Clinical Article

Clinical and Radiological Findings of Discogenic Low Back Pain Confirmed by Automated Pressure-Controlled Discography

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Objective: Few studies on the clinical spectrum of automated pressure-controlled discography (APCD)-defined positive discs have been reported to date. Thus, the present study was undertaken to analyze clinical parameters critical for diagnosis of discogenic pain and to correlate imaging findings with intradiscal pressures and pain responses in patients with APCD-positive discs.

Methods: Twenty-three patients who showed APCD-positive discs were selected for analysis. CT discogram findings and the degrees of nuclear degeneration seen on MRI were analyzed in comparison to changes of intradiscal pressure that provoked pain responses; and clinical pain patterns and dynamic factors were evaluated in relation to pain provocation.

Results: Low back pain (LBP), usually centralized, with diffuse leg pain was the most frequently reported pattern of pain in these patients. Overall, LBP was most commonly induced by sitting posture, however, standing was highly correlated with L5/S1 disc lesions ($p < 0.01$). MRI abnormalities were statistically correlated with grading of CT discogram results ($p < 0.05$); with most pain response observed in CT discogram Grades 3 and 4. Pain-provoking pressure was not statistically correlated with MRI grading. However, it was higher in Grade 3 than Grade 4.

Conclusion: APCD-positive discs were demonstrated in patients reporting centralized low back pain with diffuse leg pain, aggravated by sitting and standing. MRI was helpful to assess the degree of nuclear degeneration, yet it could not guarantee exact localization of the painful discs. APCD was considered to be more useful than conventional discography for diagnosis of discogenic pain.

KEY WORDS: Low back pain · Intervertebral disc · Discography · MRI · Discogenic pain.

INTRODUCTION

Lumbar disc pathology has long been recognized as one of the main causes of chronic low back pain (CLBP)1,2). Recent reports indicate CLBP can be caused by structure-specific etiologies including zygapophyseal joint abnormality, disc pathology, and sacroiliac joint arthropathy; and that about 26–40% of CLBP patients were known to have discogenic pain3,4,13,17). Recent development of high-resolution neuroimaging methods has facilitated diagnostic work-up for discogenic pain. This has been especially effective in the case of herniated nucleus pulposus and degenerated disc pathology if patients’ symptoms and signs are closely correlated with neuroimaging findings. However, even modern neuroimaging equipment such as magnetic resonance imaging (MRI) and computerized tomography (CT) do not reach the level of resolution needed to visualize great detail and tiny abnormalities of spinal structures. Thus, for example, the diagnosis of internal disc disruption (IDD) which contains tiny fissures in the annulus pulposus may be missed. Furthermore, as shown by Boos et al.5,6), neuroimaging abnormalities are not often correlated with pain source in patients with spinal pain.

In early days of discography, it was performed to demonstrate disc morphology and to diagnose disc herniation in CLBP. However, this technique is no longer used for diagnosing disc herniation and result do not correlate well with morphology of intradiscal degeneration21). Instead, provocation discography often referred as “disc stimulation” is currently used to stimulate individual “painful discs” to deter-
mine whether they are sources of patients' spinal pain\textsuperscript{2,25}). In modern provoked discography, slow increase of intradiscal pressure by injecting contrast media into the nucleus pulposus can produce patient's accustomed pain if the disc is painful, while stimulation of normal disc does not produce any pain. Provocation discography has been recognized to be a very specific diagnostic test for discogenic pain. However, Seo et al.\textsuperscript{23} demonstrated that there is a difference between static pressure and real time dynamic pressure if the injection speed is fast. This pressure discrepancy makes it difficult to obtain a uniform result and interpretation of the discography procedure. An automated discography device can accurately control the speed of contrast medium injection and simultaneously display the peak pressure. Therefore, it is known to decrease potential errors produced by inexperienced discographers. Further, it is assumed that automated discography devices such as APCD system will allow more accurate isolation of the subgroups of positive or negative discs. The purpose of this study was to find characteristic clinical parameters associated with pain-aggravating factors using automated discography, and to determine the relationship of MRI and CT discogram findings with intradiscal pressure and pain responses thus observed.

**MATERIALS AND METHODS**

**Patient population**

A total of 24 discs showing positive response using automated discography in 23 patients were collected from a consecutive series of 65 patients. A positive disc was defined in accordance with the criteria of International Association for the Study of Pain; \( \geq 6/10 \) concordant pain elicited at \( \leq 50 \) psi above opening pressure with at least one painless control level\textsuperscript{18}. Patients who did not meet these criteria were not included. All patients had more than 12 weeks of unremitting LBP despite appropriate conservative management. Inclusion criteria were as follows: 1) only midline LBP, 2) midline LBP with leg pain, 3) LBP confirmed not to have pain source in zygapophyseal joint or nerve root, 4) no obvious evidence of herniation and extrusion of disc on neuroimaging with corresponding neurological signs.

