Lower back pain (LBP) is a common problem, which typically resolves with conservative management. In the event that LBP persists, effective treatment requires accurate targeting of the intrinsic cause with appropriate application of specific management options. Major sources of LBP include the facet joints, sacroiliac joints, vertebrae, disks, and muscular spasm.1-4

Initially thought to be inert and insensate, the intervertebral disk is considered a recognized source of LBP.4-5 Concordant pain has been provoked from the annulus, epidural structures, and dura at laminectomy performed under local anesthesia, and innervation of the annulus as well as surrounding structures has been established.1,5-8 Disk challenge by provocation diskography can reproduce a patient’s familiar LBP (diskogenic pain) possibly due to stretch of the injured annulus or ligament, irritation of the adjacent vertebral endplate, or leakage of the inflammatory by-products of disk degeneration.5

In general, MR imaging is most effective at identifying abnormal features of a disk, including altered hydration/matrix, disk morphology, and adjacent inflammatory endplate/vertebral changes and loss of height, but these features do not reliably predict the painful disk.9-12 Studies have suggested that when MR imaging demonstrates a focal high-intensity zone in the posterior margin of the disk, the likelihood of LBP generated by that disk is high.13-16 CT, myelography, and routine radiography can identify abnormal-appearing disks but are also unable to establish which disk space is painful.15,16

Lumbar diskography is a long-standing provocative technique, which, by nature of direct disk stimulation, can identify the painful and concordant disk.5,9-12

A previous report on cervical diskography suggested that the response of a painful disk to local anesthetic can predict the response to cervical disectomy.29 We routinely administer lidocaine to significantly painful lumbar disks during diskography to relieve the induced pain and allow more effective study of subsequently evaluated levels. Although pain relief was frequently observed, in many instances—in particular in contained disks—the patient’s pain was not improved or would worsen with local anesthetic application.

In this study, we retrospectively evaluated the relationship of pain reduction after intradiskal lidocaine administration during provocation lumbar diskography with the presence or absence of diskographic contrast leakage as identified on diskographic images or postdiskographic CT.

**Materials and Methods**

During a 13-month period, 130 consecutive patients had provocation lumbar diskography at our institution by 1 of 2 experienced spine interventional neuroradiologists. Eighty-seven patients were men and 43 were women (average age, 41 years; range, 22–64 years). Pa-
patients typically presented with long-standing severe LBP unresponsive to conservative management and were referred by spine-focused orthopedic surgeons or neurosurgeons for preoperative planning. Severe LBP was accompanied by radiating hip or gluteus pain in 10% and leg pain in 50% of patients. Twenty-eight of 130 patients had undergone prior lumbar surgery.

In 111 of the 130 patients, 1 or more significantly painful disks were encountered, and intradiskal lidocaine was injected to reduce or eliminate the patient’s provoked pain. Nineteen patients were not included (normal diskogram, n = 5; volume limitation to lidocaine injection at all painful levels, n = 5; pain only at cage fused levels, n = 4; lidocaine not administered, n = 2; and uninterpretable provocation response, n = 3).

Provocation diskography was performed at 350 levels in these 111 patients. At 182 disk levels, severe pain was provoked (concordant, 154 levels; nonconcordant, 28 levels) during disk evaluation, and intradiskal lidocaine was injected for pain control. Unoperated (L5-S1, 64; L4-5, 53; L3-4, 24; L2-3, 6; L1-2, 1; T11-T12, 1) and previously operated levels (L5-S1, 16; L4-5, 17) were both studied. These 182 disk levels form the basis of this report. Severe LBP was encountered at a single level in 53 patients, 2 levels in 46 patients, 3 levels in 11 patients, and 4 levels in 1 patient. Leg or hip pain accompanied the provoked LBP in 15% of patients.

Intradiskal lidocaine was not administered in the remaining 168 levels (normal, 126; mild nonconcordant diskomfort, 30; volume limitation to lidocaine injection or extensive venous filling on disk contrast injection, 12).

Institutional review board approval was obtained for this retrospective study.

