Systematic Assessment of Diagnostic Accuracy and Therapeutic Utility of Lumbar Facet Joint Interventions

Sukdeb Datta, MD1, Marion Lee, MD2, Frank J. Falco, MD3, David A. Bryce, MD4, and Salim M. Hayek, MD, PhD5

From: 1Vanderbilt University Medical Center, Nashville, TN; 2The Pain Center at Affinity Health Group, Tifton, GA; 3Mid Atlantic Spine & Pain Specialists, Newark, DE; 4Advanced Pain Management, Middleton, WI; and 5University Hospitals of Cleveland and Outcomes Research Consortium, Cleveland, OH.

Background: Lumbar facet joints are a well recognized source of low back pain and referred pain in the lower extremity in patients with chronic low back pain. Conventional clinical features and other non-invasive diagnostic modalities are unreliable in diagnosing lumbar zygapophysial joint pain. Controlled diagnostic studies have shown the prevalence of lumbar facet joint pain in 27% to 40% of the patients with chronic low back pain without disc displacement or radiculitis, with a false-positive rate of 27% to 47% with a single diagnostic block.

Study Design: A systematic review of diagnostic and therapeutic lumbar facet joint interventions.

Objective: To determine the clinical utility of diagnostic and therapeutic lumbar facet joint interventions in managing chronic low back pain of facet joint origin.

Methods: Review of the literature for clinical studies on efficacy and utility of facet joint interventions in diagnosing and managing facet joint pain was performed according to the Agency for Healthcare Research and Quality (AHRQ) criteria for diagnostic studies and observational studies and the Cochrane Musculoskeletal Review Group criteria as utilized for interventional techniques for randomized trials. Data sources included relevant literature of the English language identified through searches of Medline and EMBASE from 1966 to December 2008 and manual searches of bibliographies of known primary and review articles. Analysis results were performed for diagnostic and therapeutic interventions separately.

Level of Evidence: The level of evidence was defined as Level I, II, or III with 3 subcategories in Level II based on the quality of evidence developed by the U.S. Preventive Services Task Force (USPSTF) for therapeutic interventions.

Outcome Measures: For diagnostic interventions, studies must have been performed utilizing controlled local anesthetic blocks. Pain relief was categorized as at least 80% pain relief from baseline pain and ability to perform previously painful movements. For therapeutic interventions, the primary outcome measure was pain relief with secondary outcome measures of improvement in functional status, psychological status, return to work, and reduction in opioid intake. For therapeutic interventions, short-term pain relief was defined as relief lasting 6 months or less and long-term relief as longer than 6 months.

Results: Based on USPSTF criteria, evidence showed Level I or II-1 for diagnostic facet joint nerve blocks. Based on the review of included therapeutic studies, Level II-1 to II-2 evidence was indicated for lumbar facet joint nerve blocks with indicated level of evidence of Level II-2 to II-3 for lumbar radiofrequency neurotomy.

Limitations: The shortcoming of this systematic review of lumbar facet joint interventions is the paucity of published literature.

Conclusion: The evidence for diagnosis of lumbar facet joint pain with controlled local anesthetic blocks is Level I or II-1. The indicated level of evidence for therapeutic lumbar facet joint interventions is Level II-1 or II-2 for lumbar facet joint nerve blocks, Level II-2 or II-3 evidence for radiofrequency neurotomy, and Level III (limited) evidence for intraarticular injections.

Key words: Chronic low back pain, lumbar facet or zygapophysial joint pain, facet joint nerve blocks, medial branch blocks, controlled comparative local anesthetic blocks, lumbar radiofrequency neurotomy, lumbar intraarticular facet joint injections

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Lumbar facet joints are a well-recognized source of low back and referred pain in the lower extremity in patients with chronic low back pain (1-13). Facet joints are well innervated by the medial branches of the dorsal rami (14-19). Neuroanatomic, neurophysiologic, and biomechanical studies have demonstrated free and encapsulated nerve endings in lumbar facet joints, as well as nerves containing substance P calcitonin gene-related peptide (14,18-28).

Kalichman et al (29) evaluated facet joint osteoarthritis and low back pain in the community-based Framingham Heart Study. They concluded that there is a high prevalence of facet joint osteoarthritis in the community-based population with a prevalence of 59.6% in males and 66.7% in females. The prevalence of facet joint osteoarthritis increased with age and reached 89.2% in individuals 60 to 69 years old with highest prevalence of facet joint osteoarthritis found at the L4/5 spinal level. Further, they showed that individuals with facet joint osteoarthritis identified by computed tomographic (CT) scan at any spinal level showed no association with low back pain. Eubanks et al (30) in a study of 647 cadaveric lumbar spines found that facet joint osteoarthritis is a universal finding. Characteristic features of osteoarthritis begin to be seen early, with more than half of adults younger than 30 years demonstrating arthritic changes in the facets, with the most common arthritic level being L4/5. Thus, the published radiological investigations report no correlation between the clinical symptoms of low back pain and degenerative spinal changes observed on radiologic imaging studies, including radiographs, magnetic resonance imaging (MRI), CT scanning, single photon emission computed tomography (SPECT), and radionuclide bone scanning (1-3,31-33). Specifically, the association between degenerative changes in the lumbar facet joints and symptomatic low back pain remains unclear and is a subject of ongoing debate (31,34,35).

Conventional clinical features are unreliable in diagnosing lumbar zygapophysial (facet) joint pain (1-7,31-33,35-46). Hancock et al (47) performed a systematic review of tests to identify the disc, sacroiliac joint, and facet joint as the source of low back pain. They found that none of the tests for facet joint pain were found to be informative (31,36-38,48-55). Consequently, controlled local anesthetic blocks of the facet joint or its nerve supply are routinely employed to diagnose facet joint pain. The rationale for these blocks is that anesthetic blockade of a painful joint will abolish pain arising from that joint for the duration of the anesthetic effect, while anesthetic blockade of a non-painful joint will not alter the pain report. The probability that the blocked joint is the actual source of pain is increased if repeating the block with an anesthetic agent that has a different duration of action reproduces the analgesic response (33). To ensure accuracy and validity, these blocks must be controlled and verified for delivery of local anesthetic agent and placebo response. Fluoroscopic guidance and controlled dual blocks eliminate or greatly reduce placebo responses. Single facet joint injections are not recommended, as they do not control for a false-positive response (1-3,31,33,36-38,51,56-65). Rubinstein and van Tulder (66) also provided a best-evidence review of diagnostic procedures for neck and low back pain. They commented that it is quite remarkable that while many named orthopedic tests of the neck and low back are often illustrated in orthopedic textbooks, there is little evidence to support their diagnostic accuracy, and therefore their use in clinical practice. Consistent with clinical experience, many studies have demonstrated that the physical examination serves primarily to confirm suspicions raised during the history. The placebo controlled technique is considered the gold standard, but has limited clinical utility due to ethical and cost implications. Controlled comparative blocks with short and long acting local anesthetics are an acceptable alternative strategy (1-3,67-70).

