Internal fixation has become an invaluable adjunct in spine surgery, allowing for rigid fixation, higher fusion rates, improved alignment, and faster mobilization of patients. Nonetheless, spinal instrumentation is not without risks. All instrumentation should be properly considered an adjunct to fusion, because all implants are subject to fatigue and will eventually fail if bone fusion has not been achieved. Short-term risks, such as malpositioning or injury to adjacent anatomical structures during implant placement, are applicable to all forms of spinal instrumentation.

Long-term risks are associated with all types of permanent indwelling hardware. Conventional spinal instrumentation is made of stainless steel or titanium alloys, and has a number of disadvantages in clinical usage. The implants’ radiopacity makes evaluating fusion difficult, and makes some imaging modalities useless for clinical purposes. Because implants are foreign bodies, there is the persistent risk of infection. The great discrepancy between the elastic modulus of metal and cortical bone can lead to stress shielding of the spine, with the subsequent risk of telescoping, screw loosening, future instrumentation failure, device-related osteopenia, and possible fracture or instability. Also, particulate debris created by motion between metallic implants has been shown to lead to an inflammatory response in rabbit models of spinal fusion, and may theoretically be the cause of late-onset inflammation and osteolysis.

Nonresorbable polymeric devices offer the advantages of radiolucency, and also possess an elastic modulus that is closer to but still higher than that of cortical bone. Thus, stress shielding remains a significant issue. Furthermore,
the risk of particulate debris may be increased, and the long-term risks of a retained foreign body are still present.

To reduce these long-term risks, bioabsorbable spinal fixation devices are being developed. Biodegradable polymers were first used as internal fixation devices in 1966 for long-bone fractures, and have been in use for more than 30 years in the form of surgical sutures. These materials are now used in orthopedic procedures for fracture fixation, osteotomy stabilization, as interference screws in anterior cruciate ligament reconstruction, to reconstruct the skull base, and as plates in neurosurgical and maxillofacial procedures. These polymers are now being investigated for use in the spine.

Like permanent implants, bioabsorbable ones are designed to provide immediate support to the weight-bearing portions of the spine, as well as to provide support for osteoconductive and osteogenic grafts, so that a durable fusion can occur. While fusion is taking place, the implants are absorbed by the body and lose strength. An increasing load is placed on the autograft, preventing stress shielding. Bioabsorbable polymers are initially radiolucent, and because the implants ultimately will be resorbed, in the long term there will be no stress shielding, no particulate debris, and no retained foreign body.

Most research has involved implants made of combinations of the alphapolyesters PLA and PGA and their stereoisomers, which are all resorbed in the body and ultimately excreted as carbon dioxide and water. There are several stereoisomers of PLA available for use: PLLA characteristically has a higher crystallinity, strength, and longer periods of degradation than a second stereoisomer, PDLLA, which has lower strength and faster rates of degradation. The strength and absorbability of the copolymer can be significantly altered by the ratio of these two forms. These two isomers have been combined in a 70–30 ratio to PLLA to PDLLA, whose strength and degradation rate can be significantly altered by the ratio of these two forms.

In general, PLA copolymers degrade over a longer period of time and produce less inflammatory reaction than PGA alone. Nevertheless, in products made of both copolymers, as the relative amount of PLA increases the degradation time decreases. An implant can be tailored to degrade in a time period appropriate for the specific fusion application.

This study was completed in 1998 and 1999, and pre-dated more recent works conducted using off-the-shelf polymers originally designed for use in nonweight-bearing applications such as craniofacial reconstruction. The questions we wanted to address with this study included the following. 1) What is the rate of resorption of two different copolymers composed of different ratios of PDLLA and PGA? 2) Would bone grow despite the acidic breakdown products? 3) Would there be a significant inflammatory response to the devices or their breakdown products? 4) Would rhBMP-2 function in the presence of the acidic breakdown products?

MATERIALS AND METHODS

Experimental Animals

Fifty-four adolescent (1–2 years old) male goats underwent a two-level anterior cervical discectomy and fusion. All animals were cared for in a program accredited by the Association for the Assessment and Accreditation of Laboratory Animal Care International. This protocol was in compliance with the regulations established by the Institutional Animal Care and Use Committee of the University of South Florida.

