions
Study characteristic Methodological quality scoring	patients and selection criteria		measures	3 months	6 months	1 year	3 months	1 year 3 months 6 months 1 year	l year	
Karm et al. (2018) [61] Randomized, active control Quality scores: Cochrane = 11/ 13 IPM-QRB = 34/ 48	44 patients Adhesiolysis with Racz or NaviCath catheter = 20 patients Adhesiolysis with inflatable balloon catheter = 24 patients	Adhesiolysis with carheter without balloon received adhesiolysis, injections of 2 ml of 1% lidocaine with steroid, 5 mg of dexamethasone, and 15 IU hyaluronidase. In the recovery room, patients received 2 ml of 10% hypertonic saline for 2 days Inflatable balloon carheter adhesiolysis followed by injection of 4 ml of 1% lidocaine, and 15 IU of hyaluronidase. After adhesiolysis, 5 mg of dexamethasone and 1% lidocaine at each target site, 2 ml each was injected. In the recovery room, a test injection of 2 ml of 1% lidocaine and additional 4 ml of 10% hypertonic saline through the Perifix carheter for 2 days	NRS, ODI, Global Perceived Effect of Satisfaction, Medication Quantification Scale III	40% adhesiolysis group 58% in inflatable balloon catheter group	group group 58% in inflatable balloon catheter group	N N N N N N N N N N N N N N N N N N N	z	z	₹ Z	In a complicated assessment, the authors compared an inflatable balloon catheter with a balloon-less catheter in central lumbar spinal stenosis The authors performed a 2 day procedure in both groups. Inflatable balloon catheter showed significantly better improvement, with 58% of the patients considered as successful responders, and 40% at 3 months and 25% at 6 months in the balloon-less catheter group Overall, this is considered as a negative study for adhesiolysis with Racz catheter, whereas it was considered as positive for inflatable balloon catheter

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Study	Number of	Interventions	Outcome	Pain relief and function	ınction		Results			Comments/conclusions
Study characteristic Methodological quality scoring	patients and selection criteria		measures	3 months	6 months	l year	3 months	6 months	1 year	
Lumbar disc hemiation Gerdesmeyer 90 et al. (2013) [58] Randomized, placebo control Quality scores: Cochrane = 13/ 13 IPM—QRB = 44/ 48	90 patients from 381 patients with chronic radicular pain lasting longer than 4 months over a period of 4 years	Randomization: Placebo control group = 44 In the placebo group, a needle and catheter were inserted through caudal approach and the needle was intentionally inserted without entering the spinal canal and the catheter was inserted into the subcutaneous tissue overlying the afflicted level 10 ml of preservative free sodium chloride solution was injected for 3 days and the catheter was removed Neurolysis group = 46 The catheter was placed through the sacral canal with injection of 10 ml of contrast with identification of filling defects. Subsequently, a Tun-L catheter was inserted through the epidural needle and advanced to the anterolateral area of the filling defect Local anesthetic, 10 ml, 0.25% bupivacaine was injected through the catheter, followed by 10 ml of preservative free sodium chloride solution containing 150 units per ml of hyaluronidase Sodium chloride solution, 10 ml, 10%, containing 40 mg of triamcinolone was then injected slowly, along with 2 ml of 0.25% bupivacaine On the second and third days, 10 ml of 0.25% bupivacaine was injected through the eatheter, of the catheter, of the catheter, of the second and third days, 10 ml of 0.25% bupivacaine was injected through the eatheter, of the catheter, of the catheter of the cath	Primary outcome measure: ODI at 3, 6, and 12 months VAS: At least 50% reduction in ODI scores and VAS scores at 3, 6, and 12 months after reatment	Placebo group = 17% (7/42) Lysis group = 58% (26/45)	Placebo group = 11% (4/37) Lysis group = 74% (31/42)	Placebo group = 35% (9/26) Lysis group = 90% (28/31)	ور	۵.	<u>e</u> .	This is the first true placebo-controlled trial, injecting an inert substance into an inert structure, yet it showed positive response in some patients. Overall, there were significant differences in the active treatment group and lysis group compared with the placebo group. Ten year follow-up also showed significant improvement and surgery was avoided in the majority of patients. Surgery was avoided in 85% of patients
		followed by slow injection of 10 ml of 10% sodium chloride solution and 2 ml, 0.25% bupivacaine								
					1					

VAS visual analog scale, IPM-QRB Interventional Pain Management techniques-Quality Appraisal of Reliability and Risk of Bias Assessment, NA not applicable, N negative, P positive, NRS Numeric Pain Rating Scale, ODI Oswestry Disability Index, RCT randomized controlled trial, GHS Gerbershagen score

and single-arm meta-analyses. Further, the results of grading utilizing the GRADE system of appraisal for determining the body of evidence showed no change in the evidence levels.

Quantitative Analysis

Pain Level at 3 Months Figure 2A–C shows the change in pain level using the Numeric Rating Scale (NRS) or Visual Analog Scale (VAS) at 3 months.

