Clinical outcome following DIAM implantation for symptomatic lumbar internal disk disruption: a 3-year retrospective analysis

Kang Lu1
Po-Chou Liliang1
Hao-Kuang Wang1
Jui-Sheng Chen1
Te-Yuan Chen1
Ruyi Huang2
Han-Jung Chen1,3

1Department of Neurosurgery, E-Da Hospital, 2Department of Family Medicine, E-Da Hospital, 3Department of Neurosurgery, E-Da Cancer Hospital, College of Medicine, I-Shou University, Kaohsiung, Taiwan

Background/objective: Internal disk disruption (IDD), an early event of lumbar disk degeneration, is the most common cause of low back pain. Since increased intradiskal pressure (IDP) is associated with symptoms and progression of disk degeneration, unloading a painful disk with an interspinous process device (IPD) is a rational treatment option. The goal of this study was to evaluate the effectiveness of dynamic stabilization with an IPD in the treatment of symptomatic IDD of the lumbar spine.

Patients and methods: Patients with symptomatic IDD were treated with implantation of an IPD, the device for intervertebral assisted motion (DIAM). Diagnosis of IDD was based on typical MRI finding of posterior annular high-intensity zone and positive provocative test on discography. IDP was analyzed intraoperatively. Axial back and leg pain was evaluated with visual analog scale, functional status with Oswestry Disability Index, and final clinical outcomes with Odom criteria. Data from 34 patients followed up for at least 3 years were collected.

Results: DIAM implantation significantly reduced IDP (n=11, \( P<0.0001 \)). All 34 patients reported symptom relief. Thirty-one patients (91%) remained symptom free until the last follow-up. Three patients (9%) experienced recurrence of pain, of which the causes were unrelated to the IDD or surgery. Disk status at the DIAM-implanted segments remained stable. Segmental flexion/extension mobility was preserved in 27 of 30 patients with preoperative mobility. No proximal or distal adjacent segment degeneration was observed. The final clinical outcomes were excellent/good in 31 and fair/poor in three patients.

Conclusion: For patients with symptomatic IDD, dynamic stabilization with DIAM provides pain relief and functional improvement. The implantation maintains disk status and prevents progression of disk degeneration, without compromising segmental flexion/extension mobility or causing adjacent segment degeneration.

Keywords: internal disk disruption, lumbar disk degeneration, DIAM, interspinous process device, intradiskal pressure

Introduction

Internal disk disruption (IDD) has been found to be the most prevalent cause of chronic low back pain (LBP), surpassing facet joint and sacroiliac joint pain.\(^2\) Even in the absence of nerve root involvement, IDD can cause pain in the back and leg.\(^2\) Though somatic LBP is the major presentation, referred thigh pain or leg pain, when present, may be misleading in diagnosis and treatment.\(^4,5\)
IDD is regarded as an early event of disk degeneration. 6 The associated pain is believed to be related to ingrowth of nerves and blood vessels through an annular fissure. 7–10 Such LBP tends to be chronic and persistent over time. 6 Despite the fact that IDD has been identified as a common cause of disabling LBP in young patients, 9 there is little consensus as to what the best treatment is. 8

Because mechanical strains and elevated intradiskal pressure (IDP) have been shown to be closely associated with the progression of lumbar disk degeneration, 10–12 unloading a painful disk in its early stage of degeneration would theoretically alleviate pain and even stop or slow down the degenerative process. A recent development in the surgical treatment of lumbar degenerative diseases is the application of interspinous process devices (IPDs). 13–15 However, little is known about the feasibility and efficacy of IPDs in the treatment of symptomatic IDD.

The aim of this retrospective study was to evaluate the efficacy of implanting an IPD, the device for intervertebral assisted motion (DIAM, Medtronic Inc., Memphis, TN, USA) implant, as the surgical treatment for patients suffering from refractory symptomatic IDD, with at least 3 years of follow-up.

Patients and methods

Patient selection

Between November 2008 and December 2012, 52 patients with symptomatic IDD were selected as candidates for DIAM implantation. Before considering surgery, all patients had received at least 3 months (mean 9 months, range 3–60 months) of failed nonsurgical treatments, which included lifestyle and work load modifications, bracing, and continuous physical therapy. Twenty-six patients had also received one or more sessions of injection or radiofrequency therapy. The treated sites were sacroiliac joints in 17, disks in eight, facet joints in five, and unknown in three patients. All patients felt that their quality of life was significantly affected by the symptoms, and they actively sought for more aggressive measures. None of them had history of malignancy, psychiatric disorders, or previous spinal trauma, infection, or surgery.

