Instrumentation in lumbar fusion improves back pain but not quality of life 2 years after surgery

A study of 1,310 patients with degenerative disc disease from the Swedish Spine Register SWESPINE

Yohan Robinson, Karl Michaëlsson, and Bengt Sandén

Background and purpose   Instrumented and non-instrumented methods of fusion have been compared in several studies, but the results are often inconsistent and conflicting. We compared the 2-year results of 3 methods of lumbar fusion when used in degenerative disc disease (DDD), using the Swedish Spine Register (SWESPINE).

Methods   All patients registered in SWESPINE for surgical treatment of DDD between January 1, 2000 and October 1, 2007 were eligible for the study. Patients who had completed the 2-year follow-up were included in the analysis. The outcomes of 3 methods of surgical fusion were assessed.

Results   Of 1,310 patients enrolled, 115 had undergone uninstrumented fusion, 620 instrumented posterolateral fusion, and 575 instrumented interbody fusion. Irrespective of the surgical procedure, quality of life (QoL) improved and back pain diminished. Change in QoL and functional disability and return to work was similar in the 3 groups. Patients who had undergone uninstrumented fusion had more back pain than the patients with instrumented interbody fusion at the 2-year follow-up (p = 0.02), although the difference was only 7 visual analog scale (VAS) units (95% CI: 1–13) on a 100-point scale. Moreover, 83% of the patients with uninstrumented fusion used analgesics at the end of follow-up as compared to 68% of the patients who had undergone surgery with one of the 2 instrumented fusion techniques.

Interpretation   In comparison with instrumented interbody fusion, uninstrumented fusion was associated with higher levels of back pain 2 years after surgery. We found no evidence for differences in QoL between uninstrumented fusion and instrumented interbody fusion.

Several studies have focused on the effects of instrumentation in posterolateral fusion. The results up to 2005 were summarized in a Cochrane Review, which concluded that instrumentation appears to lead to a higher fusion rate, but does not appear to improve quality of life (QoL) or to give reduced pain (Gibson and Waddell 2005). Recent randomized controlled trials (RCTs) have supported that conclusion (Fritzell et al. 2002, Ekman et al. 2005, Andersen et al. 2008). 2 RCTs focused on patients with DDD or post-discectomy syndrome only. In a study by Fritzell et al. (2002), no differences between the 3 methods could be seen. In contrast, a study by Neumann et al. (personal communication) showed superior results of transformaminal interbody fusion (TLIF) over instrumented posterolateral fusion (IPF) for most, but not all, of the outcome measures.

The inconsistencies between the results of these studies may be explained by differences in inclusion criteria and in the number of participants. We therefore compared the results of different fusion methods in routine clinical practice. The Swedish Spine Register (SWESPINE) is well designed for this purpose (Zanoli et al. 2006, Strömqvist et al. 2009b).

Patients and methods

SWESPINE, the Swedish Spine Register, was started in 1993. More than 80% of all surgical procedures for degenerative lumbar spine disorders in Sweden are included in the register (Strömqvist et al. 2005). Preoperative questionnaire data and postal follow-up questionnaires are completed by the patients without any assistance from the surgeon. Preoperative data completed by the patient include age, sex, smoking habits (current use/no use), working conditions, sick listing (partial/full duration), use of analgesics (regular/occasional), and walking capacity (given as 4 classes). Back and leg pain are reported on a visual analog scale (VAS) and with the Oswestry disability index (ODI). The Short Form-36 health survey (SF-36) and the European Quality of Life questionnaire (EQ-5D) should also...
In these hospitals had completed the 2-year follow-up. These failed: none of the 117 patients who had undergone surgery that had reported to the register, the follow-up procedures had 34 patients, leaving 2,324 patients. In 4 of the 38 hospitals an invalid personal identification number was registered in

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PINE. The number of levels treated did not differ statistically over the study period (Figure 2), the models were also adjusted for the year of surgery. In many of the hospitals included in the study, 1 surgical method predominated. For dichotomous dependent variables, we used a modified Poisson regression approach with robust error variance (SAS PROC GENMOD) to assess relative risks (RRs) with 95% confidence intervals (CIs) (Zhou 2004). The multivariable models were adjusted as described above.

The study was approved by the Ethics Review Board in Uppsala, Sweden (Dnr 2009/164).

