This prospective study aimed to quantify the risks and complications associated with AxiaLIF in a series of 29 patients. AxiaLIF is a fusion technique using a percutaneous retrorectal, presacral corridor approach to access the L5-S1 and L4-L5 intervertebral spaces transaxially, through the body of S1 and L5 vertebrae. The fusion rate in the present series was 92% and the reported results ranged from 68% to 100%. The only serious complication in the authors’ series was one presacral haematoma (1/29, or 3.5%). Symptomatic subsidence occurred in the stand alone group, resulting in foraminal stenosis and radiculopathy in two patients (7%) and back pain in one (3.5%). Painful radiolucent halo around the rod was noted in a spondylolytic case (1/29, or 3.5%); it resolved after transpedicular instrumentation.

AxiaLIF is a novel truly minimally invasive technique not requiring blood transfusion and can be safely performed as a day surgery. Retroperitoneal haematoma, ureteral and vascular injuries can be avoided by respecting the regional anatomical landmarks as guided by accurate fluoroscopy. Only expanding haematomas may have to be drained. Bowel perforation can be prevented by gently sweeping away the rectum from the sacrum before inserting the guide probe.

**Keywords**: minimal invasive; AxiaLIF; spinal instrumentation; complications.

**INTRODUCTION**

The conventional open surgical approaches for posterior lumbar interbody fusion (PLIF), anterior lumbar interbody fusion (ALIF) and intertransverse fusion (IF) are associated with several complications. More specifically IF is linked with extensive damage to the musculotendinous tissues predisposing to chronic back pain, long hospital stay, postoperative pain and morbidity and extensive blood loss and late return to normal activities (7,9,11,37).
Complications reported with PLIF are nerve root injuries, dural leak, dural and epidural scar tissues that predispose to remote neural complications. ALIF complications (31) include vascular and ureter damage, retrograde ejaculation, impotence, sympathetic dysfunction, deep vein thrombosis, pancreatitis, bowel injury. Furthermore, an access surgeon is required (31) by a good number of surgeons either because they are not well versed with the intricate anatomy of the region or for medicolegal reasons. For these reasons these procedures have been supplanted by three minimally invasive surgical procedures with maximized effects: transforaminal lumbar interbody fusion (TLIF), axial lumbar interbody fusion (AxiaLIF) and extreme lateral lumbar interbody fusion (XLIF). The advantages of minimally invasive procedures are decreased blood loss thus avoiding blood transfusion, decreased postoperative pain and morbidity, decreased hospital stay and medical cost and a faster return to normal activities.

Each of these procedures has its advantages and disadvantages. In this manuscript we will address the indications and pitfalls of the surgical technique associated with AxiaLIF.

AxiaLIF, a recently introduced minimally invasive spinal instrumentation, is a special threaded-rod device (TranS1, Wilmington NC, USA), designed to stabilize the lower lumbar spine, achieve fusion with minimal morbidity, shorten the hospital stay and avoid neural, vascular and musculotendinous injuries. This transacral fusion technique using a percutaneous retrorectal, presacral corridor approach to access the L5-S1 and L4-L5 intervertebral spaces transaxially, through the body of S1 and L5 vertebrae (8) is an evolution of four previously reported techniques with similar objectives. One of the techniques described in a textbook by FW Ruthke consists of paraxial introduction of a rib strut through the sacrum, L5-S1 interspace, L5 vertebra, L4-L5 interspace to L4 vertebra (35). Another technique described by René Louis in 1996 for the treatment of L5-S1 instability particularly for spondylolisthesis involves an open paraxial approach at the lumbosacral spine using a fibular strut (16,23,28). A third technique entails a midline cage placement for L5-S1 fusion in spondylolisthesis through a sacral laminectomy (4). Finally, a fourth technique was designed by McMillan et al (24) for fusion of L5-S1, through a percutaneous posterolateral approach.

The goals of this study are to evaluate the claims of safety and effectiveness of this procedure, to analyze the surgical complications encountered in our series and finally, to formulate and highlight useful information in order to render this novel approach safe and effective.

MATERIAL AND METHODS

We conducted a prospective study of 29 patients – 21 women and 8 men – treated in our institutions over the past four years. The average age was 62.5 years (38-82). Indications for surgery are listed in table I. Surgery was performed at one level (L5-S1) in 25 patients and two levels (L5-S1 and L4-L5) in 4 patients. In the one level group, four patients underwent only AxiaLIF, (stand alone), in four the procedure was enhanced with facet screws and in 17 with pedicle screws. For the two-level fusion (L5-S1 and L4-L5) AxiaLIF was supplemented in all four patients with pedicle screws. Six of the patients were obese with body mass index (BMI) greater than 40. Of the one level L5-S1 fusion group that was enhanced with tranpedicle instrumentation, 3 had previous surgical procedures. Two were treated with L4-L5 XLIF and one with transpedicular L3-L4 & L4-L5 instrumentation. The follow-up period ranged from 24 months to 4 years. At six months, twelve months and 24 months, the patients were evaluated for clinical outcomes using the Oswestry Disability Index (ODI) (17) for function and the visual analogue scale (VAS) for pain. Assessment for fusion and technical complications, such as loosening of instrumentation, subsidence, vertebral fracture, compromise of spinal alignment, haematomas and other surgical failures were investigated using post-operative thin-slice CT scans (at 6 months and 12 months) and dynamic radiographs in flexion-extension at 6 months. Comparison between the groups was made with one way ANOVA. The significance level was set at p < 0.05.