Controlled comparative blocks have been criticized and the accuracy and validity of these precision diagnostic techniques have been questioned (71-79). Although these tests control and verify for location of local anesthetic delivery, they are faulted for assuming that the report and documentation of the magnitude and quality of pain relief are accurate. Because these tests employ subjective criteria, i.e., rely on a patient’s report of presence or absence of pain following a block and ability to isolate different painful areas, or differentiate between significant and insubstantial pain relief (when pain relief is incomplete), they promote doubt about the accuracy of these procedures.

Three systematic reviews have concluded the evidence for diagnostic accuracy of lumbar facet joint nerve blocks as strong (1-3). Further, Rubinstein and van Tulder (66) concluded that there is strong evidence for the diagnostic accuracy of facet joint blocks in evaluating spinal pain.
Facet joint pain may be managed by intraarticular injections, facet joint nerve blocks, and neurolysis of facet joint nerves. However, conflicting results have been reported for the value of the different treatment modalities in systematic reviews (5,6,80-86). A narrative review by Bogduk (87) suggested that intraarticular facet joint injections were no better than placebo for chronic lumbar spine pain. Boswell et al (5,6), in systematic reviews of therapeutic facet joint interventions, showed moderate evidence for lumbar intraarticular facet joint injections for short-term improvement, but only limited evidence for long-term improvement. Geurts et al (80) concluded that there was moderate evidence that radiofrequency lumbar facet denervation was more effective for chronic low back pain than placebo, and there was only limited evidence for effectiveness of radiofrequency neurotomy for chronic cervical zygapophysial joint pain after flexion/extension injury. However, Geurts et al (80) included both medial branch neurotomy and intraarticular neurotomy in their evaluation, along with dorsal root denervation. Manchikanti et al (83) evaluated medial branch neurotomy for the management of chronic spinal pain utilizing randomized and observational reports, and concluded that there was strong evidence for short-term relief and moderate evidence for long-term relief of facet joint pain.

This systematic review is undertaken to determine the accuracy of lumbar facet joint blocks in the diagnosis of chronic low back pain and to evaluate the effectiveness of therapeutic facet joint interventions in the treatment of chronic low back pain of lumbar facet joint origin.

**Methods**

**Literature Search**

A comprehensive literature search was conducted including Medline and EMBASE from 1966 through December 2008, Cochrane database, Clinical Trial Registry, systematic reviews, narrative reviews, and cross-references to the reviews published in the English language.

The search strategy consisted of chronic low back pain, facet or zygapophysial joint pain, lumbar facet injections, lumbar facet joint nerve blocks, intraarticular lumbar facet injections, and lumbar radiofrequency neurotomy.

**Diagnostic Lumbar Facet Joint Interventions**

**Selection Criteria**

All studies published on the diagnosis of lumbar facet joint pain in patients with chronic pain of greater than 3 months duration were included for review. Only the studies utilizing controlled diagnostic blocks either placebo or comparative local anesthetic blocks under fluoroscopy were included. The criterion standard for diagnosis of lumbar facet joint pain was at least 80% pain relief for the duration of local anesthetic and ability to perform previously painful movements.

All non-clinical studies were excluded. Further, case reports, book chapters, non-evidence-based guidelines, letters, and expert opinions were excluded.

**Review Criteria**

The manuscripts meeting the inclusion criteria of 80% relief with ability to perform previously painful maneuvers were included in methodologic quality assessment. Two physician reviewers evaluated and graded articles meeting inclusion criteria for methodologic quality and grading of evidence as described by the Agency for Healthcare Research and Quality (AHRQ) for diagnostic studies (88) and any disagreements were resolved by the third physician. Consensus-based weighted scoring developed by the guidelines committee of the American Society of Interventional Pain Physicians (ASIPP) was utilized, which was refined and used in other evaluations (85,86,89-94).

The quality of individual articles was evaluated using the above criteria with application of weighted scores. For inclusion in the analysis, each study should have scored at least 50 on a scale of 0 to 100.

If there was a conflict of interest with the reviewed manuscripts with authorship or any other type of conflict, the involved authors did not review the manuscripts for quality assessment, clinical relevance, evidence synthesis, or grading of evidence.

**Level of Evidence**

There is no hierarchy of evidence described for diagnostic studies grading and quality assessment as for therapeutic interventions. Thus, almost all diagnostic accuracy studies are observational. Thus, modified quality of evidence developed by the U.S. Preventive Services Task Force (USPSTF) as illustrated in Table 1 was utilized (95).
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Table 1. Modified quality of evidence developed by USPSTF.

<table>
<thead>
<tr>
<th>Level</th>
<th>Description</th>
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<tbody>
<tr>
<td>I</td>
<td>Evidence obtained from multiple properly conducted diagnostic accuracy studies.</td>
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<tr>
<td>II-1</td>
<td>Evidence obtained from at least one properly conducted diagnostic accuracy study of adequate size.</td>
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<tr>
<td>II-2</td>
<td>Evidence obtained from at least one properly designed small diagnostic accuracy study.</td>
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<td>II-3</td>
<td>Evidence obtained from diagnostic studies of uncertainty.</td>
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<td>III</td>
<td>Opinions of respected authorities, based on clinical experience descriptive studies and case reports or reports of expert committees.</td>
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Adapted and modified from the U.S. Preventive Services Task Force (USPSTF) (95).

Therapeutic Facet Joint Interventions

Inclusion Criteria

Three types of facet joint interventions were included in this review: intraarticular facet joint injections, facet joint nerve blocks, and medial branch radiofrequency neurotomy. All studies must have provided appropriate management with outcome evaluations of at least 6 months and appropriate statistical analysis. Studies should also have met diagnostic criteria with controlled (placebo or dual) diagnostic blocks with at least 80% relief.

Reports without appropriate diagnosis and elimination of false-positive responses, non-systematic reviews, book chapters, and case reports were excluded.

Outcome Parameters

The primary outcome measure was pain relief (short-term relief up to 6 months and long-term relief greater than 6 months) at various time points reported at least over a period of 6 months. The secondary outcome measures were functional status improvement, psychological status improvement, return to work, and opioid intake.

Methodologic Quality Assessment

The quality of each individual article used in this analysis was assessed by modified Cochrane review criteria with weighted scores (96) for randomized trials and AHRQ quality criteria for assessment of observational studies (88) with weighted scoring, as developed by ASIPP (85) and utilized in multiple other systematic reviews (86,92,97-105).

Only the studies scoring at least 50 of 100 on weighted scoring criteria were utilized for analysis.

Each study was evaluated by 2 physicians for stated criteria and any disagreements were resolved by the third physician. If there was a conflict of interest with the reviewed manuscripts with authorship or any other type of conflict, the involved authors did not review the manuscripts for quality assessment, clinical relevance, evidence synthesis, or grading of evidence.