**Provocation discography**

Patients underwent discography in the operating room under guidance of a C-arm image intensifier (Series 9600, GE OEC medical systems, UT, USA). Discography injection needles were placed according to standard technique, with at least one control disc tested in addition to symptomatic discs\textsuperscript{9}. The discography needle was connected to the APCD system\textsuperscript{°} (Cybermedic Corp, Iksan, Korea) (Fig. 1). The APCD system was equipped with a programmable variable-speed syringe pump capable of pressure up to 150 psi, pressure sensors, and a data acquisition box. It included a computer system loaded with software programmed to display pressure changes throughout the time of injection, and pressure/volume curves\textsuperscript{26}. Non-ionic contrast medium (Iopamiro, Ilsung Pharmaceuticals, Seoul, Korea) was injected into each disc using a motorized syringe of the APCD system. Injection speed was held constant at 0.01 cc/sec to avoid any pressure discrepancy between syringeal pressure and intradiscal pressure. Opening pressure was set when leakage of contrast media from the tip of the needle was first seen under fluoroscopic observation. Thereafter, actual pressure (syringeal pressure minus opening pressure) was continuously displayed on the screen. The patient was asked to describe the nature, distribution, and intensity of pain on a 0-10 visual analog scale.

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**Fig. 1.** Automated discography (APCD system). Note the display of pressure/volume curve (lower graph) and change of visual analog scale (upper graph) on the monitor screen (left), and device equipped with motorized syringe pump (right).
scale (VAS) when he or she subjectively perceived pain. Whenever there were changes of VAS, they were registered via keypad by the patient or the operator. Thus, all pain responses along the injection timeline were displayed and monitored. Injection was continued until one of the following end points was reached: 1) pain ≥ 7 VAS; 2) intradiscal pressure ≥ 70 psi; or 3) injection volume ≥ 3.5 mL. After the procedure, patients were moved immediately to the CT Room (SOMATOM Sensation 4, Siemens, Malvern, PA, U.S.A.) where a post-discography CT scan was performed. Approximately, three to four slices at 5-mm thickness and 4-mm intervals were checked at each disc level in axial view with simultaneous reconstruction of sagittal views.

Positive response of discography
A discography-positive disc was defined as one where concordant pain was produced with an intensity ≥ 6/10, and abnormal morphology on CT discogram was observed, while a control disc injection at another level produced no pain.

Evaluation of pain patterns
All patients were interviewed by a pain-specialized nurse who recorded pain distribution, nature of pain and pain-related dynamic factors. Drawings of pain distribution patterns were completed for all patients before discography, and subsequently the patterns were classified into four types (Fig. 2). Type A shows pain distribution in the lower back only; Type B shows the combination of A and posterior leg pain extending above the knee joint. If the posterior leg pain extended to the ankle, in combination with Type A, it was defined as Type C. Type D shows anterior leg pain, with or without associated back pain. In addition, “dynamic factors” which can aggravate or induce pain were evaluated. Dynamic factors include flexion and extension, and lying, sitting, standing, walking and transitional movement.

Analysis of imaging findings
MRI results from twenty patients were available for analysis. Grading of disc degeneration was classified on the basis of structure, distinction of nucleus and annulus, signal intensity, and height of intervertebral disc according to Pfirrmann et al. All the MRI results were evaluated by a neuroradiologist who was blinded to the results of discography. Imaging of post-discography CT was used to determine the direction of fissure as well as to grade the extent of annular disruption according to the modified Dallas Discogram Classification (Fig. 3); grade 1 fissures reach the inner third of the annulus; grade 2 and 3 fissures reaches the middle third and outer third, respectively; grade 4 indicates the circumferential spread of fissures is greater than 30 degree; grade 5 describes the complete rupture of Grade 3 or grade 4 fissures.

Fig. 2. Pain distribution patterns of discogenic pain. Type A shows pain distribution in the lower back only; Type B shows the combination of A and posterior leg pain. Type C indicates the combination of Type A and posterior leg pain extending just above the knee joint. Type D demonstrates anterior leg pain, with or without associated back pain.

Fig. 3. Demonstration of magnetic resonance image (MRI) and computed tomography discogram grades sampled from the patients. Note the disc degenerations without hemiations in MRI. Extensions of annular tears are demonstrated in conjunction with MRI grades in Discogram-computed tomography findings.
Statistical analysis

Statistical analyses were performed using SPSS 11.0 software (SPSS, Chicago, IL, USA). Statistical analysis for MRI and Dallas discogram grade, intradiscal pressure, and pain responses was performed with ANOVA. All differences were regarded as significant if \( p < 0.05 \).