**Lumbar Diskography Technique**

Diskography was performed in a standard fashion after appropriate informed consent. The patient’s pain history, including leg-versus-back pain contribution, was clarified, and the most-severe-versus-immediate preprocedural pain level was documented using the 0–10 visual analog scale (VAS). Limited intravenous conscious sedation (fentanyl, 0.05 mg; midazolam [Versed]), 1 mg) was given before the procedure with a dose of fentanyl typically given at the end of the examination. Administration of additional fentanyl during the examination was necessary only on rare occasions when disk provocation resulted in extremely severe patient pain that was not responsive to intradiskal anesthetic, and the level of consciousness was never affected. Supplemental administration of additional conscious sedation during the diskography was generally avoided. The lower lumbar region was cleansed and draped, trajectory to target disk space was identified with C-arm fluoroscopy (OEC Series 9800; GE Healthcare, Milwaukee, Wis), and local anesthetic was applied. Disk access was achieved employing a double-needle technique using a 20-gauge guiding spinal needle followed by a curved long 25-gauge spinal needle into the center of the disk space. All needles were placed concordantly opposite the side of leg pain before individual disk injection. Anticipated normal/control disk level was typically studied first.

Disk spaces were studied and provoked by a moderate-to-rapid hand injection of 1.5- to 4.5-mL iohexol, 240 mg iodine/mL, delivered by a 3-mL syringe and a small 0.6-mL-volume connection tube under direct fluoroscopic guidance. Injection volume was dependent on 1) disk-volume end point, 2) clearly established severe pain response, or 3) exaggerated capacity in degenerative disks. Patients were kept unaware of whether a level was being provoked or which level was being studied. Response to injection was initially observed by the operator, and with a positive pain response, the features of the pain were clarified, VAS level of pain was established, and these items were recorded in a manner similar to the technique of Walsh et al.2 Concordant pain was recorded if the provoked pain was the patient’s typical or familiar pain, and nonconcordant pain was recorded if the provoked pain was not the typical pain. Fluoroscopic spot film imaging was obtained for each disk level in anteroposterior and lateral projections during and following the injections.

During disk testing, the patient’s immediate response to injection, response to injection end point (if present), and perception of provoked pain (concordant or discordant) were the primary focus of the diskographer. Syringe disk pressures were not recorded during injection.

If a significantly painful disk space (typically VAS ≥7, concordant or nonconcordant) was encountered, preservative-free lidocaine (2% strength, 1- to 1.5-mL lidocaine [Xylocaine-MPF]) was injected into the disk in an attempt to reduce the patient’s provoked pain and to allow response clarity in subsequently studied lumbar levels.

The patient was routinely questioned regarding any pain reduction; after administration of the intradiskal lidocaine, the response was recorded as either 1) complete or near-complete pain relief, 2) partial pain relief, or 3) no significant pain relief or as a specific VAS-grade reduction from the pain generated by disk provocation relative to the baseline pain, depending on patient’s ability to express the change. Responses initially reported with the VAS were converted to the 3-point scale by calculating the percentage of pain reduction relative to baseline VAS pain level (>66% pain reduction, complete or near-complete pain relief; 33%–66% pain reduction, partial pain relief; and <33% pain reduction, minimal or no significant pain relief).

Postdiskogram CT (GE Healthcare, Milwaukee, Wis) was obtained in all patients immediately following the diskogram employing bone and soft-tissue algorithms with either direct axial 3-mm-section acquisition or with a spiral technique and 3-mm axial and sagittal reformattting with isotropic voxels.

**Imaging and Lidocaine Response Analysis**

Diskographic fluoroscopic images with the postdiskographic CT of all significantly painful lidocaine-treated disk spaces were independently and blindly reviewed by 2 neuroradiologists experienced in lumbar diskography. Identification of epidural diskographic contrast leakage was primarily established by assessment of the fluoroscopic images with secondary inspection and correlation with the postdiskographic CT. Significant leakage at the disk margin around the needle-entry site was classified as a true leak, but minimal contrast identified at needle entry only after needle withdrawal was not classified as leakage. Disks were judged as either “contained” (no contrast leaking from the disk space) or “leaking” (epidural contrast leakage from the disk space due to radial tear, combined annular and radial tear, or annular and nuclear herniation), itemized, and tabulated. Discordant judgments were resolved by consensus.