Randomization of Animals

Each animal was randomly assigned to one of several cohorts. The control animals underwent a two-level anterior cervical discectomy and fusion with a tricortical iliac crest autograft; the others underwent fusion with either an 85:15 or 70:30 PLDLLA/PGA cage, which was in turn filled with either small fragments of fresh autograft or rhBMP-2.

Surgical Procedure

The animals were not allowed to eat for 24 hours before surgery; they were allowed to drink water freely. Antibiotic prophylaxis with 1 g cefazolin was administered intravenously to each goat before surgery.
Anesthesia was induced intravenously with 0.05 mg/kg atropine, 4 mg/kg ketamine, and 0.05 mg/kg xylazine. All animals were intubated, and anesthesia was maintained with 1 to 2% isoflurane mixed with room air.

The animals were placed prone, and the posterior superior iliac crest shaved, prepared with alcohol and betadine, and draped in a sterile fashion. In the eight control animals, a 1 × 2–cm tricortical iliac bone autograft was obtained; in the other 34 animals, approximately 4 cm³ of cancellous iliac crest bone was harvested to fill the cages. The iliac crest wound was closed in a multilayered fashion; no drains were used.

Each animal was then placed supine, and the anterior neck was shaved, prepared, and draped in a sterile manner. A longitudinal skin incision was made on the right side, and the standard anterolateral approach to the cervical spine was used, developing the plane with the sternocleidomastoid and carotid sheath laterally and the trachea and esophagus medially. The longus colli muscle was divided in the midline with monopolar cautery. The anterior aspects of the C2–3 and C3–4, C3–4 and C4–5, or C4–5 and C5–6 disc spaces were exposed. The anterior two thirds of the two exposed disc spaces were removed with a combination of curettes and rongeurs; no attempt was made to remove the posterior longitudinal ligament or to enter the spinal canal. Care was taken to remove the cartilaginous endplate and to decorticate partially both the superior and inferior bone endplates.

The copolymer cages were each packed with approximately 2 cm³ of cancellous bone autograft or rhBMP-2 and inserted into a slightly distracted interspace with the convex side facing posteriorly. Each wedge-shaped tricortical iliac bone autograft was similarly tamped into an interspace, with the cancellous side facing posteriorly. No internal fixation devices were used. The surgical sites were irrigated with sterile normal saline and closed in multiple layers in routine fashion. Animals received 0.005 mg/kg buphrenorphine intramuscularly to alleviate postoperative pain. The goats were fed freely, observed in a cage for 24 hours postoperatively, and then allowed to rejoin the herd. Animals were maintained without an orthosis and unrestricted activity was permitted. All goats quickly returned to normal activity and social interactions, including head butting.

Radiographs were obtained immediately after and at 1, 6, 12, and 24 weeks postsurgery. Animals received intravenous sedation with Versed before each radiograph, and were returned to the herd that day if not killed. Animals were sedated and humanely killed at 3, 6, or 12 months by intravenous administration of 80 mg/kg pentobarbital.

Interbody Device

The raw polymers 85:15 and 70:30 PLDLLA/PGA were commercially obtained and extruded into rods. The rods were milled to yield interbody devices that approximated the anterior, lateral, and vertical interbody space of the goat cervical spine. The anterior–posterior dimension was 12 mm, the maximum width was 15 mm, the anterior height 8 mm, and the posterior height 6 mm. A central 6-mm hollow chamber with four peripheral 1- to 2-mm hollow chambers were created for the bone autograft (Fig. 1).

All cages were sterilized with gamma irradiation before implantation.

Manual Stress Analysis

At 3, 6, and 12 months postoperatively, the animals were killed and their cervical spines were excised en bloc, removing all muscle but taking care to preserve all ligaments. Each spine was manually stressed by two observers who were blinded to the surgical procedure to assess the degree of motion at each of the treated interspaces compared with adjacent untreated interspaces. Each interspace was scored (Table 1) and assigned a grade of 2 if
rigid with no detectable flexion–extension motion, 1 if reduced in motion but not rigid, or 0 if typical in flexion–extension motion compared with adjacent untreated interspaces. A researcher not involved with the surgical procedures evaluated the spines manually. Because this was a pilot feasibility study, simple physical examination was used to stress the spinal segments. Any research in which it is concluded that such a device is suitable for human testing would require detailed biomechanical analysis of the implants in both in vivo and in vitro settings.

