Computerized tomography evaluation of a resorbable implant after transforaminal lumbar interbody fusion

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Object. Synthetic bioabsorbable implants have recently been introduced in spinal surgery; consequently, the indications, applications, and results are still evolving. The authors used absorbable interbody spacers (Medtronic Sofamor Danek, Memphis, TN) packed with recombinant bone morphogenetic protein (Infuse; Medtronic Sofamor Danek) for single- and multiple-level transforaminal lumbar interbody fusion (TLIF) procedures over a period of 18 months. This is a consecutive case series in which postoperative computerized tomography (CT) scanning was used to assess fusion status.

Methods. There were 22 patients (17 men, five women; 39 fusion levels) whose mean age was 41.6 years (range 23–70 years) and in whom the mean follow-up duration was 12.4 months (range 6–18 months). Bridging bone was noted as early as the 3-month postoperative CT scan when obtained; solid arthrodesis was routinely noted between 6 and 12 months in 38 (97.4%) of 39 fusion levels. In patients who underwent repeated CT scanning, the fusion mass appeared to increase with time, whereas the disc space height remained stable. Although the results are early (mean 12-month follow-up duration), there was only one noted asymptomatic delayed union/nonunion at L5–S1 in a two-level TLIF with associated screw breakage. There were no infections or complications related to the cages.

Conclusions. The bioabsorbable cages appear to be a viable alternative to metal interbody spacers, and may be ideally suited to spinal interbody applications because of their progressive load-bearing properties.

Key Words • bioabsorbable implant • polylactide polymer • bone morphogenetic protein • lumbar fusion • computerized tomography

Structural interbody support has become a well-accepted component in the armamentarium for spinal fusion. First advocated by Steffee and widely popularized by Harms, this can be accomplished via the ALIF, PLIF, or more recently via the TLIF procedure. Structural interbody support with titanium surgical mesh in combination with posterior instrumentation has been shown to yield biomechanically superior results. Furthermore, titanium surgical mesh has performed well to date in many centers, but there are still problems with imaging artifacts, potential for late implant infection, and persistent stress shielding of the fusion mass bone.

Bioabsorbable implants appear to possess the ideal characteristics of an interbody fusion device. They provide immediate postoperative stability but permit controlled load sharing over time through resorption. Furthermore, there appears to be no significant inflammatory response or foreign body reaction, which can lead to sterile abscess formation and/or an increased infection rate. In addition, fusion status is easily assessed because of the radiolucent nature of the implant.

The advantages of BMP are also well known and include decreased operating time and blood loss compared with iliac crest harvesting, decreased morbidity, and perhaps increased fusion rates. The efficacy of rhBMP-2 in interbody fusions has been reported previously in animal models and preliminarily in controlled studies. Nevertheless, few clinical studies regarding the use of bioabsorbable implants packed with rhBMP-2 have been reported. The use of BMP may also have the added advantage of decreased infection rates, as seen in the trial of its use in open tibia fractures.

CLINICAL MATERIAL AND METHODS

Patient Population

Over the past 18 months, 35 patients have undergone a single- or multiple-level TLIF (mean 1.7 levels, range 1–5 levels) at our institution with implantation of absorbable cages and segmental pedicle screw fixation with titanium multiaxial pedicle screws and 5.5-mm-diameter rods. Of...
these patients, 22 had at least a 6-month follow-up duration and are the focus of this paper. There were 17 men and five women whose mean age was 41.6 years (range 23–70 years) and in whom the mean follow-up duration was 12.4 months (range 6–18 months). The most common diagnoses were degenerative disc disease and isthmic spondylolisthesis (Table 1). Four cases involved revision surgeries, whereas two were for failed-back syndrome. Six of the 22 patients were known smokers. In addition to the standard clinical follow-up review, immediate postoperative CT scans and standing plain radiographs were obtained in all patients. Plain radiographs were also obtained at 6 weeks, at 3, 6, and 12 months and at the latest follow-up review. The CT scans were obtained at 6, 12, and/or 18 months as appropriate to assess fusion status.

Radiographs were evaluated for instrumentation complications, disc height, and evidence of bridging bone in the interbody space. The CT scans in which sagittal and coronal 3D reconstruction was used were evaluated for bridging bone in the interbody space as well as evidence of dislodgment, subsidence, and/or a sterile abscess or re-active zone around the implant.