Imaging identification of diskographic leakage in the significantly painful treated disks was compared with the 3-point response to lidocaine administration: 1) complete or near-complete relief; 2) partial pain relief; and 3) minimal or no significant relief.

**Statistical Analysis**

Statistical analysis was performed by χ² comparison between complete response and no response to lidocaine in contained and leaking disks and a logistic regression model adjusting for the disk state to
compare the 2 diskographers regarding the proportion of responses among patients (SAS release 8.2, SAS Institute, Cary, NC).

Results
The results are summarized in Table 1 and Figs 1–5. Eightytwo of 182 (45%) lidocaine-treated disks were contained disks with 100 (55%) demonstrating diskographic leakage through radial tear, radial and annular tear, or partially herniated annular or nuclear material.

In contained disks, 56 (69%) demonstrated no improvement after intradiskal lidocaine administration (Figs 1 and 2) with 9 (11%) demonstrating partial improvement (Fig 3) and 17 (20%) demonstrating complete or near-complete pain improvement. In leaking disks, 74 (74%) demonstrated complete or near-complete pain relief after intradiskal lidocaine administration (Figs 4 and 5) with 15 (15%) demonstrating partial improvement and 11 (11%) demonstrating no significant improvement.

Overall, 91 of 182 (49%) disks demonstrated complete or near-complete improvement after intradiskal lidocaine administration with 24 (13%) demonstrating partial improvement and 67 (37%) demonstrating no significant improvement. Of disks that demonstrated complete or near-complete improvement, 74 of 91 (82%) were diskographic leaking disks and only 17 (18%) were contained disks. Of disks that demonstrated no significant improvement after intradiskal lido-

![Fig 1](https://example.com/image1)

**Fig 1.** Patient is a 39-year-old man with an injury 20 years ago related to an overturned military vehicle. He has long-standing lower back pain that has recently worsened (VAS score, 8/10). Significant degenerative disk changes are present at L4-5 and L5-S1 on MR imaging, and surgery is now being considered. After 3-mL contrast injection at L5-S1, severe concordant lower back pain was provoked (VAS score, 8/10), with no improvement after 1 mL of intradiskal lidocaine administration.

A. Frontal diskographic fluoroscopic image obtained after contrast injection at L5-S1 demonstrates degenerative changes with nuclear and annular fragmentation (arrow) but no contrast leakage.

B. Axial postdiskographic CT image obtained at L5-S1 demonstrates a contained radial tear with focal disk protrusion and focal contrast accumulation without epidural leakage (arrow).

![Fig 2](https://example.com/image2)

**Fig 2.** Patient is a 36-year-old man with long-standing back pain and left leg pain (VAS score, 8/10), who had prior diskectomy at L4-5, with only limited improvement. Diskography was requested for follow-up assessment. After 2-mL contrast injection at L5-S1, severe nonconcordant pain was provoked (VAS score, 8/10) with no improvement after 1 mL of intradiskal lidocaine administration. His pain did reproduce concordantly at the diskectomy level.

A. Outside MR sagittal T2-weighted image demonstrates significant degenerative disk changes at L4-5 and L5-S1 (arrowheads) with a high-intensity zone noted along the posterior L5-S1 disk margin (arrow).

B. Lateral diskographic fluoroscopic image obtained after contrast injection at L5-S1 demonstrates degenerative changes (arrow), without evidence of contrast leakage. Contrast dilution was noted throughout the disk after lidocaine administration (not shown), but provoked pain did not improve.

C. Axial postdiskographic CT image obtained at L5-S1 demonstrates a contained radial tear with focal disk protrusion and focal contrast accumulation without epidural leakage (arrow).
caine administration, 56 of 67 (84%) were contained disks and 11 (16%) were diskographic leaking disks. In the 58 patients with 2 or more significantly painful levels, 33 (57%) had a combination of both leaking and contained painful disks.