Radiographic Analysis

Radiographs obtained immediately before planned death were scored (Table 1) by an observer blinded to treatment group, and each interspace was assigned a grade of 2 if a transvertebral osseous bridge was present, 1 if new fragmented osseous densities were evident, or graded 0 if no osseous densities had formed. A researcher not involved with the surgical procedures evaluated the radiographs.

Histological Analysis

An observer working in a blinded fashion grossly examined the excised cervical spine of each animal. Each spine was then cut along the coronal plane, beginning on the ventral surface, and fixed in 10% neutral buffered formalin. To aid in subgroup examination, one fused interspace obtained in a single animal from each of the two experimental groups was divided along the sagittal plane, beginning on the right lateral side. Whenever subgross sectioning revealed cage remnants or degradation debris, their position was noted. After fixation, each treated interspace was trimmed, decalcified, and embedded in paraffin. Contiguous thin sections were stained with H & E. Specimens were examined qualitatively to assess the extent of growth of new trabecular bone from each end-plate, remodeling of the bone autograft, and presence of residual polymeric debris or inflammatory cells. Each interspace was scored histologically (Table 1) based on the degree of new bone formation. A grade of 3 was assigned if a marked amount of newly formed bone was present, 2 if a moderate amount was present (Fig. 2 left), 1 if minor new bone was present (Fig. 2 right), or 0 if no new bone had formed.

Composite Grade

The three grades assigned according to the manual strength, radiographic, and histological analyses were added for each level. An interspace with a manual stress grade of 2, a radiographic grade of 2, and a histological grade of 3 received a composited grade of 7, which was judged to be a stable union. The histological nature of that union (fibrous compared with bone tissue) and the accompanying complications were noted for each interspace.

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**RESULTS**

All animals that underwent the surgical procedure experienced no neurological, infectious, or other sequelae. All goats were standing, ambulating, and had resumed eating within 24 hours of surgery. There were no clinical indications of local or systemic infection, nor was there any behavior consistent with pain or discomfort after the first 24 hours postoperatively. All animals resumed normal activities once returned to the herd, including head butting and jumping. All statistics were calculated using a personal computer running commercially available software (The Statistics Calculator; StatPac, Inc., Minneapolis, MN).

**Gross Evaluation**

Neither solid cages nor functional remnants (long fragments that were still in contact and embedded in both superior and inferior vertebral bodies) of the 85:15 PLDLLA/PGA cage were present at 3 or 6 months postimplantation. The 70:30 PLDLLA/PGA cage showed no signs of absorption or degradation at either 3 or 6 months, and appeared to be intact in all specimens. No remnants of either cage were found anterior to the vertebral bodies, indicating that there were no cage extrusions. A similar finding was noted in four animals followed for 12 months after placement of a 70:30 cage filled with rhBMP-2 (no other group of animals was studied for 12 months).

Gross inflammatory debris was found around the 85:15 PLDLLA/PGA cages at 3 months; less debris was qualitatively present at 6 months. Minimal inflammatory debris was present around the 70:30 PLDLLA/PGA cages at 3 months, and none was found at 6 months. There was no gross evidence of inflammation or granuloma formation in the control animals. We found no evidence of overt or gross inflammation or fibrous adhesions in the soft tissues of the neck or in the perivertebral area in any animal.

**Manual Stress Analysis**

Each cervical spine was manually stressed, and the treated levels were compared with untreated, intact levels. All values are given as the mean ± standard error of the mean. At 3 months, the control spines were assigned a grade of 1.1 ± 0.4 (seven interspaces in four goats); the spines treated with 85:15 PLDLLA/PGA cages containing autograft and rhBMP-2 were graded 0.9 ± 0.6 (10 interspaces in five goats) and 1.3 ± 0.5 (eight interspaces in four goats), respectively; the spines treated with 70:30 PLa cages filled with autograft (four interspaces in two goats) and rhBMP-2 (eight interspaces in four goats) were all graded 1 ± 0. (Table 2). There were no significant differences between the five animal cohorts. All spines exhibited more stiffness compared with an untreated intervertebral space, although with some residual flexion and extension.

As detailed in Table 3, at 6 months, the control spines were assigned a grade of 0.9 ± 1 (seven interspaces in four goats); the spines treated with 85:15 PLDLLA/PGA cages were graded 1.3 ± 0.7 when the device was filled with autograft (10 interspaces in five goats), and 0.8 ± 1.5 when it was filled with rhBMP-2 (eight interspaces in four goats); the spines treated with 70:30 cages were graded 0.4 ± 0.5 when the device was filled with autograft (eight interspaces in four goats), and 1.8 ± 0.5 when it was filled with rhBMP-2 (eight interspaces in four goats).