If there were 4 randomized trials evaluating any one of the techniques — namely intraarticular injections — facet joint nerve blocks, or radiofrequency neurotomy, observational studies were not included in the methodologic quality assessment as well as the evidence synthesis.

Clinical Relevance

Clinical relevance of the included randomized trials was evaluated according to 5 questions recommended by the Cochrane Back Review Group (82,106). Each question was scored positive (+) if the clinical relevance item was met, negative (−) if the item was not met, and unclear (?) if data were not available to answer the question.

In the updated Cochrane review of “Injection Therapy for Subacute and Chronic Low Back Pain” (82) the authors considered a 20% improvement in pain scores (107) and a 10% improvement in functioning outcomes (108) to be clinically important. This review utilized stricter criteria than previous systematic reviews. Any relief of 6 months or less was considered as short-term, whereas Cochrane reviews (82) and others (5-7) have considered 6 weeks as short-term and longer than 6 weeks as long-term. We also utilized methodologic quality assessment criteria for minimum inclusion, thus this systematic review is expected to provide robust results, and the inclusion of observational studies is expected to improve the generalizability of the systematic review (109-113).

Analysis of Evidence

Analysis was conducted using 5 levels of evidence, ranging from Level I to III, with 3 subcategories in Level II developed by the USPSTF (95).
Recommendations

Grading recommendations were based on Guyatt et al’s criteria with 6 Levels, 1A – 1C strongly and 2A – 2C weak as illustrated in Table 2 (114).

Outcome of the Studies

A study was judged to be positive if the therapeutic facet joint interventions were clinically relevant and effective, either with a placebo control or active control in randomized trials. This indicates that the difference in the effect for the primary outcome measure was statistically significant on the conventional 5% level. In a negative study, no difference between the study treatments or no improvement from baseline was found. Further, the outcomes were judged at the reference point with positive or negative results reported at 3 months, 6 months, and one year. For observational studies, a study was judged to be positive if the authors concluded that the therapeutic facet joint interventions were effective, with outcomes reported at the reference point with positive or negative results at 3 months, 6 months, and one year.

Results

Diagnostic Lumbar Facet Joint Nerve Blocks

Our comprehensive search yielded 1,782 articles (Fig. 1). Of these, 74 full manuscripts were reviewed, whereas 35 manuscripts were considered for inclusion (31,32,36-38,46,48-65,76,115-124). Other manuscripts described pain patterns, nerve supply, and therapeutic interventions.

Methodologic Quality Assessment

A total of 7 studies met the inclusion criteria for methodologic assessment utilizing controlled local anesthetic blocks with evaluation of at least 80% pain relief and ability to perform multiple maneuvers (38,56-58,60,62,65), 7 studies were excluded as they evaluated with only single block (48-50,52-55), 14 studies were excluded as the inclusion criteria was the relief of pain less than 80% (31,32,36,37,46,51,59,61,76,116-118,122,123), 3 studies (63,64,124) were excluded as these were subgroup analysis of other studies, one study (121) evaluated validity of diagnosis with a 2-
Fig. 1. The flow diagram for diagnostic studies.
year follow-up, 2 studies (119,120) evaluated the effect of sedation, and another study (115) evaluated the role of psychological factors.

Methodologic quality assessment is illustrated in Table 3. The data and prevalence of the 7 included studies are illustrated in Table 4.

Manchikanti and coauthors in multiple publications (57,58,60) evaluated prevalence and false-positive rates of diagnostic blocks. In all included studies they utilized a criterion standard of 80% pain relief with the ability to perform previously painful movements without pain utilizing 1% lidocaine. In a large study of 500 patients in which prevalence of facet joint pain in chronic spinal pain of cervical, thoracic, and lumbar regions were evaluated (58), 397 patients were evaluated for low back pain showing a prevalence of 31% (95% CI, 27%, 36%) with a false-positive rate with single blocks with lidocaine of 27% (95% CI, 22%, 32%). The second large study by Manchukonda et al (60) evaluated 438 patients with 303 patients with lumbar pain. Prevalence of lumbar facet joint pain was determined as 27% (95% CI, 22%, 33%), with a false-positive rate of single blocks in the lumbar region of 45% (95% CI, 36%, 53%).

Schwarzer et al (38) determined the prevalence of pain arising from the zygapophysial joint in patients with chronic low back pain and to determine whether any clinical features could distinguish patients with and without such pain. The results showed 20 patients 32% (95% CI; 20% to 44%) obtained greater than 50% relief of their pain following the administration of saline. Fifty seven patients completed the study; 23 of them 37% (95% CI; 25% to 49%) failed to obtain relief following the injection of saline but obtained relief following one or more intraarticular injections of local anaesthetic. None of the historical features or clinical tests could discriminate those patients with and those without zygapophysial joint pain.

Manchikanti et al (62) in an evaluation of the relative contributions of various structures in chronic low back pain evaluated 120 patients with a chief complaint of low back pain who were evaluated with facet joint

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<td>Manchikanti et al 2002 (57)</td>
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<td>Manchukonda et al 2007 (60)</td>
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Adapted and modified from West S et al. Systems to Rate the Strength of Scientific Evidence, Evidence Report, Technology Assessment No. 47. AHRQ Publication No. 02-E016 (88).
nerve blocks, provocative discography, and sacroiliac joint injections. The results of this study showed that the facet joint is the most common pain generator in chronic low back pain with identification of the facet joint in 40% (95% CL, 31%, 49%) of patients, followed by the disc in 26% (95% CL, 18%, 34%) of patients, and the sacroiliac joint in only 2% of the patients.

Manchikanti et al (56) evaluated chronic low back pain of facet joint origin based on involvement of single or multiple spinal regions. The concluded that the prevalence of lumbar facet joint pain in patients with low back only was 21% (95% CI, 14%–27%) compared to 41% (95% CI, 33%, 49%) of the patients with low back pain with involvement of other regions of the spine with controlled comparative local anesthetic blocks. A false-positive rate of 17% (95% CI, 10%, 24%) in patients with low back pain only and 27% (95% CI, 18%, 36%) in patients with involvement of multiple regions of the spine was demonstrated with single blocks.

### Prevalence

The prevalence of lumbar facet joint pain based on the controlled diagnostic blocks is shown to be 21% to 41% (95% CI, 14% to 53%) with overall prevalence of 31% (95% CI 28%–33%).

### False-Positive Rates

False-positive rates of 17% to 49% were demonstrated with CIs ranging from 10% to 59% with overall false-positive rate of 30% (95% CI 27%–33%).