The major symptom was axial low back/buttocks pain with or without leg pain. The axial pain/leg pain ratios according to patients’ own perceptions were 100/0 in six, 90/10 in seven, 80/20 in six, 70/30 in eight, 60/40 in three, and 50/50 in four cases. Corresponding neurologic deficits were detected in 13 cases, manifested as motor weakness with or without sensory impairment. None of the patients had intermittent claudication typical of lumbar spinal stenosis or symptoms and physical findings suggestive of hip joint or sacroiliac joint arthropathy.

The diagnosis of IDD was based on imaging studies and provocative discography. Excluding five patients who refused discography and nine who showed negative discographic findings or reported no concordant pain on provocative tests, a total of 34 patients (14 men and 20 women) with a mean age of 38 years (range 24–51) were included in this retrospective study (Table 1). The study protocols were approved by the Institutional Review Board of the E-Da Hospital.

Image criteria

All patients received anteroposterior, lateral flexion/extension radiographs and MRI. Images were analyzed by the operating surgeon and by an independent neuroradiology specialist. MR findings of dark signal with a high-intensity zone in the posterior annulus were regarded as diagnostic of IDD. 16 Meanwhile, the Pfirrmann classification was used for assessing the severity of disk degeneration. 17 According to the Pfirrmann criteria, five cases were classified as grade II and 29 as grade III.

Excluded were patients who showed image findings of disk extrusion or sequestered herniation, advanced disk degeneration, or significant foraminal compromise. MRI findings of treated disks are summarized in Tables 1 and 2.

Table 1 Patient demographics and preoperative MRI findings (n=34)

<table>
<thead>
<tr>
<th>Demographic data</th>
<th>Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>38 (range 24–51)</td>
</tr>
<tr>
<td>Male/female</td>
<td>14/20</td>
</tr>
<tr>
<td>Levels operated</td>
<td></td>
</tr>
<tr>
<td>L3-4</td>
<td>2</td>
</tr>
<tr>
<td>L4-5</td>
<td>25</td>
</tr>
<tr>
<td>L5-6</td>
<td>2</td>
</tr>
<tr>
<td>L3-4, L4-5</td>
<td>4</td>
</tr>
<tr>
<td>L4-5, L5-6</td>
<td>1</td>
</tr>
<tr>
<td>Symptomatology</td>
<td></td>
</tr>
<tr>
<td>Back pain only or predominant</td>
<td>27</td>
</tr>
<tr>
<td>Back pain/leg pain equivalent</td>
<td>7</td>
</tr>
<tr>
<td>Leg pain only or predominant</td>
<td>0</td>
</tr>
<tr>
<td>VAS</td>
<td>7.6 (range 5–10)</td>
</tr>
<tr>
<td>ODI</td>
<td>25.2 (range 18–36)</td>
</tr>
<tr>
<td>MRI findings of treated disks</td>
<td></td>
</tr>
<tr>
<td>Inhomogenous gray signal</td>
<td>31</td>
</tr>
<tr>
<td>Slightly decreased disk height</td>
<td>15</td>
</tr>
<tr>
<td>Annular fissure</td>
<td>26</td>
</tr>
<tr>
<td>Foraminal compromise</td>
<td>19</td>
</tr>
</tbody>
</table>

Notes: *Presence of a sixth lumbar vertebra with intact lamina and spinous process; †Back pain or leg pain predominance was based on every patient’s own perception of the locations of pain; ‡Back pain/leg pain ratios were 100:0 in 6, 90:10 in 7, 80:20 in 6, 70/30 in 8 patients; §Leg pain as the only or major symptom; ¶Compatible with grade III disk degeneration based on Pfirrmann’s MR classification (17); #Hyperintense signal within the posterior annulus fibrosus, shown on T2-weighted MR images; $Associated with annular bulging without protrusion or extrusion of nuclear pulpous.

Abbreviations: VAS, visual analog scale; ODI, Oswestry Disability Index; MR, magnetic resonance; MRI, magnetic resonance imaging.
degeneration with Modic changes, disk space collapse (Pfirrmann grades IV or V), defects of pars interarticularis, spondylolisthesis, lumbar stenosis, or disease at the L5-S1 level. Direct decompression with or without fusion was our option of choice for such patients.