Dual-Arm Meta-analysis There were seven trials [52–56, 58, 61] with 548 patients that compared percutaneous adhesiolysis with the control group in a dual-arm meta-analysis. The results showed a statistically significant difference in pain levels between these two groups [SMD -1.21 (-1.67, -0.75), p < 0.0001] (Fig. 2A).

Single-Arm Meta-analysis Figure 2B shows the results of a single-arm meta-analysis utilizing the percutaneous adhesiolysis group. There were eight trials [52–58, 61] that assessed pain scores at 3 months using NRS or VAS in patients who underwent percutaneous adhesiolysis. As shown in Fig. 2B, the pooled mean difference of pain scores from the baseline to 3 month follow-up was a 4.499 point decrease (95% CI -4.608 to -4.390, p < 0.0001).

Figure 2C shows the results of a single-arm meta-analysis utilizing a control group. There were seven trials [52–56, 58, 61] that assessed pain scores at 3 months using NRS or VAS in patients from the control group. As shown in Fig. 2C, the pooled mean difference of pain scores from the baseline to 3 month follow-up was a 2.585 point decrease (95% CI -2.750 to -2.419, p < 0.001).

Functionality at 3 Months Figure 3A–C shows the change in functionality level using the Oswestry Disability Index (ODI) at 3 months.

Dual-Arm Meta-analysis There were seven trials [52–56, 58, 61] with 548 patients that compared percutaneous adhesiolysis with a control group in a dual-arm meta-analysis. The results showed a statistically significant

difference in functionality levels between these two groups [SMD -1.10 (-1.53, -0.67), p < 0.0001] (Fig. 3A).

Single-Arm Meta-analysis Figure 3B shows the results of a single-arm meta-analysis utilizing the percutaneous adhesiolysis group. There were eight trials [52–58, 61] that assessed the functionality scores at 3 months using ODI in patients who underwent percutaneous adhesiolysis. As shown in Fig. 3B, the pooled mean difference of functionality scores from the baseline to 3 month follow-up was a 15.914 point decrease (95% CI -16.458 to -15.371, p < 0.0001).

Figure 3C shows the results of a single-arm meta-analysis utilizing a control group. There were seven trials [52–56, 58, 61] used to assess functionality scores at 3 months using ODI in patients from the control group. As shown in Fig. 3C, the pooled mean difference of functionality scores from the baseline to 3 month follow-up was a 7.819 point decrease (95% CI: 8.616 to -7.021, p < 0.0001).

Opioid Consumption at 3 Months Figure 4A–C shows the change in opioid intake using the morphine milligram equivalent scale (MMEq) at 3 months.

Dual-Arm Meta-analysis There were three trials [54–56] with 287 patients that compared percutaneous adhesiolysis with the control group in a dual-arm meta-analysis. The results showed no statistically significant difference in opioid intake between these two groups [SMD -0.32 (-0.78, 0.13) p = 0.16] (Fig. 4A).

Single-Arm Meta-analysis Figure 4B shows the change in opioid intake using the MMEq at 3 months for patients undergoing percutaneous adhesiolysis. There were four trials [54–57] with a pooled mean reduction in opioid intake from baseline to 3 months of follow-up of 13.493 MMEq (95% CI - 18.266 to - 8.720, p < 0.0001).

Figure 4C shows the change in opioid intake using the MMEq at 3 months for patients in the control treatment. There were three trials [54–56] with a pooled mean reduction in opioid intake from baseline to 3 months of follow-up

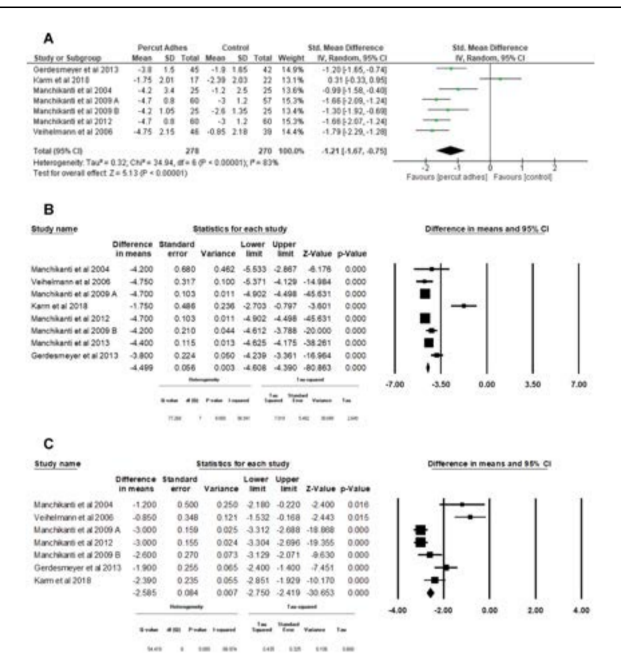


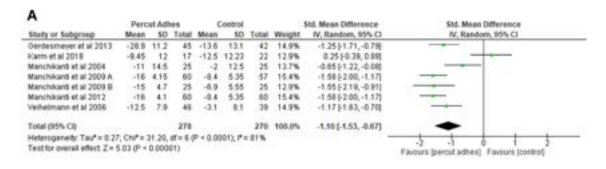
Fig. 2 Assessment of pain levels at 3 months utilizing dual-arm and single-arm meta-analyses. **A** Pain at 3 months, percutaneous adhesiolysis versus control, dual-arm meta-analysis. **B** Pain at 3 months in percutaneous