**Radiographic Evaluation**

New intervertebral osseous densities (Grade 1 or 2) were present in five of seven control animals at 3 months, for a mean grade of 0.9 ± 0.7 (seven interspaces); the grades for the 85:15 cages with autograft and rhBMP-2 were 1.3 ± 0.5 (10 interspaces) and 0.6 ± 0.5 (eight inter-

**TABLE 2**
The mean grades in the goats killed 3 months after treatment*

<table>
<thead>
<tr>
<th>Cohort</th>
<th>Manual Stress</th>
<th>Radiographic</th>
<th>Histological</th>
<th>Composite</th>
</tr>
</thead>
<tbody>
<tr>
<td>controls</td>
<td>1.1 ± 0.4</td>
<td>0.9 ± 0.7</td>
<td>1.3 ± 0.8</td>
<td>3.3 ± 1.7</td>
</tr>
<tr>
<td>85:15 cages</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>w/ autograft</td>
<td>0.9 ± 0.6</td>
<td>1.3 ± 0.5</td>
<td>1.6 ± 0.7</td>
<td>3.8 ± 1.5</td>
</tr>
<tr>
<td>w/ rhBMP-2</td>
<td>1.3 ± 0.5</td>
<td>0.6 ± 0.5</td>
<td>1.3 ± 0.5</td>
<td>3.1 ± 1.3</td>
</tr>
<tr>
<td>70:30 cages</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>w/ autograft</td>
<td>1.0 ± 0.0</td>
<td>1.1 ± 0.4</td>
<td>1.2 ± 0.5</td>
<td>3.8 ± 1.0</td>
</tr>
<tr>
<td>w/ rhBMP-2</td>
<td>1.0 ± 0.0</td>
<td>1.4 ± 0.5</td>
<td>1.4 ± 0.5</td>
<td>3.8 ± 0.9</td>
</tr>
</tbody>
</table>

* There were no significant differences between cohorts. All values are given as the mean ± standard error of the mean. Abbreviation: IS = interspace.

**TABLE 3**
The mean grades in the goats killed 6 months after treatment

<table>
<thead>
<tr>
<th>Cohort</th>
<th>Manual Stress</th>
<th>Radiographic</th>
<th>Histological</th>
<th>Composite</th>
</tr>
</thead>
<tbody>
<tr>
<td>controls</td>
<td>0.9 ± 1.0</td>
<td>1.3 ± 0.5</td>
<td>1.4 ± 0.8</td>
<td>3.7 ± 2.1</td>
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<tr>
<td>85:15 cages</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>w/ autograft</td>
<td>1.3 ± 0.7</td>
<td>1.3 ± 0.8</td>
<td>2.0 ± 0.8</td>
<td>4.6 ± 2.2</td>
</tr>
<tr>
<td>w/ rhBMP-2</td>
<td>0.8 ± 1.5</td>
<td>1.5 ± 0.5</td>
<td>1.6 ± 0.5</td>
<td>3.9 ± 1.1</td>
</tr>
<tr>
<td>70:30 cages</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>w/ autograft</td>
<td>0.4 ± 0.5</td>
<td>1.3 ± 0.5</td>
<td>1.4 ± 0.5</td>
<td>3.0 ± 0.8</td>
</tr>
<tr>
<td>w/ rhBMP-2</td>
<td>1.8 ± 0.5</td>
<td>1.9 ± 0.4</td>
<td>2.6 ± 0.5*</td>
<td>6.3 ± 1.2*</td>
</tr>
</tbody>
</table>

* The histological and composite grades of the 70:30 PLDLLA/PGA cages filled with rhBMP-2 were significantly (p < 0.05) higher than those of the other cohorts, except for the 85:15 cages filled with autograft. See text for details.
spaces), respectively. The grades in levels fused with 70:30 cages filled with autograft or rhBMP-2 were 1.1 ± 0.4 (eight interspaces) and 1.4 ± 0.5 (eight interspaces), respectively. The spines treated with 85:15 cages filled with rhBMP-2 showed significantly less radiographically confirmed fusion than spines treated with 85:15 cages filled with autograft, or those treated with 70:30 cages filled with rhBMP-2 (p < 0.05, ANOVA). There were no other significant differences between cohorts (Table 2).