### Confounding Factors

Sedation as a confounding factor was evaluated in the lumbar spine. Studies by Manchikanti et al (118,119) have demonstrated that conscious sedation may provide that 5% of subjects in the placebo group and up to 10% of subjects in the active group reported ≥ 80% pain relief and were able to perform movements that were painful prior to the administration of the intravenous agents. Application of the ≥ 80% pain reduction criterion standard instead of ≤ 50% pain reduction decreased the false-positive response from 15% to 10%. A systematic review by Smith et al (94) showed no significant evidence of the influence of sedation either with midazolam or fentanyl in the evaluation of cervical and lumbar facet joint pain with controlled cervical and lumbar facet joint nerve blocks with an indicated evidence of Level II-1, with application of stringent criteria of at least 80% pain relief and the ability to perform previously painful movements after the diagnostic blocks.

### Diagnostic Accuracy

The accuracy was evaluated in 7 studies illustrating either prevalence or false-positive rates (Table 4). A total of 1,320 patients were studied for diagnostic accuracy. The overall prevalence has been demonstrated as 27% to 40% with 16% in postlumbar laminectomy patients. The recent study (60) with large population of 303 showed a prevalence of 27% with confidence intervals ranging from 29% to 22% to 33%.

Table 4. Data of prevalence with controlled diagnostic blocks and false-positive rates in lumbar region.

<table>
<thead>
<tr>
<th>Study</th>
<th>Methodological Criteria</th>
<th>Participants</th>
<th>Prevalence</th>
<th>False-Positive Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manchikanti et al 2002 (57)</td>
<td>75</td>
<td>120</td>
<td>40% (95% CI; 31%–49%)</td>
<td>30% (95% CI; 20%–40%)</td>
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<tr>
<td>Manchikanti et al 2004 (58)</td>
<td>75</td>
<td>397</td>
<td>31% (95% CI; 27%–36%)</td>
<td>27% (95% CI; 22%–32%)</td>
</tr>
<tr>
<td>Manchukonda et al 2007 (60)</td>
<td>75</td>
<td>303</td>
<td>27% (95% CI; 22%–33%)</td>
<td>45% (95% CI; 36%–53%)</td>
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<tr>
<td>Schwarzer et al 1995 (38)</td>
<td>75</td>
<td>63</td>
<td>37% (95% CI; 25%–49%)</td>
<td>NA</td>
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<tr>
<td>Manchikanti et al 2001 (62)</td>
<td>75</td>
<td>120</td>
<td>40% (95% CI; 31%–49%)</td>
<td>47% (95% CI; 35%–59%)</td>
</tr>
<tr>
<td>Manchikanti et al 2003 (56)</td>
<td>75</td>
<td>300</td>
<td>I. 21% (95% CI; 14%–27%)</td>
<td>I. 17% (95% CI; 10%–24%)</td>
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<td></td>
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<td></td>
<td>II. 41% (95% CI; 33%–49%)</td>
<td>II. 27% (95% CI; 18%–36%)</td>
</tr>
<tr>
<td>Manchikanti et al 2007 (65)</td>
<td>75</td>
<td>117</td>
<td>16% (95% CI; 9%–23%)</td>
<td>49% (95% CI; 39%–59%)</td>
</tr>
<tr>
<td>Overall</td>
<td></td>
<td>1,420</td>
<td>31% (95% CI; 28%–33%)</td>
<td>30%* (95% CI; 27%–33%)</td>
</tr>
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</table>

CI = confidence interval; NA = not available; # Schwarzer et al (38) not included
Influence of psychological factors was also evaluated (64,115). The evaluation by Manchikanti et al (64) showed no significant differences among patients either in terms of prevalence or false-positive rates or prevalence in the diagnosis of lumbar facet joint pain. The prevalence of facet joint pain was seen in 25% of the patients with no psychopathology with a 50% false-positive rate, whereas in patients with major depression, it was 31%, with generalized anxiety disorder, 28%, and in patients with somatization disorder, prevalence was 32%, and false-positive rates were 38%, 43%, and 39% respectively.

Age as a confounding factor was evaluated. Manchikanti et al (63) showed variable prevalence of facet joint involvement in lumbar spine ranging from 18% to 44%, with significant differences noted among some groups. The results showed in Group I (18 to 30 years), the prevalence was 28% with false-positive rates of 40%; in Group II (31 to 40 years), the prevalence was 18% with false-positive rates of 50%; in Group III (41 to 50 years), the prevalence was 28% with false-positive rates of 45%; in Group IV (51 to 60 years), the prevalence was 44% with false-positive rates of 30%; in Group V (61 to 70 years), the prevalence was 21% with false-positive rates of 64%; and in Group VI (over 70 years), the prevalence was 26% with false-positive rates of 43%. Thus, the highest prevalence was seen in patients aged 51 to 60 years, whereas the lowest was seen in the age group of 31 to 40 years with 18%. Manchikanti et al (65) also showed significantly less proportion of patients with facet joint pain after lumbar surgery and occupational injury (124) with no differences based on obesity (59), based on gender, or smoking status (124).

**Criterion Standard**

No tissue diagnosis (biopsy or autopsy) techniques are available to diagnose facet joint pain and confirm specificity and sensitivity of facet joint nerve blocks. However, pain relief and stability of the diagnosis with long-term follow-up are employed as the criterion standards and are accepted across different medical disciplines (1-3,74,121).

**Validity**

Controlled lumbar facet joint nerve blocks have been established as a method to diagnose lumbar facet joint pain, either with placebo control or controlled comparative local anesthetic blocks that meet specific criteria of pain relief and functional improvement (1,3,33,36-38,46,47,51,56-66). Pain relief (74) and long-term follow-up are employed as the criterion standards and are accepted across different medical disciplines (74,110,121). Long-term relief of lumbar facet joint interventions has been demonstrated (4-7,83,85,121).

**Study Designs**

All the studies included in the methodologic quality assessment met inclusion criteria as well as study design criteria. There has been significant controversy over the study designs with some reviewers calling for randomized controlled trials (RCTs) for diagnostic interventions (6,80). However, the design accuracy of diagnostic studies involves consecutive or non-consecutive allocation and observational studies (85).

**Level of Evidence**

The level of evidence was Level I or Level II-1 evidence based on the 7 included studies.

**Therapeutic Facet Joint Interventions**

A comprehensive literature search was performed for lumbar intraarticular facet joint injections, lumbar facet joint nerve blocks, and lumbar facet joint nerve radiofrequency neurotomy (Fig. 2). The entire search yielded a total of 40 studies relevant to therapeutic lumbar facet interventions (37,47,117,121,123,125-159).

**Randomized Trials**

**Methodologic Quality Assessment**

Of the 6 randomized trials identified evaluating the effectiveness of lumbar intraarticular facet joint injections (132-137), 5 studies failed to meet inclusion criteria for methodologic quality assessment due to lack of controlled diagnostic blocks (132-136), whereas one study (137) evaluated the injections of carpometacarpal (CMC) joints of the thumb.