Assessment between complete relief versus no significant improvement after intradiskal lidocaine in leaking disks versus contained disks (Table 1, data columns 1 and 3) was statistically significant ($\chi^2, P < .001$). Interinterpreter difference was not detected by logistic regression ($P = .31$).

Results separating postoperative and nonpostoperative disks are summarized in Tables 2 and 3. Leaking disks demonstrated complete or near-complete improvement after lidocaine administration more frequently than contained disks in both postoperative levels ($P = .009$) and nonpostoperative levels ($P < .001$), similar to that of the overall study population.

Regarding postoperative disks, 17 of 33 (51.5%) were contained disks, and 16 were leaking disks (48.5%). A greater frac-
tion of painful postoperative disks demonstrated no significant improvement after intradiskal lidocaine administration (17 of 33 [51.5%], Table 2). This trend was seen in both leaking (4 of 16 [25%]) and contained (13 of 17 [76.5%]) postoperative disks. Results in the nonoperative disks paralleled the overall patient results (Tables 1 and 3).

Discussion

LBP is a significant and expensive health care problem. Sources of LBP include the facet, sacroiliac joints, intervertebral disk, vertebral injury or destruction, muscle spasm, and instability. Clearly, accurate targeting of the cause of LBP is crucial to proper application of appropriate treatment options.

Although this finding was initially controversial, the intervertebral disk is recognized as a source of LBP, with several different mechanisms (stretch of the annulus, pressurization of the endplate, and chemical irritation) considered potentially responsible.5 Free nerve endings, presumably pain fibers, have been identified in the peripheral annulus and posterior longitudinal ligament (PLL) as well as in the adjacent dura, epidural vessels, and anterior longitudinal ligament.30-38 Probing or stimulation of clinically painful disks clearly provokes a painful response when tested during laminectomy with the patient under local anesthesia.6-8 Familiar pain can be reproduced in contained disks during provocation diskography, likely related to stretch of the annulus or stimulation of the adjacent vertebral endplates.5,38 Disk material can also induce inflammation and chemical irritation; nociceptive stimulation could develop at the annular/ligamentous margin or epidural/dural structures secondary to leakage of degenerative disk by-products.40-48

Although imaging can identify abnormal-appearing disk spaces and help point to the cause of LBP or radiculopathy, provocative diskography is critical for confirming a painful disk and in particular for identifying the concordant painful disk levels in an individual patient.

Our results demonstrate several important observations with regard to the painful lumbar disk: 1) Most painful contained disks are unresponsive to intradiskal lidocaine administration, 2) some painful contained disks respond to intradiskal lidocaine, demonstrating partial or complete pain resolution, 3) most painful disks with diskographic leakage demonstrate complete or near-complete improvement after intradiskal lidocaine administration, and 4) some painful leaking disks demonstrate only partial or no pain improvement after intradiskal lidocaine. Pain improvement after lumbar intradiskal anesthetic administration has been occasionally mentioned in the diskography literature but, to our knowledge, never comprehensively assessed.49,50 These observations may suggest several different and frequently overlapping causes of diskogenic LBP.

Contained Painful Disks: No Relief

In 69% of painful contained disks, intradiskal lidocaine administration resulted in no significant improvement in their provoked pain (Figs 1 and 2). Despite the degenerative fluoroscopic and postdiskographic CT imaging appearances, pain relief was not achieved when these internal deranged surfaces (disk degeneration, annular degeneration, and annular tears) were bathed with local anesthetic. LBP in these disks is likely generated peripheral to the internal disk surfaces, perhaps in the peripheral annulus or posterior longitudinal ligament. One could question, in these instances, whether peripheral annular injury was present and not recognizable at imaging.