RESULTS

The average surgical time for one level AxiaLIF was 50 min, for a stand-alone procedure, 90 min when surgery was enhanced with a transfacet screw fixation and 110 min when supplemented with transpedicular screw instrumentation. The corresponding surgical time for a two-level stand-alone
AxiaLIF was 90 min and 3 hours when supplemented with transpedicular screw instrumentation. Blood loss for AxiaLIF alone was less than 50 ml and when augmented with the transpedicular screw instrumentation between 100 to 200 ml. Blood loss for transfacet screw fixation was minimal. The postoperative hospital stay was up to 24 hours for one level AxiaLIF, and between 48 hours to 72 hours for two-level instrumentation. The extra morbidity time can be attributed to the transpedicular screw instrumentation, since for the AxiaLIF with or without transfacet screw fixation most of the patients were ready to leave the hospital after recovery from anaesthesia.

There was an average 50 mm (p < 0.001) decrease in VAS scores (from 75 mm to 25 mm, p < 0.001) and 33 points (from 55% to 22%, p < 0.001) in Oswestry scores, between preoperative and postoperative clinical assessment at 3 months that was almost sustained at 6, 12 and 24 months follow-up in 25 patients; one patient was lost to follow-up at 12 months and three more at 18 months.

One patient developed retroperitoneal presacral haematoma 4 days after surgery. He presented with symptoms consisting of a gradual onset of abdominal cramps and discomfort associated with night sweats and low-grade fever in the evenings that lasted for two weeks. Bowel movement was not affected. CT-scan revealed presacral haematoma; ESR, CRP and complete blood count were moderately elevated. The patient was treated empirically with oral antibiotics and he recovered completely after 2 weeks.

Another patient with spondylolysis (treated with AxiaLIF and transfacet screws) developed a painful radiolucent halo around the threaded rod which manifested with low back pain four months after surgery. Complete healing of the osteolysis around the threaded rod was achieved after posterior transpedicular instrumentation and fusion (Fig. 1, 2 & 3).

Three patients from the stand-alone group developed subsidence manifesting as a back pain in one patient and L5 radiculopathy in two. The former was treated with posterior percutaneous transpedicle instrumentation and the latter two patients responded well with decompressive foraminotomy and percutaneous transpedicle instrumentation.

One obese patient exhibited superficial skin burn from inadvertent use of the cautery and this resulted in wound dehiscence, which healed satisfactorily with local wound dressings. Otherwise, all obese patients responded very well to this procedure.

**Fig. 1.** — CT images demonstrate spondylolysis that was missed preoperatively. The patient developed low back pain 3 months after surgery.

<table>
<thead>
<tr>
<th>Pathology</th>
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<tr>
<td>LBP associated with disk collapse</td>
<td>6</td>
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<tr>
<td>LBP &amp; SCIATICA associated with disk collapse</td>
<td>16</td>
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<td>Spondyloytic spondylolisthesis</td>
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<td>LBP after interspinai spacer for spinal stenosis</td>
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LBP = Low Back Pain.

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LBP = Low Back Pain.
For obese patients, AxiaLIF is a convenient procedure because it avoids extensive dissection and necrosis of fat tissue. After treating 91 obese patients (BMI > 30), including 27% patients with morbid obesity (BMI > 38), Rodgers et al concluded that AxiaLIF is safe and useful for obese patients (34).

Long scoliosis surgery with extension to the sacrum is associated with high complication rate of pseudoarthrosis (13,14). Supplementation with AxiaLIF, as an adjunct procedure, to stabilize the L5-S1 level, is a convenient expedience because of its minimal blood loss and short operating time (3) (Fig. 4 & 5) as opposed to other interbody fusion techniques (10).

Biomechanical testing adds support to the notion that this device should be avoided as a stand-alone, particularly in the presence of spondylolysis, and should be augmented with posterior instrumentation as in the ALIF, PLIF & TLIF situation (21). Bone resorption surrounding the axial rod that occurred in one of our patient (Fig. 2) was probably due to excessive motion of an unstable segment (L5-S1 spondylolysis), (Fig. 1) that healed after posterior transpedicle stabilization (Fig. 3). This can further be supported by biomechanical data suggesting that enhancement of transaxial rod fixation with posterior instrumentation results in higher construct stability and successful fusion (2,15).

DISCUSSION

AxiaLIF is a truly tissue sparing approach in terms of maintaining the lumbosacral supporting musculotendinous structures. The damage to the lumbosacral paraspinal musculature is attributed to the transpedicular instrumentation, which, however, can be of minor importance if a minimal muscle splitting technique, or percutaneous insertion is used. This procedure has advantages over other procedures, since no annular damage to the intervertebral disc is inflicted, the anterior and posterior longitudinal ligaments are not violated, and the facet joints are not compromised (8). The preservation of these structures, which are integral for spinal stability, give significant biomechanical advantage over the other interbody devices necessitating the sacrifice of crucial musculoligamento-osseous structures (20,21).