There were 7 studies evaluating therapeutic lumbar facet joint nerve blocks (37,121,135,136,139-141). Of these, one study (37) evaluated only diagnostic interventions. Another study (121) evaluated diagnostic validity of lumbar facet joint nerve blocks. Two studies (135,136) had only short-term evaluation without diagnostic blocks. One study (139) was a preliminary report of the one-year report (140). Consequently 2 studies met inclusion criteria (140,141).

There were 7 studies evaluating radiofrequency neurotomy of lumbar facet joint nerves (142-144,147,148,150,155). Of these, only one study met
the inclusion criteria (142). van Wijk et al (143), Leclaire et al (144), Gallagher et al (147), van Kleef et al (150), and Tekin et al (155) were excluded due to lack of controlled diagnostic blocks. One study (148) was excluded as it evaluated intraarticular facet joint denervation, which is not medial branch neurotomy, with lack of appropriate diagnostic criteria.

Table 5 illustrates the methodologic quality assessment of randomized clinical trials evaluating the
Table 6 illustrates clinical relevance assessment of 3 randomized trials meeting the inclusion criteria with quality assessment.

Clinical Relevance Assessment

Table 6 illustrates clinical relevance assessment of 3 randomized trials meeting the inclusion criteria with quality assessment.

Observational Studies

Multiple observational studies evaluated the role of therapeutic facet joint interventions. Of these, 15 studies evaluated the role of intraarticular injections of lumbar facet joints (41,117,125-131,137,138,151-154). However, none met inclusion criteria.

There was only one study evaluating the role of facet joint nerve blocks compared to intraarticular injections (117), which failed to meet inclusion criteria.

Methodologic Quality Assessment

Table 7 illustrates the methodologic quality assessment criteria of observational studies. Both studies met methodologic quality assessment criteria with scores of 73 for the study by Dreyfuss et al (146) and 63 for the study by Gofeld et al (145).
Intraarticular Facet Joint Blocks

Of the 5 randomized trials (132-136) and 15 observational studies (41,117,125-131,137,138,151-154), none met inclusion criteria.

Excluded Studies of Intraarticular Injections

Carette et al (132) failed to exclude placebo responders, which may account for relatively high incidence of patients in their study with presumed facet joint pain. They showed an incidence of 58% prevalence of facet joint pain based on inclusion criteria of phase 1 of their study. Failure to exclude placebo responders may have diluted the findings of true responses, making detection of differences between the study and the control group difficult. The patients in the methylprednisolone group received a greater proportion of concurrent interventions. At 6 months, 42% of patients in the steroid group showed benefit compared to 15% in the sodium chloride solution group. They concluded that there was no significant difference between the two groups. However, the effects and consequences of intraarticular placebo injections of sodium chloride are not known. Normal saline has been shown to provide better pain relief than that expected with a true placebo for a multitude of invasive procedures (160-162). Consequently, even patients with placebo injections of sodium chloride responded to treatment similar to corticosteroid injections. Further, effect of injection of placebo or any other agent in a closed space such as intraarticular injection is not known. The intraarticular injection of sodium chloride solution may produce multiple effects which are not explained at the present time.

Fuchs et al (134) conducted a study comparing intraarticular hyaluronic acid versus glucocorticoid injections for nonradicular pain in the lumbar spine. Sixty patients were included in this randomized, controlled, blind-observer clinical study and randomly assigned to 2 groups to receive 10 mg of sodium hyaluronate or 10 mg of triamcinolone acetonide per facet joint. The facet joints on both sides at levels L5-S1, L5-L4, and L4-L3 were treated once per week under CT guidance. The study visits were timed to permit assessment of the immediate effect as well as possible carryover effects at 3 and 6 months after completion of treatments. Changes in pain were assessed with a visual analogue scale (VAS) and changes in function and quality of life were assessed by the Roland-Morris questionnaire (RMQ), the Oswestry Disability questionnaire (ODQ), the Low Back Outcome Score (LBOS) and the Short-Form 36 (SF-36). Patients reported lasting relief, better function, and improved quality of life with both treatments. The disadvantages of the study include a lack of appropriate diagnosis with controlled diagnostic blocks, thus failing to exclude placebo responders which may have increased the possibility of inclusion of patients without facet joint pain. Furthermore, pain relief of 50% or greater was achieved only in the triamcinolone group with a reduction of 51.7% despite a series of injections bilaterally at 3 levels, whereas the reduction was 45.1% in the sodium hyaluronate group. RMQ scores, ODQ scores, and LBOS showed reduction in sodium hyaluronate of 43.2%, 39.1%, and 43.9% whereas in the triamcinolone group the reduction was 33.4%, 29.5%, and 34.8%. Considering that no controlled diagnostic blocks were used, and no
Table 7. Methodologic quality assessment criteria for observational studies of lumbar facet joint interventions.

<table>
<thead>
<tr>
<th>CRITERION</th>
<th>Weighted Score (points)</th>
<th>Dreyfuss et al (146)</th>
<th>Gofeld et al (145)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Study Question</td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>• Clearly focused and appropriate question</td>
<td></td>
<td></td>
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<tr>
<td>2. Study Population</td>
<td>8</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>• Description of study population</td>
<td>5</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>• Sample size justification</td>
<td>3</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>3. Comparability of Subjects</td>
<td>22</td>
<td>8</td>
<td>8</td>
</tr>
<tr>
<td>• Specific inclusion/exclusion criteria for all groups</td>
<td>5</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>• Criteria applied equally to all groups</td>
<td>3</td>
<td>-</td>
<td>-</td>
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<tr>
<td>• Comparability of groups at baseline with regard to disease status and</td>
<td>3</td>
<td>-</td>
<td>-</td>
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<tr>
<td>prognostic factors</td>
<td></td>
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<tr>
<td>• Study groups comparable to non-participants with regard to confounding</td>
<td>5</td>
<td>-</td>
<td>-</td>
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<tr>
<td>factors</td>
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<tr>
<td>• Use of concurrent controls</td>
<td>5</td>
<td>-</td>
<td>-</td>
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<tr>
<td>• Comparability of follow-up among groups at each assessment</td>
<td>3</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>4. Exposure or Intervention</td>
<td>11</td>
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<td>8</td>
</tr>
<tr>
<td>• Clear definition of exposure</td>
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<td>5</td>
<td>5</td>
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<tr>
<td>• Measurement method standard, valid and reliable</td>
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<td>3</td>
<td>3</td>
</tr>
<tr>
<td>• Exposure measured equally in all study groups</td>
<td>3</td>
<td>-</td>
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<tr>
<td>5. Outcome measures</td>
<td>20</td>
<td>15</td>
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<tr>
<td>• Primary/secondary outcomes clearly defined</td>
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<tr>
<td>• Outcomes assessed blind to exposure or intervention</td>
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<td>-</td>
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<tr>
<td>• Method of outcome assessment standard, valid and reliable</td>
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<td>5</td>
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<tr>
<td>• Length of follow-up adequate for question</td>
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<td>5</td>
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<tr>
<td>6. Statistical Analysis</td>
<td>19</td>
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<tr>
<td>• Statistical tests appropriate</td>
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<tr>
<td>• Multiple comparisons taken into consideration</td>
<td>3</td>
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<td>3</td>
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<tr>
<td>• Modeling and multivariate techniques appropriate</td>
<td>2</td>
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<td>• Power calculation provided</td>
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<td>• Assessment of confounding</td>
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<tr>
<td>• Dose-response assessment if appropriate</td>
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<td>7. Results</td>
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<tr>
<td>• Measure of effect for outcomes and appropriate measure of precision</td>
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<td>5</td>
<td>5</td>
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<tr>
<td>• Adequacy of follow-up for each study group</td>
<td>3</td>
<td>3</td>
<td>3</td>
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<td>8. Discussion</td>
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<tr>
<td>• Conclusions supported by results with possible biases and limitations</td>
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<td>5</td>
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<tr>
<td>taken into consideration</td>
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<tr>
<td>9. Funding or Sponsorship</td>
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<td>5</td>
</tr>
<tr>
<td>• Type and sources of support for study</td>
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</tr>
<tr>
<td>TOTAL SCORE =</td>
<td>100</td>
<td>73</td>
<td>63</td>
</tr>
</tbody>
</table>