Contained Painful Disks: Partial or Complete Relief

Complete pain relief was seen in 20% of painful contained disks, and partial pain relief occurred in 11% (Fig 3). This observation strongly suggests that the visualized internal degenerative features are related to pain provocation for that disk level. The existence of pain nociceptive fibers in the disk has been long debated. Free nerve endings suggesting pain fibers have been documented in the peripheral one third of the annulus and PLL, most commonly associated with small penetrating blood vessels.1,31,34,35 Free nerve endings have been observed extending as far as the peripheral half of the annulus in patients with disk herniation.32 Neovascularization and substance P immunoreactive innervation (nociceptive pain-mediator fibers) of the inner one third of the annulus (46%) and nucleus (22%) have been described in operative biopsy of painful disk levels52 and recently reconfirmed with calcitonin gene-related peptide-expressing nociceptor innervation demonstration in the outer and inner annulus of postoperative specimens.53 Local anesthetic action on these annular or nuclear pain fibers could be responsible for the pain resolution or improvement noted in this subset of our patients with painful contained disks.

Painful Disks with Diskographic Leakage: Complete Response

In 74% of painful leaking disks, complete or near-complete resolution of LBP occurred after intradiskal lidocaine administration (Figs 4 and 5). In these lumbar disks, local anesthetic was either reaching the location of the LBP generator or blocking its central neural transmission.

LBP in these patients could be generated at the annular/PLL surface at the radial tear or herniation. Alternatively, irritation of epidural structures (vessels) or dura could be the source of pain and relieved by the local anesthetic. It is well known to diskographers that leaking disks can reproduce severe and concordant LBP after only minimal contrast injection or leakage. Patients are also encountered in whom non-

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specific/semiconcordant LBP is provoked with initial disk injection, but exact replication is recognized only when greater contrast volume and pressure are used and focal contrast leakage finally occurs.

The irritant effects and by-products of disk degeneration such as phospholipase A2, substance P, inflammatory cells, and hydrogen ions have been suggested as a source of LBP and radiculopathy.40-48 Unmyelinated fibers (somatic and autonomic) are known to be present in the peripheral annulus/PLL but are also encountered in epidural blood vessels and the ventral dura. Radiating pain patterns (radiculopathy) at times resemble a nerve root distribution and at times are more distinct, variable, or referred, perhaps reflecting the distribution of the somatic (somatic and autonomic innervation including dermatome, myotome, and sclerotome), and this distribution has been termed the “radiating zones of head.”2 A vascular pain fiber (epidural, paravertebral vessel, or dural) source of pain fiber irritation and LBP might explain a portion of the response to lidocaine in painful leaking lumbar disks.

Painful Disks with Diskographic Leakage: Partial Response or No Response

The fourth important observation is related to the absent improvement (11%) or partial improvement (15%) to local anesthetic noted in the painful leaking disks. Considering the response of contained disks, it is likely that some leaking disks also have internal derangement or peripheral annular injury as the source of their pain, which does not come in contact with the intradiskal local anesthetic. Absent pain improvement (11%) despite leakage suggests that the leak pathway and leaking substances may not be the pain generator and that pain is being produced elsewhere in the disk. Partial pain improvement (15%) suggests that the leak pathway and leaking products may partially contribute.

Pain origin in leaking disks, therefore, could be related to the visualized internal derangement, peripheral annular margin injury, leakage path, or irritant effects of epidural leakage of inflammatory material. Although a simple epidural anesthetic mechanism of pain extinction could be suggested for relief in the leaking disks, clearly complete pain ablation is not encountered in all instances. In addition, in many leaking disks, the leak is small and focal, and widespread leakage of contrast (and presumably lidocaine) is not present.

We have not embraced injection-pressure measurement during provocation lumbar diskography, despite familiarity with the technique. In our experience, pressure measurements have rendered limited insight into confirmation of or characterization of (concordant-versus-discordant) objectively provoked diskogenically painful beyond a perceived end point to injection. This result is in part supported by the lack of clear postoperative improvement when pressure-controlled diskography is used in patient selection.34-56 Of interest, a statistically significant difference in surgical outcome between discectomy/intradiskal fusion in low-pressure (leaking?) disks compared with fusion only in high-pressure (contained non-leaking?) disks has been demonstrated in a subset comparison within 1 study.56

Several limitations are present in this retrospective study, despite the large number of patients and disks evaluated. In contained disks, internal disk derangement is complex, flow of injected material is unpredictable, and uniform spread of the injected anesthetic may not occur, leading to a reduced rate of improvement in these disks. In leaking disks, early epidural contrast reabsorption may lead to under-recognition in subtle cases with small leaks, not originally appreciated at fluoroscopy. Second, many leaking disks cannot be completely distended, and areas of annular injury may not be fully tested with anesthetic or adequately provoked without more aggressive biomechanical strain. Third, anesthetic leakage, if excessive, could potentially mask the response at another level, but in our experience, adjacent levels can be correctly tested after anesthetic administration.