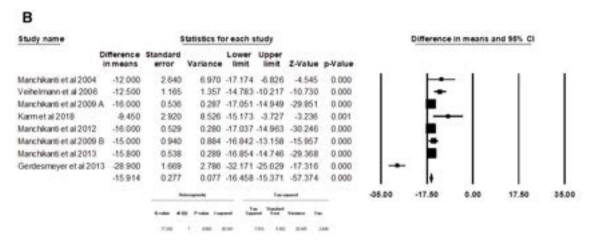
adhesiolysis groups with single-arm meta-analysis. C Pain at 3 months in control groups with single-arm meta-analysis

of 2.736 MMEq (95% CI -7.406 to 1.935, p = 0.251).

Overall, at 3 months, there was a significant improvement with percutaneous adhesiolysis utilizing dual- and single-arm meta-analyses with pain and function. In reference to opioid

consumption, while there was no significant difference with the dual-arm analysis, with the single-arm analysis, opioid consumption was decreased by 13.5 MMEq in percutaneous adhesiolysis groups, whereas in the control groups, it was reduced by 2.7 MMEq. Further,





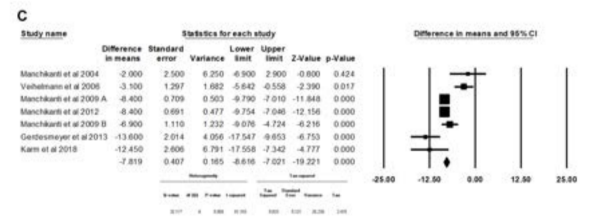


Fig. 3 Assessment of functional status at 3 months utilizing dual-arm and single-arm meta-analyses. **A** Functionality at 3 months, percutaneous adhesiolysis versus control, dual-arm meta-analysis. **B** Functionality at

3 months in percutaneous adhesiolysis groups with single-arm meta-analysis. ${f C}$ Functionality at 3 months in the control single-arm meta-analysis

there was significant decrease in pain relief of 4.5 points in adhesiolysis groups compared with 2.6 points in control groups.

Pain at 6 Months: Percutaneous Adhesiolysis versus Control Figures 5A-C showed the

change in pain level using the NRS or VAS at 6 months.

Dual-Arm Meta-analysis There were seven trials [52–56, 58, 61] with 504 patients that compared percutaneous adhesiolysis with a

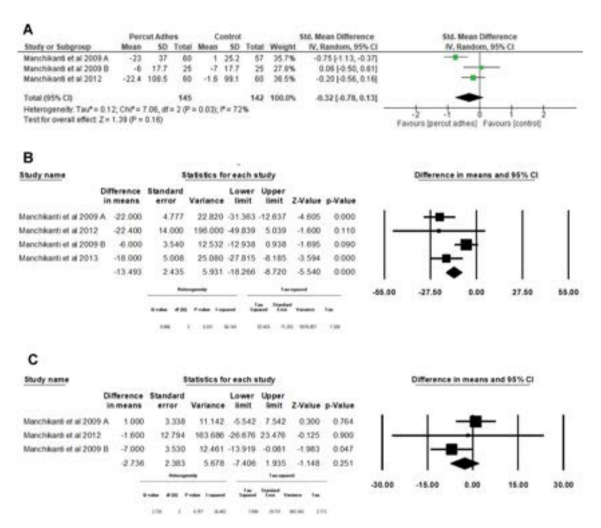


Fig. 4 Assessment of opioid consumption at 3 months utilizing dual-arm and single-arm meta-analyses. **A** Opioid consumption at 3 months, percutaneous adhesiolysis versus control, dual-arm meta-analysis. **B** Opioid consumption at

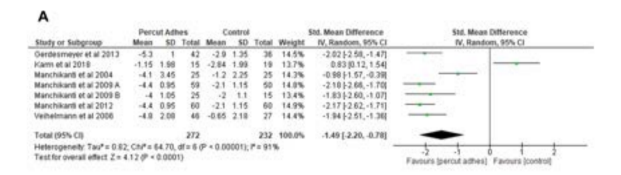
3 months in percutaneous adhesiolysis groups with singlearm meta-analysis. C Opioid consumption at 3 months in control groups with a single-arm meta-analysis

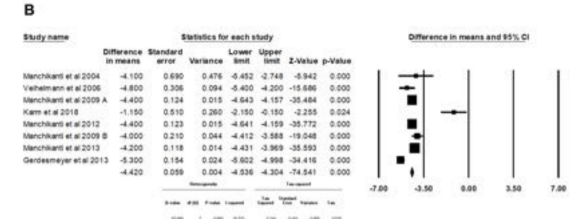
control group in a dual-arm meta-analysis. The results showed a statistically significant difference in pain levels between these two groups [SMD -1.49 (-2.20, -0.78), p < 0.0001] (Fig. 5A).