**Resorbable Interbody Cages**

All procedures in this series were performed using the bioabsorbable noncrystalline copolymer implant HYDROSORB (Medtronic Sofamor Danek, Memphis, TN), which has a 70:30 ratio of poly(L-lactide) to poly(D,L-lactide). The device made from this material has an elastic modulus of 3.15 GPa, an ultimate compressive strength of 100 MPa, an ultimate tensile strength of 58 MPa, and a ductility of 5% elongation to failure (Fig. 1). It can withstand a compressive load to failure of 8230 N after sterilization, which exceeds the load necessary to crush most vertebral bodies. It degrades by surface dissolution and does not form crystalline degradation products. This resorbable form has significantly reduced the incidence of sterile abscess formation and the concerns about infection that were first noted during the early attempts to use bioabsorbable implants. The HYDROSORB cages come in various heights (5–20 mm) and in various diameter configurations (11, 13, 19, and 26 mm). The wall thickness is a standard 2 mm for the smaller-diameter components, and 4 mm for the 19- and 26-mm components. Insertion is completed using a specialized straight or angled inserter. In this study, the typical implant was 11 or 13 mm in diameter and approximately 9 mm in height (range 5–11 mm height), depending on the vertebral level and diagnosis. Routinely, the implants were placed in a unilateral fashion via a transfemoral approach, but they can obviously be placed via a bilateral, PLIF, or ALIF approach. Degradation occurs over 18 to 36 months, with a minimum of 50% of the initial structural strength maintained at 1 year postoperatively. The implant is also well visualized on CT scans prior to degradation. Nonetheless, some degradation is still influenced by host conditions, as well as by the implant size and geometry.

**Bone Morphogenetic Protein—2**

All procedures were also completed using rhBMP-2, which was packed into the cages before insertion. A standard small Infuse bone graft collagen sponge (Medtronic Sofamor Danek) is adequate for a single-level TLIF® completed with two interbody spacers. In most cases, part of the sponge was placed anteriorly to the two cages as well. Nevertheless, a large collagen sponge is more cost effective when performing a fusion at two or more levels, because a single large kit can adequately cover a two-level interbody fusion. A cost analysis of BMP in spinal fusion has previously been published.

**RESULTS**

Eight single-level and 14 multiple-level (two–five levels) fusions were performed (39 fusion levels, Table 2). The mean operating time was 4.2 hours (range 2.25–8.9 hours) or 2.3 hours per fusion level (range 2.25–3.8 hours/level). The mean blood loss was 380 ml (range 150–800 ml) or 223.5 ml per level. The mean hospital stay was 4.7 days (range 3–9 days). Although there was no control group in this study, these parameters appear to be less than or consistent with those in previously reported series. There were also 15 active-duty soldiers in the study, all of whom remained on active duty at the last follow up.

There were no infections, allergic reactions, deep venous thromboses, or implant complications. There were three postoperative neurological changes, however, two of which were considered to be transient neurapraxia or mild motor weakness that was considered secondary to nerve root retraction at the operative level. Both of these deficits resolved within 6 weeks postoperatively. This is a fairly common finding after TLIF procedures. The changes were not considered to be secondary to the position or insertion of the implant. The third neurological postoperative change was a persistent right-sided L-5 neuropraxia, which also occurred at the operative level in an L4–S1 two-level TLIF for Grade II L5–S1 spondylolisthesis treated with partial reduction. At 6 months postoperatively, this patient underwent a selective L5–S1 foraminal decompression, which relieved the radiculopathy. There was also one intraoperative dural tear, which was primarily repaired without sequelae. There were no complications related to the resorbable cages or rhBMP-2.

**Radiographic Findings**

Standing lateral plain radiographs were evaluated for
instrumentation complications, posterior disc height, and evidence of bridging bone in the interbody space. All measurements were performed on a digital PAX machine (Siemens AG, Karlsburg, Germany) equipped with digital calipers accurate to 0.01 mm. One broken screw was found in a 33-year-old man who underwent an L4–S1 two-level TLIF for chronic back pain caused by degenerative disc disease (Fig. 2). At the 1-year follow-up visit he was asymptomatic and appeared to have a solid arthrodesis on CT scanning. He is also a smoker. There were no other instrumentation complications.

The mean preoperative disc height at each treated level (39 levels), measured at the posterior vertebral corner, was 5.2 mm (range 2.5–8.5 mm). In the case of spondylolisthesis, a tangential line along the inferior vertebral body was projected posteriorly to depict the estimated disc height without listhesis. Measured on standing radiographs obtained immediately postoperatively, the mean treated disc space height was 8.6 mm (range 4.5–10.5 mm), or +3.3 mm (range +2 to 5.8 mm). At the 3-month follow-up review, the mean disc height was 8.2 mm (range 4.4–10.5 mm), possibly indicating slight settling of the implant; however, this difference was not statistically significant (p = 0.31). At the last follow-up review (mean 12.4 months), the mean posterior disc height was unchanged at 8.1 mm (p = 0.42).

The bioabsorbable cages are not visualized on plain radiographs; therefore, the interbody fusion status can be accurately assessed. We considered 34 (87.2%) of 39 fusion levels to have a solid arthrodesis of the interbody space without instrumentation failure, as confirmed by the presence of continuous bridging bone observed on the lateral radiographs (Fig. 3). In the patient who had a broken pedicle screw, this was considered to be a pseudarthrosis. The mean breadth of the bridging bone also appeared to widen with time, although admittedly this is an inherently unreliable finding because each radiograph is not consistently obtained in an exact lateral projection. A sentinel sign, however, was present in only 10 (25.6%) of 39 levels, which is consistent with other series.