Adapted and modified from West S et al. Systems to Rate the Strength of Scientific Evidence, Evidence Report, Technology Assessment No. 47. AHRQ Publication No. 02-E016 (88).
mention was made of at least an 80% relief of pain following a diagnostic block, this study was excluded from the final evaluation.

**Level of Evidence**

The evidence based on the present evaluation is Level III or limited.

**Recommendations**

Due to lack of significant evidence, the recommendation is that intraarticular injections are 2C/very weak recommendation or recommendation not to provide intraarticular injections.

**Lumbar Facet Joint Nerve Blocks**

Our search strategy included 2 randomized trials of facet joint nerve blocks meeting methodologic assessment criteria (140,141). There were no observational studies meeting inclusion criteria with evaluation of lumbar facet joint nerve blocks.

Manchikanti et al (140) in a randomized, double blind controlled trial evaluated the role of lumbar facet joint nerve blocks in managing chronic facet joint pain. The study was conducted to determine the clinical effectiveness of therapeutic local anesthetic lumbar facet joint nerve blocks with or without steroid in managing chronic function-limiting low back pain of facet joint origin. The study included 60 patients in Group I with local anesthetic and 60 patients in Group II with local anesthetic and steroid. The inclusion criteria were based on the positive response to the diagnostic controlled comparative local anesthetic lumbar facet joint blocks. Outcome measures included numeric pain scores, Oswestry Disability Index (ODI), opioid intake, and work status. All outcome assessments were performed at baseline, 3 months, 6 months, and 12 months. The results showed significant improvement with significant pain relief (≥ 50%) and functional improvement (≥ 40%) were observed in 82% and 85% in Group I, with significant pain relief in over 82% of the patients and improvement in functional status in 78% of the patients. Based on the results of the present study, it appears that patients may experience significant pain relief 44 to 45 weeks of one year, requiring approximately 3 to 4 treatments with an average relief of 15 weeks per episode of treatment. While limitations of this study include a lack of placebo control, the study included an active control in a randomized equivalence or non-inferiority controlled trial, and the study met all the criteria with 60 patients in each group with appropriate outcome measurements.

Manchikanti et al (141) in a randomized clinical trial evaluated 200 patients with controlled diagnostic blocks for the presence of facet joint pain. Eighty-four patients, or 42% were determined to have lumbar facet joint mediated pain. These patients were randomly allocated into 2 groups: Group I receiving therapeutic injections with local anesthetic and Sarapin, and Group II receiving therapeutic injections with a mixture of local anesthetic, Sarapin, and methylprednisolone. A total of 73 patients were treated with lumbar facet joint nerve blocks under fluoroscopy. Results showed that patients underwent multiple procedures over a period of 2½ years. The mean number of procedures or interventions was 2.5 ± 0.09 from 1 to 3 months, whereas it was 4 +/- 0.13 for 4 to 6 months, 6.1 ± 0.21 for 7 to 12 months, and 8.4 ± 0.31 for 13 to 32 months. Cumulative significant relief with one to 3 injections was 100% up to 1 to 3 months, 82% for 4 to 6 months, 21% for 7 to 12 months, and 10% after 12 months, with a mean relief of 6.5 ± 0.76 months. There was significant improvement noted in overall health status with improvement not only in pain relief, but also with physical, functional, and psychological status, as well as return-to-work status. Even though this is an RCT, it was not blinded. Further, there was no placebo group.

**Radiofrequency Neurotomy**

One randomized trial (142) and 2 observational studies (145,146) met inclusion criteria with methodologic quality assessment for evidence synthesis.

Nath et al (142) in a randomized control trial of 40 patients with chronic low back pain (20 active and 20 controls) found that the active treatment group showed improvement accompanied by significantly greater improvements in paravertebral tenderness,
Lumbar Facet Joint Interventions

various movements, quality of life, and use of analgesics. The pain relief was however, only monitored for 6 months, as it was felt that patients who received placebo treatment could not be left untreated for longer than 6 months. Bogduk (163) in a “Point of View” editorial on the Nath et al’s study commented that: “Nath et al’s study was the first to use controlled diagnostic blocks to select patients, and the first to use correct technique. He did not select ideal patients, free of comorbidity, with already good function, and no depression. He drew his patients from a pain clinic population. This treatment did not relieve every pain that Dr. Nath’s patients had. Nevertheless, the index pain was relieved, and corroborated by improvements in function.”

Dreyfuss et al (146) reported that 87% of 15 patients obtained at least 60% pain relief 12 months status post radiofrequency denervation, with 60% of the patients achieving at least 90% relief. In addition to stringent inclusion criteria, the authors used 16 gauge electrodes and assessed the efficacy of radiofrequency denervation by performing electromyography of the multifidus muscle. Another flaw that pervades most radiofrequency studies is that sensory stimulation (usually less than 0.5 V) is used to corroborate proximity of the electrode to the targeted medial branch. Whereas sensory stimulation is almost certain to be perceived when the electrode is placed on or adjacent to a neural structure, many patients can perceive concordant sensory stimulation at 0.5V or less even when the electrode is purposefully placed in muscle, as during a sham procedure. An attractive alternative to sensory stimulation is to instead attempt to elicit multifidus muscle contraction, as the same medial branch that innervates the facet joint also innervates this paraspinal muscle. In the 2 studies in which the medial branch was identified by motor stimulation of the multifidus muscle, both reported a positive outcome (146,150).