Surgical procedure
All operations were performed by the same surgeon (KL). After midline skin incision, the paraspinal muscles were dissected and retracted to expose the target rostral and caudal spinous processes and laminae. The interspinous and the interlaminar ligaments were removed, while the supraspinous ligament was preserved. Epidural fat, if present, was kept as intact as possible. The DIAM implant was placed between the two spinous processes and laminae under the supraspinous ligament. The implant size was determined by insertion trials (8 mm, 10 mm, 12 mm, and 14 mm). The implant was firmly anchored in place by the two tethers passed around the two adjacent spinous processes, and fixed with crimps.

IDP measurement
IDP was measured intraoperatively in the first 14 cases using an intracranial pressure monitoring device (Codman Microsensor ICP Transducer; Codman & Shurtleff, Inc., Raynham, MA, USA). Written informed consent was obtained from patients to participate in the study. After removing the interlaminar ligament, a limited laminotomy was performed under an operating microscope to expose the nerve root. The nerve root was gently retracted medially. A small puncture of the annulus fibrosus was made with an 18 G needle or a No. 11 scalpel blade. The tip of the microsensor was inserted through the annulus into the center of the disk. A baseline resting IDP reading was obtained first, and then subsequently at three different time points, including distracting between the spinous processes and laminae with a distractor, inserting trial templates, and after DIAM implantation (Figure 1). The sensor was removed before wound closure.

Follow-up assessments
All patients were followed up at 3 weeks, 3 months, 6 months, 12 months, and then annually after surgery. Back and leg pain intensity was assessed with the visual analog scale (VAS) and severity of functional impairment with the Oswestry Disability Index (ODI). Final clinical outcomes were evaluated with the Odom criteria.

Radiographs were obtained at each follow-up to evaluate the implant position, status of the disk, segmental mobility at the DIAM-implanted level, and the proximal and distal adjacent disk levels.

Statistical analysis
Preoperative and postoperative data of VAS and ODI were compared using the paired t-test. Comparisons of IDP data among intraoperative events were made using the Shapiro–Wilk test. All analyses were performed with SPSS 15 (SPSS Inc., Chicago, IL, USA).

Figure 1 Intraoperative IDP recording using a microsensor for intracranial pressure monitoring.
Note: The monitoring was done at a resting state (A), on distraction between the two spinous processes with a distractor (B), a template (C), and after DIAM implantation (D).
Abbreviations: DIAM, device for intervertebral assisted motion; IDP, intradiskal pressure.
Results

IDP data

IDP monitoring was successfully done in eleven patients and abandoned in three patients. The reasons for aborting pressure monitoring included failure to pass the sensor through the annulus and mechanical problems of the sensors or monitors. The mean resting pressure before any interspinous distraction was 267.2±37.7 mmHg. Distraction between spinous processes with the distractor caused an abrupt drop of pressure (mean 81.5±34.0 mmHg). The pressure rebounded on replacing the distractor with a template (mean 0.6±57.7 mmHg). The mean final pressure after securing the DIAM implant was 91.9±38.7 mmHg. Compared to the initial resting IDP, the readings recorded on distraction, template insertion, and DIAM implantation were all significantly lower (P<0.0001; Figure 2).

As the pattern of intraoperative IDP changes was so distinctive and constant, pressure monitoring was not performed in the rest of the cases.

Illustrative case

A 28-year-old woman presented with LBP with radiation to buttocks for 3 years. The pain was dull and persistent. Soreness was felt on her buttocks and posterior thighs. The pain did not radiate to lower legs. There was no numbness on the legs. Her tolerance to upright positions, including standing and sitting, was poor. Her ability to walk was normal. Her perception of pain distribution was 80% on the low back and buttocks. Her tolerance to upright positions, including standing and sitting, was poor. Her ability to walk was normal. Her perception of pain distribution was 80% on the low back and buttocks, and 20% on the legs. Her VAS was 6–8, depending on the time of day. Her ODI was 36. She showed normal posture and gait pattern. Neurological examination disclosed no motor or sensory deficits. She had received radiofrequency therapy twice at two other hospitals, 1 year and 2 months prior to her visit, respectively. Both sessions gave her temporary pain relief, but symptoms soon recurred.