Findings on CT Scans

The CT scans in which sagittal and coronal 3D reconstruction was used were evaluated for bridging bone in the

<table>
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<td><strong>Number of TLIFs performed at each level</strong></td>
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<td><strong>Level</strong></td>
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* According to a study by Harms and Jeszenszky.
interbody space, as well as evidence of dislodgment, subsidence, and/or sterile abscess or reactive zone around the implant. Bridging bone was noted as early as the 3-month postoperative CT scan when obtained; however, this was not considered robust (Fig. 4). At the last follow-up visit, a continuous bridging bone was noted in 38 (97.4%) of 39 fusion levels. On coronal scans, this typically measured 6 to 9 mm in width per cage, consistent with the internal diameter of an 11- or 13-mm cage minus the 2-mm cage wall thickness. For the several patients with repeated scans, the fusion mass appeared to increase in width after 1 year of follow up.

The cages were easily identified on coronal images, but became less distinct on follow-up scans. On sagittal 3D reconstructions, there often appeared to be a small area of radiolucency at the caudad and cephalad ends of the implant; however, this was not directly measured when the plain radiographs were evaluated for settling of the implant. We believe that this finding is real, but minimal, as noted on the lateral radiographs. This may represent a fibrous margin that is present all around the implant.24 There were no instances of a destructive inflammatory reaction or sterile abscess formation.

For those patients who underwent repeated scanning, the fusion mass appeared to increase with time, whereas the disc space height remained stable. Additionally, other authors have found similar results with the use of BMP, increasing the rate of interbody fusion over time.7

**DISCUSSION**

The science of bioabsorbable implants continues to evolve as clinical evidence grows. This series is one such study, in which the early surgical and clinical results are available.

**Postoperative Complications**

With newer generation of composite polymers, early concerns about sterile abscess or sinus tract formation, osteolysis, allergic reactions, or hypertrophic fibrous encapsulation have been almost eliminated.2–4,20 This has occurred because of the use of poly-L-lactic acid stereoisomers.22 Consequently, the incidence of infection appears to be low, and is probably consistent with allograft or titanium interbody spacers (unpublished data). In our series there were no superficial or deep infections at a mean follow-up duration of 12.4 months. A quiescent fibrous tissue layer containing dispersed foreign body giant cells, however, has been noted in animal studies,24 and this also appears to be consistent with our postoperative CT scanning. Nonetheless, no device-related complications were noted.

**Radiographic Findings**

The radiographic fusion rate was 87.2% (34 of 39 fusion levels) at a mean of 12.4 months of follow up, and there was evidence of one instrumentation failure. This is consistent with other studies, in which radiographs have been used to determine fusion success.15,16,18,23 Although plain radiographs are considered to be inherently unreliable for assessments of fusion status, bridging bone was conclusively identified on the majority of radiographic images. On 3D CT reconstruction scanning, the fusion rate was considered to be 97.3% (100% for single-level fusions, eight cases). This is considerably better than in most series, as has been noted in multiple studies in the literature (mean 92% fusion success for single-level and 76% fusion success for two-level fusions). Notably, it is lower than the 98.9% overall fusion rate and the 100% fusion rate in two-, three-, and four-level PLIFs report by Brantigan, et al.6
Perhaps more importantly, there appears to be a fusion mass buttressing effect, which occurs in the bioabsorbable cages packed with rhBMP-2. This is most likely caused by the time-patterned resorption and controlled “dynamization” of the resorbable cage. Ideally, the fusion mass would have to be solid by 18 months, because the cage resorption is complete by 18 to 36 months. If buttressing does not occur, then perhaps an inadequate fusion mass will eventually fail with increased activity, or more likely the instrumentation will fail.

Investigators have found that a cross-sectional area of 30% is needed to achieve a solid interbody arthrodesis.10 Because our study involves early findings (mean 12.4 months of follow up), this has not been noted. In the case of the broken screw, the 1-year follow-up CT scans revealed a solid arthrodesis. Perhaps this is an example of uncontrolled resorption or dynamization; nonetheless, the patient fortunately attained fusion and is currently asymptomatic.

CONCLUSIONS

Bioabsorbable interbody spacers do not appear to lead to an increased incidence of complications, and they appear to possess ideal interbody fusion characteristics because their slow resorption, compatibility with postoperative imaging, and gradual transfer of anatomical loads to the developing fusion mass. Postoperative fusion success is excellent, as confirmed on CT scanning. The devices reported in this series are not approved by the Food and Drug Administration for use in the spine. These devices have been cleared by the Food and Drug Administration, however, to maintain the relative position of weak bone tissue in trauma and reconstructive orthopedic procedures when they are used in conjunction with traditional rigid fixation.

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