Single-Arm Meta-analysis Figure 5B shows the results of a single-arm meta-analysis utilizing a percutaneous adhesiolysis group. There were eight trials [52–58, 61] that assessed pain scores at 6 months using NRS or VAS in patients who underwent percutaneous adhesiolysis. As shown in Fig. 5B, the pooled mean difference of

pain scores from the baseline to 6 month follow-up was a 4.420 point decrease (95% CI -4.536 to -4.304, p < 0.0001).

Figure 5C shows the results of a single-arm meta-analysis with a control group. There were seven trials [52–56, 58, 61] used to assess pain scores at 6 months using NRS or VAS in patients from the control group. As shown in Fig. 5C, the pooled mean difference of pain scores from the baseline to 6 month follow-up was a 2.141 point decrease (95% CI -2.313 to -1.970, p < 0.0001).





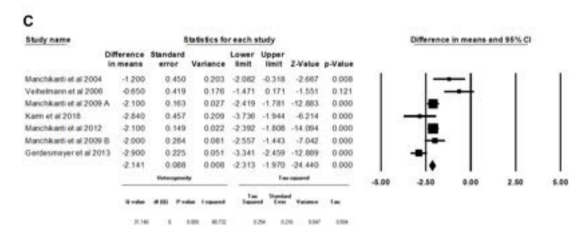


Fig. 5 Assessment of pain levels at 6 months utilizing dual-arm and single-arm meta-analyses. A Pain at 6 months, percutaneous adhesiolysis versus control, dual-arm meta-analysis, single-arm meta-analysis. B Pain at

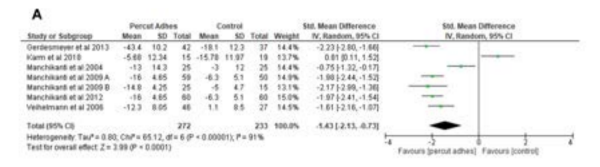
Functionality at 6 Months

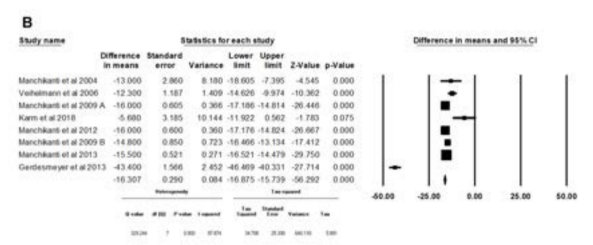
Figure 6A–C shows the change in functionality level using the ODI at 6 months.

Dual-Arm Meta-analysis There were seven trials [52–56, 58, 61] with 505 patients that

6 months in percutaneous adhesiolysis groups with singlearm meta-analysis. C. Pain at 6 months in control groups with single-arm meta-analysis

compared percutaneous adhesiolysis with a control group in a dual-arm meta-analysis. The results showed a statistically significant difference in functionality levels between these two groups [SMD -1.43 (-2.13, -0.73), p < 0.0001] (Fig. 6A).





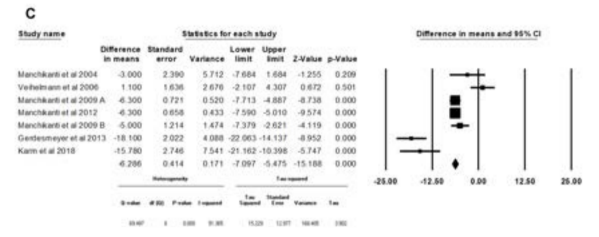


Fig. 6 Assessment of functional status at 6 months utilizing dual-arm and single-arm meta-analyses. A Functionality at 6 months in percutaneous adhesiolysis versus control, in a dual-arm meta-analysis. B Functionality at

6 months in the percutaneous adhesiolysis group, singlearm meta-analysis. **C** Functionality at 6 months in the control group, single-arm meta-analysis

Single-Arm Meta-analysis Figure 6B shows the results of a single-arm meta-analysis of percutaneous adhesiolysis. There were eight trials [52–58, 61] that assessed functionality scores at 6 months using ODI in patients who

underwent percutaneous adhesiolysis. As shown in Fig. 6B, the pooled mean difference of functionality scores from the baseline to 6 month follow-up was a 16.307 point decrease (95% CI -16.875 to -15.739, p < 0.0001).