Results

Of the 1,310 patients included in the analysis, 115 underwent UIF, 620 IPF, and 575 IIF. The choice of UIF procedure

be completed. Surgical data, including diagnosis, are recorded by the surgeon without access to the patient’s questionnaires. The current protocol of the register, which has been validated in a test-retest situation, can reliably detect postoperative improvements between large groups of patients such as in a registry (Zanoli et al. 2006, Strömqvist et al. 2009b).

For this register-based study, the population of interest was all patients who had been operated on for painful DDD using any posterior method for lumbar fusion. Data were obtained for all patients registered in SWESpine between January 1, 2000 and October 1, 2007. Other conditions, such as central or lateral spinal stenosis, disc herniation, ischemic spondylolisthesis, postoperative instability, and degenerative scoliosis are mutually exclusive diagnoses in the register and they were therefore excluded from the present analysis.

The fusion methods included were uninstrumented fusion (UIF), instrumented posterolateral fusion (IPF), posterior lumbar interbody fusion (PLIF), and transfemoral lumbar interbody fusion (TLIF). PLIF and TLIF were analyzed as 1 group under the name instrumented interbody fusion (IIF). The reasons for treating the posterior interbody techniques as 1 group were partly the low number of TLIF procedures and partly our suspicion that different modifications of the PLIF procedure had been used in many of the patients described as PLIF in the register. For the UIF procedure, cancellous bone grafting was performed after decortication of posterior bony structures, followed by 3 months of lumbar bracing. Anterior fusion methods were not included in our study because the number of patients was low; in addition, most of the anterior procedures were performed during the early years of SWESPINE. The number of levels treated did not differ statistically significantly between the groups; it ranged from 1 to 5 levels of lumbar fusion.

Of the 2,358 patients who fulfilled the inclusion criteria, an invalid personal identification number was registered in 34 patients, leaving 2,324 patients. In 4 of the 38 hospitals that had reported to the register, the follow-up procedures had failed: none of the 117 patients who had undergone surgery in these hospitals had completed the 2-year follow-up. These patients were therefore excluded. An additional 225 patients had been operated for lumbar fusion more than once during the study period, making it difficult to evaluate the result of one separate procedure. Consequently, these 225 patients were excluded, leaving 1,982 eligible patients (Figure 1). The distribution of the 3 surgical methods in the 225 patients who were excluded was similar to the distribution of the surgical methods in the patients who had completed the 2-year follow-up (25 patients had undergone UIF, 96 patients IPF, and 104 patients IIF). Of the remaining 1,982 patients, 1,310 (66%) had completed the 2-year follow-up.

VAS for back pain or leg pain at baseline, smoking habits, duration of symptoms, and the distribution of the different fusion methods were similar in the 672 patients who did not complete the 2-year follow-up and in the patients with complete follow-up. Factors that negatively affected the response rate were low age, male sex, previous spine surgery, low EQ-5D at baseline, and high ODI at baseline (data not shown).
depended on the hospital the patient was operated in. In 1 hospital, 72% of the patients underwent UIF, while this procedure was not performed at all in several other hospitals.

The patients in the IIF group were slightly younger, whereas the proportion of smokers in this group was smaller. The number of patients who had undergone previous spine surgery was lower in the IIF group (Table 1).

**Generic and condition-specific outcome measures**

All groups improved from baseline with regard to EQ-5D and ODI (all p < 0.01). The results 2 years after surgery were similar for the 3 fusion methods studied, as measured with the EQ-5D and the ODI (Table 2).

**Back and leg pain**

Pain was recorded at the 2-year follow-up using VAS. All groups improved from baseline to follow-up with regard to both back pain and leg pain (all p < 0.01). The patients who had undergone UIF generally had more back pain than the patients with IIF at the 2-year follow-up (p = 0.02), although the difference was only 7 VAS units (CI: 1–13) on the 100-point scale. Leg pain was similar in the 3 groups (Table 2).

**Use of analgesics and return to work**

At the 2-year follow-up, use of analgesics was more frequent in the UIF group (83%) than in the other 2 groups (68% on average: IPF 70% and IIF 65%), corresponding to a multivariable adjusted RR of 1.2 (CI: 1.1–1.3) for those treated with UIF.