AxiaLIF is also quite effective for correcting and stabilizing grade I and II spondylolisthesis as demonstrated by Rodgers et al in a series of 79 patients (33).

Two patients demonstrated inadequate fusion at L5-S1 interbody space, as observed at one and two years follow-up. However, they were asymptomatic in their clinical assessment. Therefore, the fusion rate was 92% (23 out of 25 patients). In this series no other serious or minor complications were encountered.
in order to allow a differential distraction between anterior (greater distraction) and posterior (lesser distraction) intervertebral disc space.

Subsidence, as occurred in our patient, may provoke compressive radiculopathy. Posterior stabilization with pedicle screws may have some effect but apparently cannot prevent altogether some asymptomatic subsidence, which apparently is not an infrequent occurrence (32) (Fig. 7). Potentially, subsidence may be prevented by purchasing securely the base of the threaded rod in the cortical bone of the S2 sacral segment. Furthermore, overdistraction should be avoided particularly in osteoporotic patients.

Presacral or retroperitoneal haematoma is not strange occurrence in anterior spine procedures or

The lumbar lordotic curve, at L5-S1 level, can be improved by placing the AxiaLIF threaded rod, which has a variable thread count, more anteriorly
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various general surgical, urological and gynecological procedures. Most of the time, small amounts of blood will be re-absorbed in a matter of a few days to weeks (5). Less than 1% of patient is expected to develop symptomatic haematoma consisting of decreased haemoglobin and haematocrit, bloody discharge, sacral pain even compression of the rectum and urinary bladder (1). On CT-scan the haematoma is usually demonstrated as a hyper-dense mass in the presacral area (22). The origin of presacral haematoma in AxiaLIF surgery has not been documented with certainty. Small vessels can be disrupted when dissecting through the pre-sacral fat tissue (1). Premature draining of the haematoma may risk further excessive bleeding caused by rupture of the tamponated haematoma. Only an expanding pelvic haematoma with unstable vital signs should be addressed aggressively. In a series of 285 patients, reported by Smith et al (36), only two out of 5 patients who developed retroperitoneal haematoma required re-exploration and drainage. It is important for the surgeon to be familiar with the anatomical landmarks in the region. The midpresacral coronal safe zone dissection is limited laterally by the iliac vessels and ureter, which should not be violated during the process of instrumentation.

For S1-S2 levels this ranges between 6.5 cm to 6.9 cm for males and 5.4 cm to 6.8 cm in females. Distal to S1-S2 the hypogastric nerve and parasympathetic nerves are located several centimeters lateral to the midline, and therefore are not at risk (26,38). The mean sagittal distance from the anterior sacral margin to the rectum at S1, S2, and S3 levels was reported for men to be 16.2 mm, 14.9 mm and 13 mm respectively and the corresponding figures for women are 11.9 mm, 12.2 mm and 10.6 mm (29).

The most dreadful complication of AxiaLIF is rectal perforation. Documented statistics have not been published except in one case report (6) and one case series of 50 patients (5). The signs and symptoms of rectal injury are hypogastric pain, nausea and melaena (6). Compatible findings with rectal perforation on CT-scan are presacral soft tissue fluid density with fat strading, extraluminal rectal contrast and gas within the areas of soft tissue enhancement (6).

Certain medical situations such as previous rectal or perirectal surgery, pelvic radiotherapy and anterior or transperitoneal or retroperitoneal spinal approach are considered absolute contraindications for AxiaLIF. Relative contraindications are Crohn’s practi-
tis, ulcerative colitis, diverticulitis and full-thickness rectal prolapse. A history of intra-abdominal infections such as appendicitis, or a pelvic inflammatory disease predispose to rectal injury (5,6,25,30). Open or minimally invasive anterior and retroperitoneal approaches to the lumbar spine have also been reported to provoke intestinal adhesions (12,18,19,27). Smith et al have not encountered rectal injuries, in a large series of 285 consecutive cases (36). Preoperatively, all patients should undergo full mechanical bowel preparation the day before the procedure. Parenteral prophylaxis against both aerobic and anaerobic microbial organism should be given prior to skin incision. Rectal injury was reported to respond satisfactorily with temporary diversionary ileostomy or colostomy and a course of IV antibiotics (5,29). Primary colonoscopic repair can be successful if the diagnosis of bowel injury is made preoperatively. The main objective of this clinical study was to scrutinize the complications of this procedure in our series in order to establish technical guidelines for avoiding potential pitfalls and render AxiaLIF a safer procedure. It remains to determine its level of effectiveness in relation to other techniques in controlled randomized studies. The postoperative morbidity is minimal and the patient can be ambulatory soon after the recovery period from anaesthesia and hospitalization is less than 24 hours for one level fusion. Its minimal tissue damage also renders this procedure safe for the elderly patients (36). Since in our series we encountered no serious complications, and the fusion rate is relatively high, this suggests encouraging perspectives.

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