At 6 months, the mean radiographic grade in the control cohort was 1.3 ± 0.4 (seven interspaces). The grades in the 85:15 cohorts with autograft and rhBMP-2 averaged 1.3 ± 0.8 (10 interspaces) and 1.5 ± 0.5 (eight interspaces). The mean radiographic grades in the groups treated with 70:30 devices filled with autograft or rhBMP-2 were 1.3 ± 0.5 and 1.9 ± 0.4 (eight interspaces in each cohort), respectively (Table 3).

Although the mean radiographic grades were comparable in all groups, as expected, these grades were higher in animals examined at 6 months. Furthermore, in all animals but one in the cohort that underwent fusion with the 70:30 PLDLLA/PGA cage and rhBMP-2, the radiographic grade was 2, meaning that the fusion mass crossed the interspace. Osseous bars were seen both within and anterior to the cage in this goat. The group treated with 70:30 cages and rhBMP-2 demonstrated significantly more radiographically confirmed fusion than those treated with 85:15 cages containing autograft (p < 0.05, ANOVA), and trended to more fusion (0.05 < p < 0.10) than control animals and the group treated with 70:30 PLa cages containing autograft.

**Histological Evaluation**

For all cohorts at 3 months, fibroplasia was the principal histological feature (Fig. 3). Connective tissue was interposed between vertebral endplates, which in these cases prevented development of a solid bone union. This deposit was composed of fibrous tissue surrounded by numerous macrophages, epithelioid cells, scattered lymphocytes, and rare multinucleated giant cells. Underlying this tissue was new bone, which had formed along the endplates of each interspace peripheral to the fibrous tissue. Osteoid was deposited around bone trabeculae and islets of calcifying cartilage, mostly peripheral to the fibrous tissue, outside the cages. The 70:30 cages were intact; there were a few solid remnants of the 85:15 cages remaining in the interspaces, but no functional fragments. A scant inflammatory reaction was noted adjacent to the cages and their remnants, which were solidly incorporated into the surrounding tissue. The cages and their remnants had to be manually removed from the tissue before it could be prepared for histological analysis.

The histological pattern was different in animals examined at 6 months. In the animals whose interspaces were treated using tricortical autograft, copolymer cages (both 85:15 and 70:30) filled with autograft, and 85:15 cages filled with rhBMP-2, there was quantitatively less fibrous tissue, and increased bone formation, than was found in the animals killed at 3 months (Fig. 4). The new bone was growing from both vertebral endplates. Again, osteoid was deposited around bone trabeculae and islets of calcifying cartilage, but hypertrophic osteoblasts were found surrounding long spicules of bone. In the cohort of animals that underwent fusion with 70:30 cages and rhBMP-2, the primary histological reaction was of a fibrocartilaginous bone union extending from the superior to the inferior bone ridge, both inside and outside the cage (Fig. 5). Again, the 70:30 cages were intact, whereas there were no remnants of the 85:15 cages.

Despite the increased amount of bone formation in these latter groups, the mean histological grade of new bone formation was similar and not significantly different from control values.
between the animals killed at 3 and 6 months, which ranged from 1.3 to 2 (the complete results are described in Tables 2 and 3). Only the cohort that underwent fusion with 70:30 PLDLLA/PGA cages filled with rhBMP-2 (Fig. 6) and killed at 6 months, one of seven control interspaces was judged to have a stable union, whereas none of the other interspaces had a union. The mean composite grade for all five cohorts was between 3.1 and 3.8, with no significant differences (Table 2). At 6 months, one of seven control interspaces was judged to be rigid (Grade 2) on manual inspection, to be connected by a new osseous intervertebral bridge (Grade 2) on radiographic studies, and to have marked new bone formation (Grade 3) on histological investigations, for a composite grade of 7. That interspace had successfully arthrodesed, and was bridged by bone without intervening soft tissues or inflammation.

Of the interspaces fused with cages filled with autograft, three (30%) of 10 of the spines treated with 85:15 devices were graded 7, whereas none of the spines treated with 70:30 PLDLLA/PGA cages were graded 7. Of the interspaces treated with cages filled with rhBMP-2, none of the spines treated with 85:15 PLDLLA/PGA cages were graded 7, whereas five (63%) of eight of the spines treated with 70:30 cages were graded 7. A union consisting primarily of bone stabilized each of the five cage-implanted interspaces graded 7. The mean combined grade at 6 months showed no significant differences between any of the cohorts, whose grades ranged from 3 to 4.6 (Table 3), except for the graph treated with 70:30 cages filled with rhBMP-2, which the goats had a mean composite grade of 6.3 ± 1.2 (p < 0.05, ANOVA).