Gofeld et al (145) evaluated in a large clinical audit, extending from 1991 to 2000, 209 patients, with 174 completing the study. They included only the patients with an appropriate response to comparative double diagnostic blocks. Of the 174 patients with complete data, 55 (31.6%) experienced no benefit from the procedure, 119 patients (68.4%) had good to excellent pain relief lasting from 6 to 24 months. They concluded that proper patient selection and anatomically correct radiofrequency denervation of the lumbar zygapophysial joints provides long-term pain relief in a routine clinical setting.

Characteristics of Excluded Studies

Gallagher et al (147) randomly assigned 41 patients based on their response to diagnostic intraarticular blocks (equivocal or good response) to either sham or true denervation. A statistically significant difference in outcome was observed at one month only between sham and true radiofrequency denervation in those patients who obtained a definitive response to diagnostic blocks. This difference persisted for the duration of the 6 month follow-up.

A prospective, randomized, placebo-controlled double-blind trial (n = 31) was conducted by van Kleef et al (150) to assess the efficacy of percutaneous radiofrequency denervation of the lumbar facet joints in reducing pain, functional disability, and physical impairment. All patients had chronic back pain for at least one year, suggestive of facet joint origin and positive response to diagnostic nerve blocks. Patients in the radiofrequency treatment group (n = 15) received an 80 C radiofrequency lesion of the dorsal ramus of the segmental nerve roots L3, L4, and L5. Patients in the control group (n = 16) underwent the same procedure without use of radiofrequency current. A blinded investigator assessed patients’ physical impairment, rating of pain, the degree of disability, and quality of life. Eight weeks after treatment, there were 10 success patients in the radiofrequency group and 6 in the sham group. Three, 6, and 12 months after treatment there were more success patients in the radiofrequency group compared to the sham group. At 12 months, 7 of 15 patients in the radiofrequency group were judged as successes versus 2 of 16 in the sham group. The authors concluded that radiofrequency results is significant for short-term and long-term alleviation of pain and functional disability in a select group of patients with chronic low back pain. The study was excluded from the final analysis as it utilized a single diagnostic block utilizing lidocaine 1% 0.75 mL and considered patients who had at least 50% pain relief to be eligible for study participation.

Leclaire et al (144) in 2001 conducted a randomized, double-blind placebo-controlled trial to assess the efficacy of RFA for low back pain. Patients included in the study (n = 70) had low back pain of at least 3 months duration and had a good response to intraarticular facet injections. Patients were randomly assigned to receive either RFA under fluoroscopic guidance (n = 36) or the same procedure without denervation (sham procedure) (n = 34). Outcome measures were functional disability as assessed by the
Oswestry and Roland-Morris scales and pain as indicated on a VAS. At 4 weeks, the Roland-Morris score had improved by a mean of 8.4% in the RFA group and 2.2% in the placebo group. At 4 weeks, no significant treatment effect was reflected in the Oswestry score or VAS. At 12 weeks, no functional disability treatment effect was reflected in the Oswestry or Roland Morris scale, and no pain reduction was seen as measured by VAS scores. The authors concluded that, although RFA may provide short-term improvement in functional disability, the efficacy of the treatment has not been established. This study differed from other RFA studies in that a functional inventory was the primary outcome, only VAS scores were used to assess pain, and facet injections were used to identify the affected locations. Other studies used diagnostic nerve blocks to identify affected locations. The Leclaire study invited criticism as it failed to define the study population and had inappropriate diagnostic criteria (use of intraarticular injections to identify patients for radiofrequency neurotomy). Patients were evaluated with a single diagnostic block with 50% pain relief as a criterion standard. They considered any relief of one day duration during a 7-day period following a single diagnostic intraarticular injection as significant. Such an effect may be a result of many factors, including natural sequence. Thus, any results or conclusions based on this study would be erroneous.

van Wijk et al (143) conducted a randomized, double-blind sham lesion-controlled trial to determine the efficacy of radiofrequency facet joint denervation as it is routinely performed. The study was designed to reflect common practice in that although no interventions between trial treatment and 3 months follow-up were performed, further radiofrequency or injection procedures were allowed after this period if the initial treatment did not sufficiently alleviate pain. Patients were randomized to radiofrequency (n = 40) or a sham treatment (n = 41). The primary outcome was determined with a combined outcome measure comprised of VAS, physical activities and analgesic intake, secondary outcome measures were the separate diary parameters, global perceived effect (i.e., complete relief, ≥50% relief, no effect) of pain increase, and SF-36 Quality of Life Questionnaire. There was no difference between the 2 groups in the combined outcome measure or VAS, although both groups showed improvement in VAS scores. The global perceived effect, however, improved in the radiofrequency group. The researchers observed that the lack of improvement in physical function despite reduction in pain scores underlines the need to combine these procedures with subsequent structured rehabilitation programs. The authors concluded that in selected patients, radiofrequency facet denervation appears to be more effective than sham treatment. However, the van Wijk article was excluded from the review in consideration of several weaknesses. The van Wijk study failed to utilize comparative, controlled diagnostic blocks and reasonable pain relief criteria. The article generated 2 letters to the editor (164,165). Bogduk (164) commented that the radiographs published by van Wijk et al (143) indicates not only that the electrodes were placed perpendicular to the target nerves, but also that they were lateral to the actual location of the nerve. In these locations the lesions produced were destined to fail to coagulate the nerves adequately, if at all. Consequently, the study of van Wijk et al (143) amounts to comparing one sham with another. Gofeld (165) also pointed out that uncontrolled single blocks, such as used in the van Wijk study, yields 27% false-positive results. Furthermore, he emphasized that the electrodes were 1) definitely not positioned “parallel the nerves”; 2) placed at the base of the transverse process, and not at the base of the superior articular process, and therefore too lateral from the nerve and 3) too far posteriorly as in the lateral view (that is, on the mamilloaccessory ligament) which insulated medial branches and L5 posterior primary ramus from radiofrequency electrodes. Lastly, the authors had accepted that 22 gauge electrodes with 5-mm active tip could produce insufficient lesion size, but managed to execute lesions using this electrode without position adjustments, thus generating very limited lesions (165).

Level of Evidence

The indicated level of evidence for radiofrequency neurotomy is Level II-2 to II-3 based on one randomized trial (142), and 2 observational studies (145,146).

Recommendations

Based on Guyatt et al’s criteria (114), the recommendation is 1B or C/strong for radiofrequency neurotomy.
With comprehensive review of the literature, this systematic review provides current evidence for the effectiveness of lumbar facet or zygapophysial joint interventions in managing chronic low back pain of facet joint origin. Based on the results of this evaluation, the evidence for diagnostic facet joint blocks utilizing 80% pain relief with controlled diagnostic blocks as a criterion standard is Level I or II-1 based on the USPSTF criteria, utilizing 7 studies which met the inclusion criteria. Based on this evaluation, the prevalence of chronic lumbar facet joint pain related to low back pain is common ranging from 16% to 41% with CI ranging within 9% and 49%. The prevalence of false-positive rate with a single block was studied in 6 controlled diagnostic evaluations with a prevalence ranging from 17% to 49%, with CI ranging within 10% and 59%. A large study (58) showed prevalence of 31%, (95% CI 27%–36%) with a false-positive rate of 27% (95% CI 22%–32%) with a single block. Overall prevalence (n = 1,420) and false-positive rate of all studies is 31% (95% CI; 28% to 33%) and 30% (95% CI; 27% to 33%) respectively.

