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Comparison of 2-Level Versus 1-Level Total Disc Replacement: Results From a Prospective FDA-Regulated Trial

Jack E. Zigler, MD,* and Donna D. Ohnmeiss, DrMed†

ABSTRACT

Background
Fusion has been the traditional surgery for painful disc degeneration unresponsive to nonoperative care. Fusion rates may decline in multilevel procedures. Also, fusion may force additional stress onto adjacent discs. This effect may be amplified in multilevel procedures. Single-level total disc replacement (TDR) has been found to be as effective as fusion. There have been few published reports addressing 2-level TDR. The purpose of this study was to compare results of TDR at 2 levels to 1-level procedures.

Methods
This report included the first consecutive 86 patients who had reached 24-month follow-up from among those enrolled in the ProDisc-L investigational device exemption (IDE) study of patients undergoing TDR at 1 level (N = 54) or at 2 levels (N = 32). Clinical outcome measures included visual analog scales (VAS) assessing pain, Oswestry Disability Index, satisfaction measured by VAS, and responses to the question regarding whether the patient would have the same surgery again.

Results
Operative time and length of hospitalization were significantly less in the 1-level cases compared to 2 levels (61.6 min vs 97.8 min; and 1.89 days vs 2.44 days; P < .05). There was a trend for less blood loss in single-level cases (59.0 mL vs 79.2 mL) (.05 < P < .09). VAS and Oswestry scores were significantly improved in both groups postoperatively (by approximately 50%). At no follow-up were there significant differences in VAS, Oswestry, or patient satisfaction scores between the single- and 2-level patients. At all follow-ups, the mean satisfaction in both groups was greater than 7.5 on a scale of 0 to 10.

Conclusions
Patients undergoing 2-level TDR improved significantly postoperatively based on VAS and Oswestry scores, and there were no significant differences in outcome scores when comparing 1- and 2-level TDR.

Clinical Relevance
This study suggests that 2-level TDR can be undertaken in appropriately selected patients and achieve results similar to single-level cases.


INTRODUCTION
In the treatment of painful disc degeneration, lumbar spinal fusion has been the primary treatment after patients have not gained acceptable relief from nonoperative measures. While the results have been acceptable, the percentage of patients who experience good outcomes appears to hit a ceiling of about 75%, regardless of the innovations in fusion technique made during an approximate 20-year period.1 These results are undoubtedly influenced by a variety of factors including the specific surgical technique, surgeon skill and experience, patient personality, and the specific origin of each patient’s symptoms. Some studies have reported that fusion rates decrease, or that there is at least a trend for such, as the number of operated levels increases.2,4 This decline in fusion rates may lead to less-desirable clinical outcomes in patients undergoing fusion for multilevel degenerative disc disease.

One of the potential advantages of TDR over fusion is a reduction in the likelihood of accelerating the degeneration of the transitional adjacent segment. While the impact of adjacent segment degeneration is not yet consistently described clinically,5-10 there are biomechanical studies supporting that fusion increases the pressure within the adjacent disc.11-15. It has also been reported that progressively greater changes in
lumbar segment kinematics correlate with a greater number of instrumented levels. Computer modeling has found that the stress on the disc adjacent to an interbody fusion increases with the number of levels fused. These studies suggest that 2-level fusion may have a greater potential for detrimental consequences at adjacent discs.

While the results of TDR have been found to be favorable in most studies, there has been little investigation of 2-level TDR. One article reviewed 10 2-level and 15 3-level TDR procedures and another noted results of 13 multilevel cases, including 11 2-level procedures. While the results in these series were favorable, they represent a relatively small number of patients. The purpose of this study was to evaluate the outcome of 2-level TDR and to compare the results to single-level procedures using the same implant.

MATERIALS AND METHODS
The study was based on the consecutive series of the first 86 patients undergoing TDR with ProDisc-L (Synthes Spine, West Chester, Pennsylvania) at one center as part of an FDA-regulated clinical trial. There were 32 2-level disc replacements and 54 single-level replacements. All procedures were performed using a retroperitoneal approach to the lumbar spine. The only demographic difference identified between the 2 groups was that the mean age was significantly greater among patients undergoing 2-level replacement (Table 1). All patients were treated for symptomatic disc degeneration unresponsive to a minimum of six months of nonoperative care. All patients had undergone MRI and the majority had discograms. Only patients with a score of at least 40% on the Oswestry Disability Index were included. Patients with greater than Grade I spondylolisthesis, previous lumbar fusion, or clinically relevant facet joint changes were excluded from study enrollment.

Data were collected preoperatively, perioperatively, and postoperatively at 6 weeks, and at 3, 6, 12, 18, and 24 months. Only patients who reached a minimum 24-month follow-up were included in this study. Outcome data included visual analog scale (VAS) assessing pain, Oswestry Disability Index, patient satisfaction (VAS ranging from 0 to 10), and patient responses to whether they would have the same surgery again.

Data Analysis
Means were compared between the 2 groups using independent t-tests and change in preoperative to postoperative mean scores on the VAS and Oswestry were compared using paired t-tests. Chi-square analyses were used to compare proportional data. Statistical analyses were performed using SPSS software (Chicago, Illinois).

RESULTS
Perioperative Data
Operative time and length of hospitalization were significantly less in the 1-level cases compared to the 2-level cases (Table 2). The additional level required approximately 30 minutes of operative time. There was also a trend for less blood loss in single-level cases, although the difference was not significant.