Radiographs showed that she had a sixth lumbar vertebra, and that there was a loss of L5–6 disk height (Figure 3A–C). MRI showed dark signal in the disk with a posterior high-intensity zone (Figure 3D and E). As the L6 level possessed lamina and spinous process that were large enough, a DIAM implant was secured between L5 and L6. She reported relief of the LBP and leg pain immediately after surgery. She remained symptom-free at the 3-year follow-up. Postoperative dynamic radiographs showed preserved flexion/extension mobility and enlarged neuroforamen (Figure 3F–H).

Clinical outcome

All 34 patients reported improvement of pain and functional status at the first follow-up 1 week after surgery. Thirty-one cases (91%) remained free of symptoms and of any requirement for medications until the last follow-up. Mean preoperative VAS and ODI were 7.8±1.5 and 24.6±3.5, while values at the last follow-ups were 1.3±2.2 and 6.3±7.6, respectively (P<0.0001) (Figure 4).

Three patients reported recurrence of back and/or leg pain during follow-ups despite an initial improvement after surgery. MRI was repeated in these patients, which unequivocally revealed resolution of the posterior annular high-intensity zone and widened neuroforamens (Figure 5). The back pain in the first case, a 45-year-old man who underwent L3–4, L4–5 DIAM implantations, started after a fall in a motor vehicle accident. It soon subsided with medical treatment. The second case, a 40-year-old man, suffered new right leg pain caused by L5–S1 disk herniation after relief of LBP following L4–5 DIAM implantation. Microdiscectomy had to be performed 4 months after the initial operation to relieve his symptoms. The third case, a 42-year-old woman, despite initial pain relief after surgery, suffered from recurrence of back and leg pain 3 months later. Imaging and electrophysiological studies did not show any abnormalities. Medications and physical therapy failed to relieve the pain. Meanwhile, she had sleep disorder and menstrual irregularity and visited other specialists for narcotics frequently. The cause of her pain was suspected as psychiatric. A consultation with a psychiatrist was proposed, but refused by the patient. This was the only case who reported poor clinical outcome.

Overall, based on the Odom’s criteria, the final clinical outcomes were excellent in 24 cases, good in seven, fair in two, and poor in one case.
Radiographic outcome

At the DIAM-implanted segments, follow-up radiographs showed no progression of disk height loss, endplate changes, appearance of vacuum clefts, or segmental instability. Before surgery, obvious segmental flexion/extension mobility was present in 30 cases. At the final follow-up, this was preserved in 27 cases, but restricted in two, and lost in one case. The four cases without obvious preoperative flexion/extension mobility remained the same. There was no radiographic evidence of degeneration at any proximal or distal adjacent segments next to the DIAM-implanted segments.

Repeated MRI was obtained in the three aforementioned cases experiencing recurrent pain. Consistently, all the MR images showed increased neuroforaminal widths and resolution of annular fissures at the DIAM-implanted segments (Figure 5).

Complications

No fracture of spinous processes or dural tear occurred during the surgical procedures. No patient developed neurologic deficit after surgery. Superficial wound infection occurred in one patient, which subsided with proper wound care. All patients were ambulatory in a lumbar corset on the day of surgery or the next day. Hospital stay was no more than 3 days in all cases.

Discussion

Different from more advanced lumbar disk degeneration, IDD lacks obvious image findings to justify any aggressive treatment. Even if a patient suffers from severe symptoms, many
would not view them as candidates for surgery. However, IDD may cause chronic disabling symptoms, and may progress to more advanced degeneration. The idea of easing a patient’s symptoms, making their quality of life better, and halting the process of degeneration in its early stage is appealing.

We selected only those who presented with predominantly axial low back/buttock pain with no or relatively mild leg symptoms as candidates for stand-alone DIAM implantation. The diagnosis of IDD was based on MRI and discographic findings. Our results indicate that DIAM implantation is an effective and durable surgical option for IDD. We excluded patients with advanced disk degeneration (Pfirrmann grades IV and V), overt lumbar stenosis, spondylolisthesis, and instability. The rationale was the belief that the DIAM implant provides axial distraction but does not relieve anteroposterior or circumferential compression from hypertrophic ligamentum flavum and facet joints present in advanced disk degeneration and lumbar stenosis. Moreover, such cases often have associated disk space collapse and ankylosis of the facet joints, which would significantly limit the applicability of the DIAM implant and its mobility after implantation.