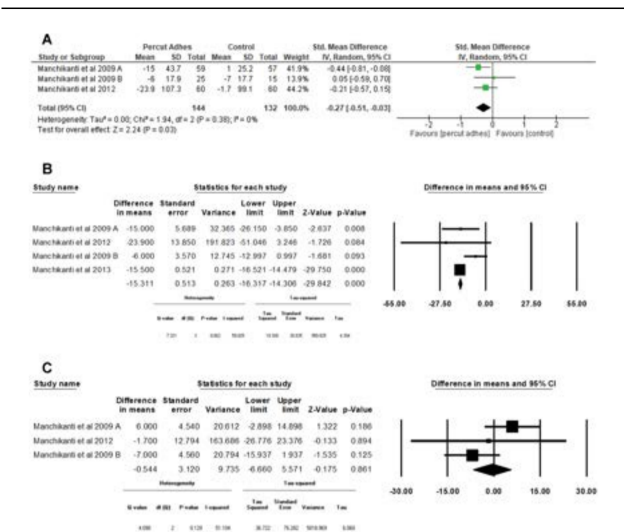


Fig. 7 Assessment of opioid consumption at 6 months utilizing dual-arm and single-arm meta-analyses. **A** Opioid consumption at 6 months, percutaneous adhesiolysis versus control, dual-arm meta-analysis. **B** Opioid consumption at

6 months in percutaneous adhesiolysis groups with a single-arm meta-analysis. **C.** Opioid consumption at 6 months in control groups with a single-arm meta-analysis

Figure 6C shows the results of a single-arm meta-analysis from the control group. There were seven trials [52–56, 58, 61] that assessed functionality scores at 6 months using ODI in patients from the control group. As shown in Fig. 6C, the pooled mean difference of functionality scores from the baseline to 6 month follow-up was a 6.286 point decrease (95% CI -7.097 to -5.475, p < 0.0001).

Opioid Consumption at 6 Months Figure 7A–C shows the change in opioid intake using the MMEq at 6 months.

Dual-Arm Meta-analysis There were three trials [54–56] with 276 patients that compared percutaneous adhesiolysis with a control group in a dual-arm meta-analysis. The results showed a statistically significant difference in opioid intake between these two groups [SMD -0.27 (-0.51, -0.03) p = 0.03] (Fig. 7A).

Single-Arm Meta-analysis Figure 7B shows the change in opioid intake using the MMEq at 6 months for patients undergoing percutaneous adhesiolysis. There were four trials [54–57], with a pooled mean decrease in opioid intake from

baseline to 6 months of follow-up of 15.311 MMEq (95% CI -16.317 to -14.306, p < 0.0001).

Figure 7C shows the change in opioid intake using the MMEq at 6 months for patients from the control group. There were three trials [54–56] with a pooled mean decrease in opioid intake from baseline to 6 months of follow-up of 0.544 MMEq (95% CI -6.660 to 5.571, p = 0.861).

Overall, at 6 months follow-up, pain and function improved significantly on dual- and single-arm analyses. Marked changes were observed with the single-arm analysis from baseline to the treatment. In addition, opioid consumption at 6 months also showed a significant difference with dual-arm analysis. However, these differences were significant with single-arm analysis, with a decrease of 15.3 MMEq in percutaneous adhesiolysis groups compared with 0.5 MMEq in control groups.

Pain at 12 Months Figure 8A–C shows the change in pain level using the NRS or VAS at 12 months.

Dual-Arm Meta-analysis There were six trials [52–56, 58] with 362 patients that compared percutaneous adhesiolysis with a control group in a dual-arm meta-analysis. The results showed a statistically significant difference in pain levels between these two groups [SMD -1.71 (-2.19, -1.22), p < 0.0001] (Fig. 8A).

Single-Arm Meta-analysis Figure 8B shows the results of a single-arm meta-analysis utilizing a percutaneous adhesiolysis group. There were seven trials [52–58] used to assess pain scores at 12 months using NRS or VAS in patients who underwent percutaneous adhesiolysis. As shown in Fig. 8B, the pooled mean difference of pain scores from the baseline to 12 month follow-up was a 4.226 point decrease (95% CI: -4.352 to -4.099, p < 0.0001).

Figure 8C shows the results of a single-arm meta-analysis utilizing the control group. There were six trials [52–56, 58] used to assess pain scores at 12 months using NRS or VAS in patients who from the control group. As shown in Fig. 8C, the pooled mean difference of pain scores from the baseline to 12 months follow-up

was a 2.156 point decrease (95% CI -2.409 to -1.904, p < 0.0001).

Functionality at 12 Months Figure 9A–C shows the change in functionality level using the ODI at 12 months.

Dual-Arm Meta-analysis There were six trials [52–56, 58] with 362 patients that compared percutaneous adhesiolysis with a control group in a dual-arm meta-analysis. The results showed a statistically significant difference in functionality levels between these two groups [SMD -1.65 (-2.09, -1.21), p < 0.0001] (Fig. 9A).