Conclusion

The provoked pain response of lumbar disks to intradiskal lidocaine administration was different for disks demonstrating diskographic leakage versus disks without contrast leakage. This observation suggests that the primary pain generator related to the disk is in the outer annulus or extrinsic to the disk. In painful disks without diskographic leakage, though some patients improved after anesthetic injection, most did not, and it is possible that injected lidocaine simply cannot reach the appropriate nerves. These observations suggest more than 1 cause of diskogenic LBP, and these causes may overlap in the patient or at an individual disk level.

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References

1. Bogduk N. The innervation of the lumbar spine. Spine 1983;8:286–93
2. Jinkins RJ. The pathoanatomic basis of somatic and autonomic syndromes originating in the lumbosacral spine. Neuroimaging Clin N Am 1993;3:443–63
3. Schneiderman G, Flannigan B, Kingston S, et al. Magnetic resonance imaging in the diagnosis of disk degeneration: correlation with diskography. Spine 1992;17(6 Suppl):S164–71
4. Kuslich SD, Ulstrom CL, Michael CJ. The tissue origin of low back pain and sciatica: a report of pain response to tissue stimulation during operations on the lumbar spine using local anesthetics. Orthop Clin North Am 1991;22:181–87
5. Guyer RD, Ohnmeiss DD. Lumbar diskography: position statement from the North American Spine Society Diagnostic and Therapeutic Committee. Spine 1995;20:2048–59
6. Fakoner MA, McGeorge M, Begg C. Observations on the cause and mechanism of symptom-production in sciatia and low back pain. J Neurol Neurosurg Psychiatry 1948;11:13–26
7. Smyth MB, Wright V. Sciatica and the intervertebral disk: an experimental study. J Bone Joint Surg 1958;40-A:1401–18
8. Kuslich SD, Ulstrom CL, Michael CJ. The tissue origin of low back pain and sciatica: a report of pain response to tissue stimulation during operations on the lumbar spine using local anesthetics. Orthop Clin North Am 1991;22:181–87
9. Schneiderman G, Flannigan B, Kingston S, et al. Magnetic resonance imaging in the diagnosis of disk degeneration: correlation with diskography. Spine 1987;12:276–81
10. Zucherman J, Derby R, Hsu K, et al. Normal magnetic resonance imaging with abnormal diskography. Spine 1988;13:1335–59
11. Gunzburg R, Parkinson R, Moore R, et al. A cadaveric study comparing diskography, magnetic resonance imaging, histology, and mechanical behavior of the human lumbar disk. Spine 1992;17:417–26
12. Horton WC, Daftari TK. Which disk as visualized by magnetic resonance imaging is actually a source of pain? A correlation between magnetic resonance imaging and diskography. Spine 1992;17:6 Suppl:S164–71
13. Aprill C, Bogduk N. High-intensity zone: a diagnostic sign of painful lumbar disk on magnetic resonance imaging. Br J Radiol. 1992;65:361–69
14. Schellhas KP, Pollei SR, Gundy CR, et al. Lumbar disk high-intensity zone: correlation of magnetic resonance imaging and diskography. Spine 1996;21:79–86
15. Grubb SA, Lipscomb HJ, Guilford WB. The relative value of lumbar roentgenograms, metrizamide myelography, and diskography in the assessment of patients with chronic low-back syndrome. Spine 1987;12:282–86
34. Bogduk N, Tynan W, Wilson AS.