It is arguable whether an intervertebral disk in its early stage of degeneration should not be treated surgically at all. One of the major findings of this study is that in IDD, the symptoms can be severe enough to affect quality of life badly, even though the disk may appear only mildly degenerated.

While waiting for a longer period for conservative treatment to work is always an option, the patient may have to continue tolerating the symptoms and futile therapy. In addition, there is the risk of progressive degeneration and dysfunction of the disk and facet joints over time to a state where dynamic stabilization with a DIAM implant would no longer be optimal, and fusion might become the only choice.

The patterns of IDP changes were strikingly similar among the measured cases. The data provide biomechanical evidence supporting our treatment. Distraction between two adjacent spinous processes and laminae, whether from a distractor, a trial template, or a DIAM implant, unequivocally reduced IDP (Figure 2). Cadaveric and finite element studies have demonstrated that IDPs significantly unload implanted intervertebral disks. Given the association between elevated IDP and the progression of disk degeneration, our findings imply disk unloading as a mechanism for the clinical benefits of DIAM implantation.

The maintenance of disk height on follow-up radiographs suggests that the DIAM implant was effective in preventing the progression of disk degeneration. Besides, its limitation on flexion/extension mobility appears minimal, provided such...
mobility is present before surgery. Although one biomechanical feature of IPDs is to act as an “extension block”, it was not obvious in our cases. This might have been due to the relatively young age of our patients, as well as the elastic and compressible property of the silicon-based material of the DIAM implant.

The absence of any adjacent segment degeneration, either proximal or distal, suggests that DIAM implantation has little adverse impact on adjacent segments. This is in accordance with previous reports showing that IPDs do not alter the loading at the adjacent segments. Even in the four patients with no preoperative segmental flexion-extension mobility, and the three cases who lost normal mobility after surgery, radiographic changes of adjacent segment degeneration were not observed. It is likely that even if a segment is immobilized by a DIAM implant, it does not become “fixed”. Therefore, the impact on its adjacent segments should be quite different from that caused by rigid fusion.

According to a recent review, placement of IPDs is associated with relatively high complication and failure rates. Common complications included sinusous process fractures and dural tears. Reasons for revision surgery were lack of improvement, worsening of pain, or implant dislocation. In contrast, none of these adverse events occurred in our patients. Several factors might explain the difference. First, our patients were relatively young with very low risk of osteoporosis. Second, an operating microscope was routinely used for removal of the interspinous and interlaminar ligaments, while preserving the epidural adipose tissue if present. This practice should have helped preventing violation of the dura.

Our study has several limitations. First, the number of cases was relatively small. However, the small case number actually reflects our stringency in patient selection. We virtually treated only cases showing early disk degeneration but suffering intractable symptoms. Second, even though MRI and discographic evidence was available for diagnosis, other sources of pain, such as the facet joints and the sacroiliac joints, might have probably existed in some patients. We did not take specific diagnostic measures to exclude them as pain sources. Third, we are unable to provide data concerning facet status before and after surgery. In this regard, even though we have provided evidence of disk unloading (IDP reduction) as underlying the DIAM-mediated symptom relief, we cannot be sure whether facet joint unloading also played a role, though it has been demonstrated in cadavers that IPDs reduce facet loading. Finally, postoperative MRI was done in only the three patients who had recurrence of pain. The National Health Insurance Administration of our country dictates that MRI done without a just cause is liable to be denied reimbursement or even fined. As the causes of recurrent pain were proved to be unrelated to the initial disk degeneration, the MR findings of resolution of annular fissure and enlargement of neuroforamens can be regarded as image evidence supporting the benefits of DIAM implantation for IDD, even though the number of cases was small.

Conclusion

In IDD with intractable symptoms, DIAM implantation provides pain relief, functional improvement, and satisfactory clinical outcomes. Such dynamic stabilization appears effective in resolving annular fissures and preventing progression of disk degeneration. Segmental flexion/extension mobility, when present, tends to be maintained. The implant does not lead to either proximal or distal adjacent segment degeneration. These beneficial effects, together with disk unloading and increased neuroforaminal dimensions, may be the mechanisms for the observed satisfactory clinical outcome.

Disclosure

The authors declare no conflicts of interest in this work.

References