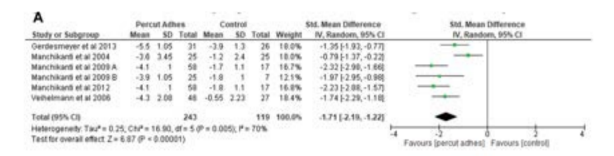
Single-Arm Meta-analysis Figure 9B shows the results of a single-arm meta-analysis in patients undergoing percutaneous adhesiolysis. There were seven trials [52–58] used to assess functionality scores at 12 months using ODI in patients who underwent percutaneous adhesiolysis. As shown in Fig. 9B, the pooled mean difference of functionality scores from the baseline to 12 months follow-up was a 15.881 point decrease (95% CI -16.485 to -15.277, p < 0.0001).

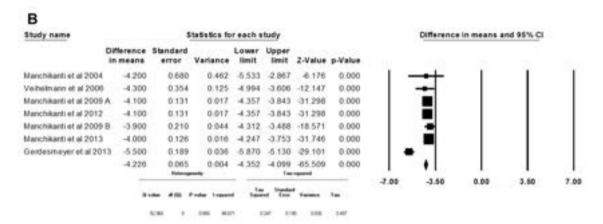
Figure 9C shows the results of a single-arm meta-analysis utilizing a control group. There were six trials [52–56, 58] used to assess functionality scores at 12 months using ODI in patients from the control group. As shown in Fig. 9C, the pooled mean difference of functionality scores from the baseline to 12 months follow-up was a 5.387 point decrease (95% CI -6.646 to -4.129, p < 0.0001).

Opioid Consumption at 12 Months Figure 10A–C shows the change in opioid intake using the MMEq at 12 months.

Dual-Arm Meta-analysis There were three trials [54–56] with 182 patients that compared percutaneous adhesiolysis with the control group in a dual-arm meta-analysis. The results showed no statistically significant difference in opioid intake between these two groups [SMD -0.31 (-0.69, 0.06) p = 0.10] (Fig. 10A).

Single-Arm Meta-analysis Figure 10B shows the change in opioid intake using the MMEq at





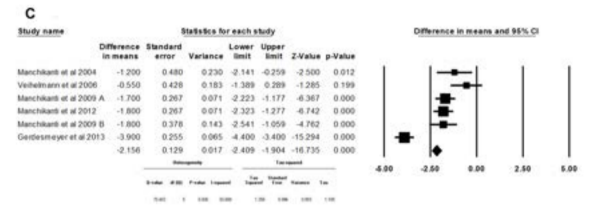
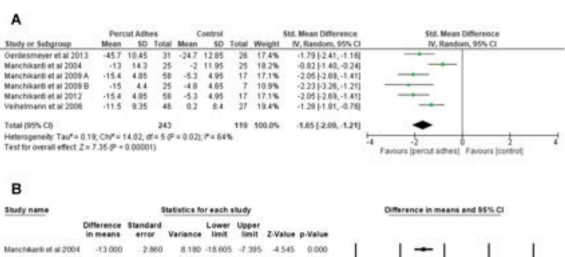


Fig. 8 Assessment of pain levels at 12 months utilizing dual-arm and single-arm meta-analyses. **A** Pain at 12 months, percutaneous adhesiolysis versus control, dual-arm meta-analysis. **B** Pain at 12 months in

percutaneous adhesiolysis groups with single-arm meta-analysis. ${f C}$ Pain at 12 months in control groups with single-arm meta-analysis

12 months for patients undergoing percutaneous adhesiolysis. There were four trials [54–57] with a pooled mean decrease in opioid intake from baseline to 12 months of follow-up of 15.094 MMEq (95% CI -16.141 to -14.048, p < 0.0001).

Figure 10C shows the change in opioid intake using the MMEq at 12 months for patients from the control group. There were three trials [54–56] with a pooled mean decrease in opioid intake from baseline to 12 months of



hidy name			itatistics fo	or each s	hudy				Difference	in means a	and 95% CI	
	Difference in means	Standard error	Variance	Lower limit	Upper limit	Z-Value	p-Value					
lanchkanti et al 2004	-13.000	2.860	8.180	-18.605	-7.395	4.545	0.000	1	1 -	- 1	1	- 1
eihelmann et al 2006	-11.500	1.379	1.902	-14.203	-8.797	-8.339	0.000		- 55	-		- 1
lanchikanti et al 2009 A	-15.400	0.637	0.406	-16.648	-14.152	-24.176	0.000					- 1
lanchikunti of all 2012	-15,400	0.637	0.406	-16.648	-14.152	24 176	0.000			i I		
lanchikanti et al 2009 (-15.000	0.880	0.774	-16.725	-13.275	-17.045	0.000			i		
lanchkanti et al 2013	-15.200	0.544	0.296	-16.266	-14.134	-27.941	0.000					
erdesmeyer et al 201.	45.700	1.877	3.523	49.379	42.021	24347	0.000	-	1 7			
	-15.881	0.308	0.095	-16.485	-15.277	-51.536	0.000	1,000	1 1		- 1	- 1
		Hatanganek			-			-55.00	-27.50	0.00	****	55.00
	5-97			-	thested			-99.00	47.00	0.00	27.50	99.00