The frequency of returning to work was analyzed for those patients who were less than 65 years of age at the time of follow-up, and who had been working before surgery. The RR for returning to work in the UIF group was 0.97 (CI: 0.8–1.2), it was 1.04 (CI: 0.9–1.3) in the IPF group, and it was 0.97 (CI: 0.8–1.2) in the IIF group, indicating no significant differences between the groups.

**Discussion**

The patients in this study showed significant improvements in back pain, function, and QoL 2 years after surgery when measured with VAS, ODI, and EQ-5D regardless of surgical method (Table 2). Due to the large sample size, however, statistical significance could be achieved with small improvements that are not clinically relevant. In the annual report of SWESPINE, the minimal clinically important difference (MCID) for VAS after fusion surgery was estimated to be 14 points, and the MCID for EQ-5D was estimated to be 0.2 (Strömqvist et al. 2009a, Gatchel et al. 2010). For surgical interventions, an

### Table 2. Outcome 2 years postoperatively related to surgical method. Values are adjusted means (95% CI) a

<table>
<thead>
<tr>
<th></th>
<th>UIF b</th>
<th>IPF b</th>
<th>IIF b</th>
</tr>
</thead>
<tbody>
<tr>
<td>EQ-5D</td>
<td>0.61 (0.54–0.69)</td>
<td>0.62 (0.57–0.67)</td>
<td>0.64 (0.59–0.69)</td>
</tr>
<tr>
<td>Oswestry disability index</td>
<td>28 (23–34)</td>
<td>28 (24–33)</td>
<td>27 (23–32)</td>
</tr>
<tr>
<td>Back pain, VAS</td>
<td>40 (33–47)</td>
<td>34 (30–39)</td>
<td>33 (28–38)</td>
</tr>
<tr>
<td>Leg pain, VAS</td>
<td>32 (24–39)</td>
<td>29 (24–34)</td>
<td>29 (23–35)</td>
</tr>
</tbody>
</table>

a The estimated mean values have been adjusted for differences in age, sex, smoking, previous spine surgery, duration of symptoms, year of surgery, hospital, and differences at baseline for the variable studied.

b For abbreviations, see Table 1
MCID of 15 points for ODI has been suggested (Roland and Fairbank 2000). In SWESPINE, however, the MCID for ODI after fusion surgery was estimated to be 8 points (Strömqvist et al. 2009a). Thus, the patients in our study experienced clinically important improvements in back pain, QoL, and functional disability after surgery.

However, the patients who had undergone UIF reported more back pain 2 years after surgery than the patients treated with an interbody fusion method, but the difference could not be regarded as clinically important. With UIF, there was an indication of an increased probability of use of analgesics compared to IPF and interbody fusion. There were higher levels of back pain after UIF despite a higher consumption of analgesics. Apart from these findings, there were no significant differences evident between the 3 surgical techniques.

The clinically and functionally superior improvement from instrumented fusion as compared to uninstrumented fusion was possibly due to a hypothetically greater rate of fusion. Furthermore, the ability to address the patient’s sagittal balance with instrumentation could improve the long-term results compared to uninstrumented cases. The finding that the postoperative results were similar in all 3 groups could have been biased due to the fact that only a few surgeons were performing many UIFs, thus being highly specialized in this technique, and possibly performing meticulous bone grafting and postoperative bracing. Of course, most surgeons have a preference for one surgical method over the other, whether it is based on evidence or simply on belief. This reasoning is rather hypothetical, but it must be taken into account when evaluating the results of the present study.

The number of patients who were lost to follow-up was a limitation of our study. Of the 1,982 patients included in the study, 1,308 (66%) completed the 2-year follow-up. The patients who were lost to follow-up had a higher frequency of previous spine surgery, had a higher chance of being male, and were somewhat younger than the patients who completed the follow-up. At baseline, the patients who were lost to follow-up generally had an inferior QoL and somewhat higher functional disability. The attendance rate was, however, equal for the 3 surgical methods and the characteristics of the patients who were lost to follow-up were equivalent in the 3 groups. Furthermore, the distribution of the 3 fusion methods was similar in the 225 patients who were excluded because of repeated fusion surgery during the study period. In a recent study from Norway based on a local register for degenerative lumbar surgery, the results from the non-responders were compared with the results from the responders. In that study, 22% of the patients were lost to a 2-year follow-up. These patients were subsequently traced and interviewed by telephone. There were no statistically significant differences in outcome between the responders and the non-responders (Solberg et al. 2011). Unfortunately, the nationwide Swedish spine register has a slightly worse degree of loss to follow-up (34%), which might be due to worse register logistics, insufficient discipline of the registering surgeons, or worse patient-reporting morale. However, both our statistical dropout analysis and the results from Norway suggest that the loss to follow-up probably did not affect the external validity of this study.