<table>
<thead>
<tr>
<th>Table 1. Descriptives of the 1- and 2-Level TDR Subgroups</th>
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<tr>
<td>1-Level (n = 54)</td>
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</tr>
<tr>
<td>Gender</td>
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<tr>
<td>Male</td>
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<tr>
<td>Female</td>
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<tr>
<td>Age (years):</td>
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<td>Mean</td>
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<td>Levels Operated</td>
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<td>L3-4</td>
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<td>L5-S1</td>
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<td>L3-4 &amp; L4-5</td>
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<tr>
<td>L4-5 &amp; L5-S1</td>
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<tr>
<td>Smoking</td>
</tr>
<tr>
<td>Smoker</td>
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<tr>
<td>Non-smoker</td>
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<tr>
<td>Previous Surgery</td>
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<tr>
<td>Yes</td>
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<td>No</td>
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<tr>
<td>Insurance</td>
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<td>Private Insurance</td>
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The only significant difference was the mean age in the 2-level cases was significantly greater.

<table>
<thead>
<tr>
<th>Table 2. Perioperative Data for 1-Level and 2-Level Subgroups</th>
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<tbody>
<tr>
<td>1-Level</td>
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<td>---------</td>
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<tr>
<td>Blood loss (ml)</td>
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<tr>
<td>Operative time (min)</td>
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<td>Length of hospitalization (days)</td>
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The mean blood loss, operative time and length of hospitalization were all less in single-level cases (p < .05; t-test).

VAS Pain Scores
VAS scores assessing pain improved significantly in both groups from preoperative to postoperative measurements. The improvement was approximately 50% (Figure 1). At no visit was there a significant difference in mean VAS score between the 1-level and 2-level groups.
Oswestry Disability Scores

The Oswestry scores followed a pattern similar to the VAS scores. The postoperative Oswestry scores improved significantly compared with the preoperative scores in both groups (Figure 2), with no significant differences between groups at any of the evaluation periods.

Patient Satisfaction

Patient satisfaction was assessed using 2 different methods. A 10-point VAS was completed at each follow-up visit, with greater scores indicating greater satisfaction. As seen in Figure 3, presenting the mean VAS satisfaction scores, patients in both groups indicated a high level of satisfaction. There were no significant differences between groups at all follow-up periods.

There was no significant difference in the proportional distribution of responses to the question, “Would you have the same treatment again?” (P > .25) when comparing 1- vs 2-level TDR.

DISCUSSION

This study found that clinical results of TDR were not diminished as the number of operated levels increased from 1 to 2 adjacent levels in the lower lumbar spine. Perioperative data indicated that 2-level TDR was associated with statistically significantly greater operative time and length of hospitalization. The addition of the second level required approximately 30 minutes more than the single-level replacements. The mean operative times in our series were quite a bit less than those reported by Bertagnoli et al. using the same implant for 1- and 2-level procedures (median of 81 minutes for 1-level and 135 minutes for 2-level).19,20
difference may be related to operative technique including the approach used to gain access to the anterior lumbar spine.

Our clinical outcomes for single-level cases were similar to those reported in other TDR trials involving single-level TDR. Although Bertagnoli et al. reported median values, rather than mean values, the outcome scores in their series of 10 patients undergoing 2-level TDR appear to be similar to the 2-level cases in our larger series. Collectively these studies indicate that there is stability in the published clinical outcomes of TDR.

In a study of patients who underwent fusion for spondylolisthesis, Wimmer et al. reported that the incidence of translation at the segment adjacent to a fusion was greater among patients with multilevel fusions compared to single-level fusion. Another study reported that the number of levels fused was not related to a greater rate of reoperation at the adjacent segment. However, as the authors discussed, in single-level cases, there were more levels at risk of reoperation than in multilevel fusions. Also those authors noted a trend for reoperation at an adjacent segment that was degenerating at the time of the index surgery. This may have also skewed the data concerning reoperation (if only one of several degenerated discs was operated, this may have artificially increased the reoperation rate for single-level index cases, making it similar to the reoperation rate in multilevel cases).

The results of previous studies investigating adjacent segment deterioration, as well as their designs and populations studied, vary greatly. Biomechanical studies and computer modeling have consistently found altered kinematics and increased disc pressure at segments adjacent to a fusion. In addition, some data suggest that kinematics and disc pressure worsen as the number of fused segments increases. The impact of dynamic stabilization on adjacent segment deterioration has yet to be the topic of sufficient investigation. However, one report, using the Graf system, did find that this motion-sparing technology was associated with a decreased rate of adjacent segment deterioration compared to fusion. Recent biomechanical studies in the cervical spine found that, unlike fusion, TDR did not alter the kinematics of the adjacent motion segment and it did not increase intradiscal pressure. Similar results were found in a study of the lumbar spine. More biomechanical studies as well as clinical data are needed to determine if TDR can have a similar protective impact in the lumbar spine, particularly in multilevel procedures.

The results of this prospective study suggest that 2-level TDR with ProDisc-L is an effective treatment for 2-level symptomatic disc degeneration in appropriately selected patients with pain unresponsive to nonoperative care. While the additional level did increase operative time and length of hospitalization, the clinical outcomes were as favorable as single-level procedures.

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REFERENCES
LUMBAR ARTHROPLASTY