In contrast to the diagnostic interventions, this systematic review found variable evidence for therapeutic facet joint interventions. There was no evidence available for therapeutic intraarticular lumbar facet joint injections. However, the evidence is moderate to strong for therapeutic facet joint nerve blocks and radiofrequency neurotomy with Level II-1 to Level II-3 with 1B or C/Strong recommendation for both short and long-term improvement of chronic low back pain.

In the included studies, for both therapeutic modalities, patients were selected based on a positive response to controlled diagnostic blocks and 80% pain relief as the criterion standard.

Facet arthrosis has been suggested as a cause of low back pain for decades (29,30). However, the exact source of pain in the facet joints is ambiguous. Theories on the generation of pain range from mechanical alterations to vascular changes and molecular signaling. While disc degeneration can clearly cause low back pain, some patients may not experience pain until degenerative changes in the facet joints alter mechanical alignment sufficiently to produce “articular” low back pain (166). Eubanks et al (30) and others (167,168) concluded that evidence of facet arthrosis appears early and can be linked to the amount of heavy work done before age 20. Indeed, it appears that facet arthrosis starts early, with nearly 60% of adults showing some signs of degenerative changes by the time they reach age 30. After this early rise in arthritic changes, subsequent degeneration appears to steadily increase until the seventh decade when the evidence of arthrosis becomes ubiquitous (30).

A systematic review is defined as, “the application of scientific strategies that limit bias by the systematic assembly, critical appraisal, and synthesis of all relevant studies on a specific topic” (169,170). Systematic reviews are labor intensive and require expertise in both the subject matter and review methods. Thus, expertise in one or other areas is not enough and may lead to inaccurate conclusions in turn leading to inappropriate application of the results (109-113). Thus, this systematic review has provided not only expertise in the subject matter which is crucial, but also knowledge in review methodology. A systematic review differs from a narrative review because a systematic review attempts to minimize bias by the comprehensiveness and reproducibility of the search and selection of articles for review and provides assessment of the methodological quality of the studies (109-113).

In this systematic review, we attempted to answer specific narrow clinical questions in depth – the diagnostic accuracy and validity of facet joint blocks and the level of evidence with recommendation for therapeutic facet joint interventions. A systematic searching, selecting, appraising, interpreting, and summarizing of data from original studies was performed. The study summaries were qualitative and quantitative. In this review we have also searched for other types of integrative evidence including other systematic reviews and cost effectiveness studies.

The rationale behind using 50% pain relief as criteria to proceed to a therapeutic radiofrequency neurotomy was outlined by Schwarzer et al (31) who cite the high incidence of concurrent spinal pathology occurring with lumbar facet joint degeneration as the primary reason. Fujiwara et al (168) found that whereas lumbar degenerative disc disease frequently occurs in absence of lumbar facet joint degeneration, patients with severe lumbar facet joint osteoarthritis virtually always have radiologic evidence of degenerative disc disease and/or other spinal pathology.

Considering that facet joint nerve blocks are inherently nonspecific, even when low volumes are injected under fluoroscopic guidance, a strong case can be made for increasing the criteria to a more stringent 80% pain relief. A study by Dreyfuss et al (171) found that using 0.5 mL low volume facet joint nerve
block using conventional landmarks resulted in con-
trast spread into the epidural space or intervertebral
foramen in 16% of cases, and between the cleavage
plane of the multifidus and longissimus muscles in all
injections. Obviously, if 50% or more relief is used,
the prevalence of lumbar facet joint pain will be-
come much higher than the reported values and will
range at approximately 60 to 70% in the lumbar spine
(1-3,56-58,60,62,65).

Cohen et al (123) emphasized that one reason
that double blocks were not used for their study on
the success of lumbar zygapophysial joint radiofre-
quency denervation as a function of diagnostic block
relief was that the use of controlled blocks was not
cost-effective. Manchikanti et al (172) commented
that the whole concept of single blocks resulting with
50% or more relief followed by radiofrequency de-
nervation creates many questions regarding the reli-
ability of diagnostic blockade, increased health care
costs, and coverage for facet joint nerve blocks and
radiofrequency neurotomy. Schwarzer et al (38) us-
ing 90% relief of pain as a standard showed that the
prevalence of lumbar zygapophysial joint pain in 37%
of patients. The same authors showed a placebo re-
sponse in 32% of the patients receiving normal saline.
Most publications agree that 2 diagnostic blocks must
be performed before radiofrequency denervation and
many payors are requiring 80% or more pain relief.
Consequently, a single block will definitely increase
costs of care as the single diagnostic block will lead
to an increase in number of radiofrequency denerva-
tions, which are more expensive and time consuming.
Cost effectiveness of controlled, comparative, local an-
esthetic facet joint nerve blocks has been evaluated
and found to be superior to an algorithmic approach
starting with discography in axial pain (62).

The most common and worrisome complications
of facet joint interventions are related to needle place-
ment and drug administration. Potential complications
include dural puncture, spinal cord trauma, infection,
intraarterial or intravenous injection, spinal anesthe-
sia, chemical meningitis, neural trauma, pneumotho-
rax, radiation exposure, facet capsule rupture, hema-
toma formation, and steroid side effects (173-182).

Potential side effects with radiofrequency de-
nervation include painful cutaneous dysesthesias, in-
creased pain due to neuritis or neurogenic inflamma-
ation, anesthesia dolorosa, cutaneous hyperesthesia,
pneumothorax, and deafferentation pain. Uninten-
tional damage to a spinal nerve during medial branch
radiofrequency, causing a motor deficit, is also a com-
plication of a neurolytic procedure (183).

**Conclusion**

Diagnostic lumbar facet joint nerve blocks are
safe, valid, and reliable. The strength of evidence for
diagnostic facet joint techniques is Level I or II-1 based
on multiple controlled trials available in the diagno-
sis of spinal pain in the lumbar region.

For therapeutic interventions, the indicated evi-
dence for lumbar facet joint nerve blocks is Level II-
1 or II-2 with a 1B or C/strong recommendation. For
radiofrequency neurotomy, the indicated evidence is
II-2 to II-3 with a recommendation of 1B or 1C. The
evidence for lumbar intraarticular injections is Level
III (limited) with a recommendation of 2C/very weak
recommendation or recommendation not to provide
intraarticular injections.

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