31. Malinsky J.

33. Edgar MA, Ghadially JA.

32. Jackson HC 2nd, Winkelmann RK, Bickel WH.

20. Sachs BL, Vanharanta H, Spivey MA, et al.

35. Korkala O, Gronblad M, Liesi P, et al. Immunohistochemical demonstration of nociceptors in the ligamentous structures of the lumbar spine. Spine 1985;10:156–57

36. Groen GJ, Baljet B, Drukker J. The innervation of the spinal dura mater: anatomy and clinical implications. Acta Neurochir (Wien) 1988;92:39–46

37. Groen GJ, Baljet B, Drukker J. Nerves and nerve pleures of the human vertebral column. Acta Anat 1990;138:282–96

21. McFadden JW.

17. Simmons EH, Segil CM. An evaluation of diskography in the localization of symptomatic levels in diskogenic disease of the spine. Clin Orthop Relat Res 1975;108:57–69

19. Viderman T, Malimivaara A, Mooney V. The value of the axial view in assessing diskograms: an experimental study with cadavers. Spine 1987;12:299–304

29. Roth DA.

28. Moneta GB, Videman T, Kaivanto K, et al. Lumbar diskography followed by computed tomography: reevaluation of inflammatory cells in disk herniation tissue. Spine 1990;15:690–707

24. Weinstein J, Claverie W, Gibson S. The pain of diskography. Spine 1988;13:1344–48

27. Walsh TR, Weinstein JN, Spratt KF, et al. Lumbar diskography in normal subjects: a controlled, prospective study. J Bone Joint Surg Am 1990;72:1081–88

23. Yasuma T, Ohno R, Yamauchi Y. False-negative lumbar diskograms: correlation of diskographic and histological findings in postmortem and surgical specimens. J Bone Joint Surg Am 1988;70:1279–90

26. Bernard TN Jr. Lumbar diskography followed by computed tomography: refining the diagnosis of low-back pain. Spine 1990;15:690–707

25. Vanharanta H, Guyer RD, Ohnmeiss DD, et al. Autologous nucleus pulposus induces neurophysiologic and histologic changes in porcine cauda equina nerve roots. Spine 1993;18:1425–32

20. McFadden JW, McCall IW, Williams L, et al. Provocation diskography as a guide to planning operations on the spine. J Bone Joint Surg Br 1988;70:267–71

22. Colhoun E, McCullough K, Williams L, et al. Incision of the annulus fibrosus induces nerve root morphologic, vascular, and functional changes: an experimental study. Spine 1996;21:2539–43

19. Viderman T, Malimivaara A, Mooney V. The value of the axial view in assessing diskograms: an experimental study with cadavers. Spine 1987;12:299–304

20. Sachs BL, Vanharanta H, Spivey MA, et al. Dallas diskogram description: a new classification of CT/diskography in low-back disorders. Spine 1987;12:287–94

21. McFadden JW.

23. Yasuma T, Ohno R, Yamauchi Y. False-negative lumbar diskograms: correlation of diskographic and histological findings in postmortem and surgical specimens. J Bone Joint Surg Am 1988;70:1279–90

26. Bernard TN Jr. Lumbar diskography followed by computed tomography: refining the diagnosis of low-back pain. Spine 1990;15:690–707

27. Walsh TR, Weinstein JN, Spratt KF, et al. Lumbar diskography in normal subjects: a controlled, prospective study. J Bone Joint Surg Am 1990;72:1081–88

28. Moneta GB, Viderman T, Kaivanto K, et al. Reported pain during lumbar diskography as a function of annular ruptures and disk degeneration: a re-analysis of 833 diskograms. Spine 1994;19:5688–74

29. Roth DA. Cervical analgesic diskography: a new test for the definitive diagnosis of cervical radiculopathy. Spine 1990;15:690–707

30. Pedersen HE, Blunck CF, Gardner E. The anatomy of lumbosacral posterior rami and meningeal branches of spinal nerve (sinu-vertebral nerves); with an experimental study of their functions. J Bone Joint Surg Am 1988;70:1279–90

31. Malinsky J. The ontogenic development of nerve terminations in the intervertebral disks of man. Acta Anat (Basel) 1959;38:96–113

32. Jackson HC 2nd, Winkelmann RK, Bickel WH. Nerve endings in the human lumbar spinal column and related structures. J Bone Joint Surg Am 1966;48:1272–81