RESULTS

Demographic profile

The demographic characteristics of 14 male and 9 female patients are summarized in Table 1. The average age of males was 33.36 ± 13.73 (range 22-63 years) and that of females was 43.33 ± 14.0 years (range 20-58 years). Average duration of pain before discography among all the patients was 52.93 ± 61.04 months. The total number of discs tested was 63, and concordant pain responses were observed in 24 discs in 23 patients because one patient showed concordant pain responses in two levels. There were one positive disc in L1/L2, two in L3/L4, nine in L4/L5, and twelve in L5/S1 intervertebral level.

Clinical parameters

Analysis of pain patterns in APCD-positive patients revealed 16 (60%) with Type C pattern, 4 (17%) with Type A, 2 (9%) with Type B, and 1 (5%) patient with Type D pain pattern. Low back pain in these patients showed a tendency of centralization around midline and leg pain was diffuse. Neurological examination disclosed no neurological deficit. Among dynamic factors, pain was most commonly induced by sitting posture (91%), followed by standing (65%), flexion (61%), and walking (52%) (Table 2). Standing was quite closely correlated with L5-S1 disc lesions \( (p < 0.01) \), compared to other levels of disc lesions. In contrast, lying and transitional movement were dynamic factors least correlated with pain production all the lumbar levels of disc lesions.

MRI and CT discogram findings

Table 3 summarizes the morphological MRI grading and CT discogram results. MRI and CT discograms showed relatively concordant grading in grades 2 and 4. However, they did not in grade 3 and 5. The grade of CT discogram was higher in more-degenerated discs. MRI abnormalities were statistically correlated with grading of CT discogram \( (p < 0.05) \). However, other CT discogram findings (including direction and angle of annulus fissures) were not correlated with pain responses and pain patterns.

Intradiscal pressure and pain response in relation with imaging abnormality

Average opening pressure was 15.98 psi and average pressure to provoke pain was 33.63 psi. Opening pressure was not correlated with degree of nuclear degeneration shown in MRI. Although most pain response was observed in grade 3 (30%) and grade 4 (65%), pain response was not statistically correlated with MRI grading. Pressure to provoke the pain response was also not statistically correlated with MRI grading, however, the pain-provoking pressure was higher in grade 3 than grade 4.

Complications after discography

There were no discography-related infection or hemorrhage seen in any of the patients. Although pain aggravation after discography was frequently noted, it was easily controlled with medication and disappeared within 48 hours.

DISCUSSION

Although discogenic pain is well known to clinicians dealing with back pain, it is very difficult to diagnose because it lacks characteristic clinical findings by which reliable clinical diagnosis can be made. Recently, several studies have postulated that discogenic pain is induced by stimulation of a painful disc[1,2,27]. When annular fissures extend into the boundary of a disc, branches of sinuvertebral nerve have

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Table 1. Demographic profile

| Characteristic          | Value          |
|-------------------------|----------------|
| Sex (M : F)             | 14 : 9         |
| Age in years, mean ± SD (range) |               |
| Male                    | 33.36 ± 13.73 (22-63) |
| Female                  | 43.33 ± 14.00 (20-58) |
| Mean duration of pain (mo) | 52.93 ± 61.04  |
| No of disc level tested | 63             |
| No of discs tested per person | 2.74 ± 0.54   |
| No of positive disc level | 24 (38%)      |

Table 2. Frequency of dynamic factors aggravating discogenic pain depending on the levels

| Dynamic factor       | L1/2 | L3/4 | L4/5 | L5/S1 | Total (%) |
|----------------------|------|------|------|-------|-----------|
| Flexion              | 0    | 1    | 4    | 9     | 14 (61)   |
| Extension            | 0    | 1    | 4    | 2     | 7 (30)    |
| Transitional movement| 1    | 1    | 2    | 4     | 8 (35)    |
| Lying                | 1    | 1    | 4    | 3     | 9 (39)    |
| Sitting              | 1    | 2    | 9    | 9     | 21 (91)   |
| Standing             | 0    | 1    | 3    | 11*   | 15 (65)   |
| Walking              | 1    | 2    | 2    | 7     | 12 (52)   |

*p-value < 0.01
been observed growing inside, possibly into nucleus pulposus. Mechanical pressure due to long standing or sitting or injection of contrast medium into the nucleus pulposus can stimulate the sensitive peripheral nerve endings of ingrowing sinuvertebral branches, thereby producing discogenic pain. Provocation discography was known to simulate the mechanical pressure by increasing the intradiscal pressure, however, its stimulation method is not the same as mechanical stimulation imposed by physical posture and movement. In addition, there are other technical sources of errors during the procedure as well as in interpretation of discography results.