A further concern with our study is that only 9% of the patients registered received the uninstrumented procedure, which could indicate that this treatment strategy was only used for highly selected patients. However, the baseline data were similar for all 3 treatment groups with regard to most of the variables registered. Furthermore, the choice of UIF procedure depended on the hospital the patient was operated in. To minimize effects of confounding and possible selection bias, outcome values were adjusted for age, sex, smoking, previous spine surgery, duration of symptoms, hospital, and differences at baseline for the variables under study.

Apart from these factors, our analysis included an adjustment for the year of surgery. The reason for this adjustment was partly that we wanted to minimize the influence of any learning factor and partly our assumption that changes in Sweden’s social security system could influence the results. The frequency of sick listing has decreased in Sweden since 2003, probably because the authorities have made a massive effort to promote early return to work. The number of laborers on long-term sick leave in 2008 was less than half of the number in 2002 (Jonsson 2009). Because these changes appeared during the study period and because the different fusion methods were not evenly distributed during this period, it was obvious that the analysis of returning to work required adjustment for year of surgery. This adjustment not only influenced return to work but also all of the other variables studied.

The results of fusion surgery in Sweden have improved during the past decade, as measured by EQ-5D or Global Assessment (Strömqvist et al. 2009b). This improvement can probably be partly explained by improved surgical techniques and improved selection of patients for the procedure. Moreover, it is known that changes in the compensation system influence registered disability and well-being. This phenomenon was first described in 1879 (Parker 1977), and there have been several reports of the influence of compensation systems—not only on return to work, but also on functional disability and QoL in people with low back pain (Haddad 1987, Greenough and Fraser 1989, Sanderson et al. 1995, Carreon et al. 2010). The improvement in the results of fusion surgery in Sweden during the past decade could be explained by an improved experienced QoL and function due to a greater level of return to work. These factors should be considered when results of different studies are compared.

We did not analyze the complication rate in this study. We felt that the quality of complication data in SWESPINE is not optimal. In general, more complicated methods lead to higher frequencies of complications, indicating that instrumentation with pedicle screws and interbody fusion techniques generate more complications than posterolateral fusion techniques (Fritzell et al. 2003).
Several RCTs on different methods of instrumentation have been performed. In most of these studies, different diagnoses were included (e.g. isthmic spondylolisthesis, degenerative olisthesis, and DDD). In a Danish study a combination of IPF and anterior fusion did not lead to better functional outcome 2 years after surgery compared to IPF alone (Christensen et al. 2002). However, at follow-up 5–9 years after surgery in the same patients, IPF combined with anterior fusion showed superior results (Videbaek et al. 2006). In two studies comparing IPF with posterior interbody lumbar fusion (PLIF), no differences in clinical outcome could be found (Kim et al. 2006, Cheng et al. 2009).

The divergent results of randomized studies on different fusion methods can (at least to some degree) probably be explained by patient selection. Inclusion criteria such as “pain emanating from L4-L5 and/or L5-S1” (Fritzell et al. 2001), “disabling back and/or leg pain….refractory to at least 6 weeks of conservative treatment” (Kim et al. 2006), and exclusion criteria such as “psychosocial instability” (Christensen et al. 2002) or "secondary gains from surgical fusion” (Kim et al. 2006) are not well defined, and the patient groups in different randomized studies have probably been quite heterogeneous. The inclusion criterion in our study was the diagnosis DDD, as put by the surgeon. This diagnosis is chosen if disk degeneration is evident on MRI-imaging but if no other cause of pain (i.e. central spinal stenosis, foraminal stenosis, or disk herniation) can be identified and manual provocation of the degenerated segment induces pain.

As previously reported (Strömqvist et al. 2009b), factors other than the type of surgery (e.g. sex, smoking, duration of symptoms, and previous spine surgery) influence the outcome. We believe that further improvements in the results after fusion surgery are more likely to appear by appropriate selection of patients rather than by the use of increasingly demanding surgical procedures.

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