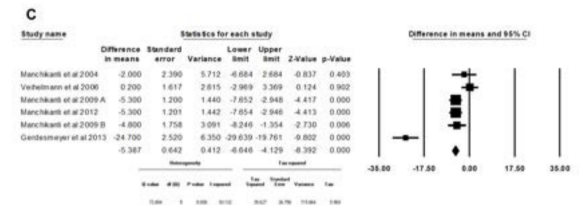


Fig. 9 Assessment of functional status at 12 months utilizing dual-arm and single-arm meta-analyses. A Functionality at 12 months, percutaneous adhesiolysis versus control, dual-arm meta-analysis. B Functionality at

12 months in percutaneous adhesiolysis groups with single-arm meta-analysis. **C** Functionality at 12 months in control groups with single-arm meta-analysis

follow-up of 2.664 MMEq (95% CI -11.399 to 6.070, p = 0.550).

Overall, at 1 year of follow-up, pain and function improved significantly in percutaneous adhesiolysis groups compared with the control groups with dual- and single-arm analyses. Opioid intake was not significantly

different with dual-arm analysis between both groups, even though single-arm analysis showed a significant difference, with a decrease of 15 MMEq in percutaneous adhesiolysis groups compared with a decrease of 2.7 MMEq in control groups.

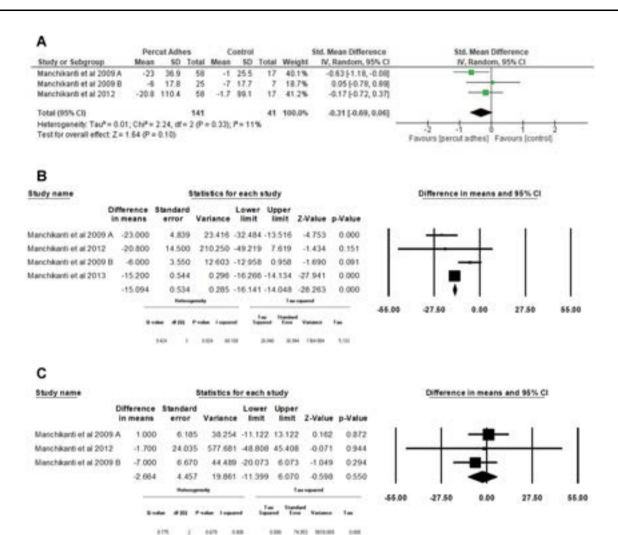


Fig. 10 Assessment of opioid consumption at 12 months utilizing dual-arm and single-arm meta-analyses. A Opioid consumption at 12 months, percutaneous adhesiolysis versus control, dual-arm meta-analysis. B Opioid

consumption at 12 months in percutaneous adhesiolysis groups with single-arm meta-analysis. C Opioid consumption at 12 months in control groups with single-arm meta-analysis

DISCUSSION

The present systematic review and meta-analysis of percutaneous adhesiolysis for low back and lower extremity pain secondary to post-surgery syndrome, central spinal stenosis, and chronic disc herniation showed level I–II or strong to moderate evidence with nine relevant RCTs with moderate to strong strength of recommendation. The RCTs were from six trials studying postsurgery syndrome, two trials studying spinal stenosis, and one randomized placebo-controlled trial studying disc

herniation. All other trials were active controlled. Past analysis of evidence synthesis based on individual conditions showed level I evidence for postsurgery syndrome [1, 3, 5, 6] and level II evidence for central spinal stenosis [1, 2] and chronic disc herniation [1, 3]. In contrast to previous reviews, the present meta-analysis combines all updates with utilization of all nine RCTs for three conditions [1–4].

Failed back surgery syndrome (FBSS) was defined by the International Association for the Study of Pain (IASP) as a phenomenon of persistent or recurrent pain, mainly in the lower back or legs, even after previously anatomically successful surgeries [77]. FBSS has been described extensively [72, 78–82]. The most common causes of FBSS have been identified as epidural fibrosis, arachnoiditis, recurrent disc herniation, and lateral and central spinal stenosis. Spinal stenosis is the result of abnormal narrowing of the spinal canal, lateral recess, or the intervertebral foramina, resulting in pressure on the spinal cord and/or nerve roots [83–86].

Among the studies that met the inclusion criteria, seven trials provided 1 year results and all of them reported positive results. However, at 6 months follow-up of the nine trials, one trial reported negative results [61]. The one negative trial [61] essentially compared percutaneous adhesiolysis with an inflatable balloon catheter. The inflatable balloon catheter had better results.

The results of this systematic review are in agreement with the majority of previous studies [1–7], except for one notable exception [5, 8]. The systematic review performed by Brito-García et al. [8] did not include a meta-analysis. Since the publication of the Brito-García review [8], other RCTs have been published. While multiple systematic reviews showed positive evidence ranging from level I to II [1–6], Cho et al. [7] showed a higher level of evidence for adhesiolysis than SCS.