We used an APCD to validate the procedure and interpretation of discography, since the term ‘discography’ does not ensure that the procedure is performed consistently with correct interpretation. Conventional discography may not provide the same results under the same conditions for several reasons: interobserver variability during discography; differences in experience and technique in performing the procedure; and normal biological variations in the test subjects. Furthermore, a false negative response is a common problem for inexperienced discographers because they are likely to inject contrast media at high speed, which leads to a discrepancy between real intradiscal pressure and manometric pressure. The APCD system is helpful in overcoming inherent drawbacks of discography by injecting the contrast media less than 0.01 cc/sec to minimize the difference between postyrisgeal pressure and intradiscal pressure. The APCD system also visualizes intradiscal pressure dynamically and at the end of the procedure, and all data are saved for later accurate interpretation. It should be noted that this low speed of injection is almost impossible to achieve by human hands. Therefore, our method of provocation discography contributes to lowering rates of false negative response and thus has a more valid basis compared to other studies.

Lack of characteristic clinical symptoms and signs is an important element in misleading the clinician’s diagnosis of discogenic pain. Considering this problem, we attempted to classify pain patterns in our patients to guide diagnosis of discogenic pain; however, this classification did not prove to correlate significantly with clinical factors and demographic data. We found that back pain combined with posterior leg pain was most common, followed by back pain only, and back pain with buttock pain. Axial back pain was observed in most patients and has the tendency to be localized in the midline, comparable to diffuse or band-like back pain observed in lumbar zygapophyseal joint arthropathy. The leg pain was diffuse extending to knee joint or ankle, comparable to narrow cord-like sciatica observed in radicular pain of disc herniation. These pain patterns are consistent with other previously published studies. Our results showed that these pain patterns can be induced by various kinds of movements (dynamic factors). Sitting, sitting posture, flexion and walking movement were all closely correlated with induction of discogenic pain, while long standing was statistically associated with L5/S1 disc pathology and walking was least associated with L4/5 disc lesions.

If strict classification guidelines are guaranteed, MRI is an excellent tool to define disc degeneration. In our study, grading of MRI abnormality was correlated with that of CT discogram except grade 5 (p < 0.05). However, it was difficult to match both grading systems in the group higher than grade 3. Painful degenerated disc often appears to maintain disc contour in MRI, yet annular fissures extending into the canal can facilitate leakage of contrast dye. If disc degeneration is severe, it is difficult to raise the intradiscal pressure up to the level to provoke a pain response. Consistent with this idea, our results show pain-provoking pressure is lower in grade 4 than grade 3 of MRI abnormality. This implies that a false negative response may occur due to leakage of contrast media, and subsequent failure to increase intradiscal pressure. Development of a new method to eliminate this limitation of discography is a challenge for the future.

In our series, it is not possible to elucidate the rate of false positive responses. However, all the patients have other clinically relevant information such as clinical symp-tomatology and imaging data directly related or indirectly related with discogenic pain. Several factors including psychological comorbidity, complex pain syndrome, narcotic usage, and secondary gain issues have been suggested to correlate with false positive responses. Carragee et al. suggested that the false positive rate in asymtomatic subjects is about 25%. While in contrast, Bogduk and Modic hypothesized that low pressure disc injection can effectively eliminate false positive discography risks. We observed that the smooth progress of the procedure using the APCD system helped patients to be emotionally more stable during discography.

### Table 3. Comparison of disc degeneration on MRI with CT discogram classification

| Grade | MRI grading | CT discogram |
|-------|-------------|--------------|
| 1     | 0           | 0            |
| 2     | 1           | 1            |
| 3     | 6           | 3            |
| 4     | 13          | 12           |
| 5     | 0           | 7            |
| Total | 20          | 23           |

CT: computed tomography, MRI: magnetic resonance image
and thus very likely contributed to a decrease in the false positive rate.

This study has some drawbacks. The sample size is not large enough to cover the diverse clinical findings of discogenic pain. In addition, a small number of the patients could not be appropriately assigned depending upon their difference of pressure in the patients with positive response. A larger cohort study is thus required for further detailed analysis. However, this is a pioneering study to elucidate and correlate clinical findings in patients with positive disc response defined by automated discography.

CONCLUSION

We have analyzed the patients who showed positive response during automated discography. Centralized midline back pain with diffuse leg pain was commonly observed in these patients with discogenic pain. Sitting was most commonly correlated with pain aggravation, and standing was correlated with pain production particularly in the L5/S1 disc lesion. MRI and CT discogram findings were significantly correlated with each other, however, imaging findings were not helpful to determine which would be the painful disc. One distinct benefit in the use of automated discography is validating the contents of discography and contributing to the accurate interpretation of discography results. In addition, it may lower false positive or negative responses compared with conventional discography.

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