33. Edgar MA, Ghadially JA. Innervation of the lumbar spine. Clin Ortho Relat Res 1976;115:35–41

34. Bogduk N, Tynan W, Wilson AS. The nerve supply to the human lumbar intervertebral disks. J Anat 1981;132(Pt 1):39–56

35. Korkala O, Gronblad M, Liesi P, et al. Immunohistochemical demonstration of nociceptors in the ligamentous structures of the lumbar spine. Spine 1985;10:156–57

36. Groen GJ, Baljet B, Drukker J. The innervation of the spinal dura mater: anatomy and clinical implications. Acta Neurochir (Wien) 1988;92:39–46

37. Groen GJ, Baljet B, Drukker J. Nerves and nerve pleures of the human vertebral column. Acta Anat 1990;138:282–96

38. Konttinen YT, Gronblad M, Antti-Poika I, et al. Neuroimmunohistochemical analysis of peridiscal nociceptive neural elements. Spine 1990;15:383–86

39. Bogduk N, Modic MT. Controversy in lumbar diskography. Spine 1996;21:402–04

40. Olmarker K, Rydevik B, Nordborg C. Autoagous nucleus pulposus induces neurophysiologic and histologic changes in porcine cauda equina nerve roots. Spine 1993;18:1425–32

41. Groenblad M, Virri J, Tolonen J, et al. A controlled immunohistochemical study of inflammatory cells in disk herniation tissue. Spine 1994;19:2744–51

42. Saal JS. The role of inflammation in lumbar pain. Spine 1995;20:1821–27

43. Olmarker K, Blomquist J, Stromberg J, et al. Inflammagoteric properties of nucleus pulposus. Spine 1995;20:665–69

44. Kayama S, Konno S, Olmarker K, et al. Incision of the annulus fibrosus induces nerve root morphologic, vascular, and functional changes: an experimental study. Spine 1996;21:2539–43

45. Haro H, Shinomiya K, Komori H, et al. Upregulated expression of chemokines in herniated nucleus pulposus resorption. Spine 1996;21:1647–52

46. O’Donnell JL, O’Donnell AL. Prostaglandin E2 content in herniated lumbar disk disease. Spine 1996;21:1653–55

47. Kang ID, Stefanovic-Racic M, McIntyre LA, et al. Toward a biochemical understanding of human intervertebral disk degeneration and herniations: contributions of nitric oxide, interleukins, prostaglandin E2, and matrix metalloproteinases. Spine 1997;22:1065–73

48. Weiler C, Andreas GN, Bachmeier BE, et al. Expression and distribution of tumor necrosis factor alpha in human lumbar intervertebral disks: a study in surgical specimens and autopsy controls. Spine 2005;30:44–54

49. Park WM, McCullough K, Brien JP, et al. Fissuring of the posterior annulus fibrosus in the lumbar spine. Br J Radiol 1997;52:382–87

50. Jaffray D, O’Brien JP. Isolated intervertebral disk resorption: a source of mechanical and inflammatory back pain? Spine 1986;11:397–401

51. Yoshizawa H, O’Brien JP, Smith WT, et al. The neuropathology of intervertebral disks removed for low back pain. J Pathol 1980;102:95–104

52. Freemont AJ, Pepperock TE, Goupille P, et al. Nerve ingrowth into diseased intervertebral disk in chronic back pain. Lancet 1997;350:178–81

53. Ozawa T, Ohbira S, Isome G, et al. The degenerated lumbar intervertebral disk is innervated primarily by peptide-containing sensory nerve fibers in humans. Spine 2006;31:2418–22

54. Madan S, Gundanna M, Harley JM, et al. Does provocative diskography screening of diskogenic back pain improve surgical outcome? J Spinal Disord Tech 2002;15:243–51

55. Carragee EJ, Linclon T, Parmar VS, et al. A gold standard evaluation of the “diskogenic pain” diagnosis as determined by provocative diskography. Spine 2006;31:2115–23

56. Derby R, Howard MW, Grant JM, et al. The ability of pressure-controlled diskography to predict surgical and nonsurgical outcomes. Spine 1999;24:364–71