Systematic reviews and meta-analysis are performed to meet the goals of evidence-based medicine using the best available evidence in determining clinical care for an individual patient or population. While systematic reviews and meta-analyses are expected to provide information from high-quality research, they may vary and do not guarantee high methodological and reporting rigor [39]. In the scientific world, multiple biases may be present, with publication bias, outcome reporting bias, multiple publication bias, place of publication bias, citation bias, and interpretation bias, which appear to be crucial and relevant to systematic reviews in interventional pain management [39]. Of importance is the interpretation bias referring to the researchers' or reviewers' abilities to synthesize and objectively judge and weigh the results found in a study. Consequently, two researchers of different backgrounds might look at the same result in a different way, thus drawing different conclusions based on their own background [87–89]. This is common when the data are debatable or qualitative, leading to some conclusions being overstated while others are understated [89]. The major issue is the erroneous classification of trials as "pragmatic" and "real world". Dal-Ré et al. [50] described that a genuinely pragmatic RCT should fulfill at least two fundamental features, including conduct of the study, which should resemble usual clinical practice, and the applicability of the results to multiple other settings, i.e., real world, not only the one where the trial was conducted. They also showed that some RCTs overtly deviate from usual clinical care and pragmatism, yet many RCTs are classified as pragmatic for purposes of convenience since pragmatic trials are set to represent realworld evidence. A recent publication of epidural steroids in disc herniation and sciatica in response to Cochrane review [90] illustrated multiples of these issues [38, 39]. Further, the role of placebo also has been a seeming source of continuous debate and has been the cause of discordance [40]. In fact, Manchikanti et al. [5] performed a systematic analysis of findings of systematic reviews in post lumbar surgery syndrome showing high compliance in only one systematic review [6] and moderate compliance with two systematic reviews [3, 7], whereas, one systematic review showed negative results with low compliance rate [8] with the PRISMA checklist. A Measurement Tool to Assess Systematic Reviews (AMSTAR) scoring also showed similar results, with high compliance for three systematic reviews [3, 6, 7] and poor compliance for one systematic review [8]. They also evaluated with Scottish Intercollegiate Guidelines Network (SIGN) scoring system showing similar results, thus one systematic review [8] consistently showed lower methodological quality. The present systematic review showed that all the trials included resembled clinical practice, with applicability of the results in a real-world setting.

Epidural steroid injections have been used extensively in managing low back and lower extremity pain [1, 10–16, 19, 22, 37–39, 91, 92]. Causes of chronic radicular pain include

mechanical compression of nerve roots, as well as different proinflammatory substances [1] that trigger ectopic neuron firing [1]. Chronic radicular pain secondary to postsurgery syndrome, central spinal stenosis, and disc herniamanaged with mechanical are decompression around the compressed nerve root, and inhibition of the inflammatory mediators by injecting targeted steroids into the epidural space or around the affected nerve. While results of studies of epidural injections continue to be debated and differ, the proportion of patients who failed to respond to epidural steroid injections are candidates for percutaneous epidural adhesiolysis [1, 42, 91, 92]. Thus far, the evidence has been in favor of percutaneous adhesiolysis in managing post lumbar surgery syndrome, spinal stenosis, and chronic disc herniation [1-6]. The mechanism described in percutaneous adhesiolysis is the combined effect of local lavage of proinflammatory cytokines, reduction of swelling, lysis of adhesions, desensitization and modification of neuromodulation, and local anesthesia. The presence of epidural adhesions may be diagnosed with magnetic resonance imaging (MRI), followed by epidurography based on filling defects. These filling defects by epidurography are minimized in size after successfully performing epidural lysis of adhesions. The epidural space is opened by injection of solutions if the catheter is placed directly into the zone of adhesions. However, the mechanical effect of adhesiolysis with a catheter has been debated [83]. It has been shown that the catheter itself was not able to have a significant mechanical effect in an experimental study setup. However, the authors themselves discussed the obvious limitations that the experimental setup did not represent the real clinical and anatomical environment. However, based on extensive clinical experience, we believe that the mechanical effects are real.

The limitations of this systematic review include the continued paucity of literature despite nine eligible trials that looked at various conditions separately (i.e., postsurgery syndrome, central spinal stenosis, and disc herniation). The other limitation is the lack of

placebo-controlled trials despite significant differences noted among the active-controlled trials utilizing epidural injection as control.

CONCLUSIONS

This systematic review with meta-analysis utilizing appropriate methodology with qualitative and quantitative evidence synthesis with conventional dual- and single-arm analyses shows level II–I, or moderate to strong evidence of effectiveness based on five high-quality and two moderate-quality RCTs with 1 year follow-up, showing improvement in pain and function as well as a decrease in opioid consumption.

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Compliance with Ethics Guidelines. This article is based on previously conducted studies and does not contain any studies with human participants or animals performed by any of the authors.

Data Availability. The data used in this review are available from multiple sources, including public databases PubMed, Google Scholar Cochrane library, US national Guidelines clearing house and are included in the references section of this article.

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