**DISCUSSION**

Degradation of both PGA and PLA occurs through non-specific hydrolytic cleavage of their ester bonds. As it breaks down, PGA is degraded into glycolic acid and glyoxylic acid, which may be excreted in the urine, or further converted to glycine, then to serine, and finally to pyruvic acid, which enters the tricarboxylic acid cycle, and is excreted as water and carbon dioxide. Although PGA is degraded rapidly, in approximately 2 to 4 months, PLA degradation, in contrast, takes several years. As it breaks down, PLA is degraded to lactic acid, which is further converted to pyruvic acid, and again enters the tricarboxylic acid cycle. The rapid absorption of PGA has been noted to be the cause of the stronger inflammatory response observed with this material as opposed to PLA.

The absorption rate of biopolymers is influenced by multiple factors. The ratio of the different polymers and stereoisomers in each compound has a profound effect on the absorption rate. In mixed PLA/PGA implants, decreasing the amount of PLA will (counterintuitively) tend to increase the degradation rate. The polylactides have the capacity to be molded into various shapes and sizes, which can also significantly affect the degradation rate. A higher surface/volume ratio of a biodegradable construct will result in more rapid degradation. Thus, a
screw, with a higher surface/volume ratio, will be absorbed faster than an interbody device made of the same material. The relative environment will also significantly affect the absorption of such devices. Faster degradation rates have been found in liquid environments and at higher temperatures. Materials have sometimes been found to absorb faster in vivo than results of in vitro studies had led one to expect. The process by which the implantable device is manufactured and sterilized may also influence the absorption rate. In previous studies it has been reported that bone can be formed in both nonspinal and spinal applications. Thus, osteogenesis can occur in the acidic environment created by the breakdown of these polymers.

In this study, placement of a bioabsorbable cage filled with either autograft or rhBMP-2 led to similar stable interbody union rates as in interspaces fused with tricortical autograft. In contrast to the bone fusion elicited by the autograft, the cages initially elicited a primarily fibrous response at 3 months after implantation, with minimal new bone deposit at the periphery of the fibrous mass. At 6 months after implantation, the fibrous mass was being replaced by new bone, which was growing into the inter-space from both vertebral bodies. This bone fusion was occurring both inside and outside the cage, indicating that an initial fibrous union will not necessarily prevent a final bone union from occurring. The 70:30 PLDLLA/PGA cages filled with rhBMP-2 elicited more bone tissue in the fusion histologically than the other cohorts, and also led to a stronger fusion mass than in the other cohorts, including the controls. Bone morphogenetic protein has been previously shown to lead to a stronger, more rapid fusion than in controls when titanium cages are used; our results indicate that this is true when rhBMP-2 is used with more slowly dissolving cages as well. None of the cages extruded, despite the fact that no orthoses were used, nor was internal fixation, and the goats engaged in head-butting activity. Nevertheless, this activity may account for why our fusion rates are lower than reported previously in the literature.

Further studies are needed to determine if these cages are suitable for clinical use, but recent animal and clinical studies are promising. Toth et al. studied HYDROSORB (70:30 PLLA/PDLLA) cages filled with autograft or rhBMP-2 that were implanted in the sheep lumbar spine, with follow-up durations of up to 24 months post-surgery. Biomechanical testing in their study revealed that the fused spines were stiffer than nonfused spines. Histological, radiographic, and Faxitron high-resolution radiographic studies revealed that bone fusion was occurring between the superior and inferior vertebrae through, anterior to, and sometimes posterior to the cage, and that eventually the separate fusion masses were extending through the anterior margins of the device, indicating that bone was replacing the degrading cage. There was a trend of increased fusion stiffness and of radiographically and histologically confirmed fusion over time. At 6 months, in two of four animals fusion was confirmed histologically, in four (67%) of six it was confirmed at 1 year, and five (100%) of five exhibited fusion at more than 1 year. Only a mild-to-moderate inflammatory response was present.

Wuisman, et al. studied PLLA interbody devices in goat lumbar spines, with a follow-up duration of 36 months. Their results showed that there is an increased rate of bone formation in the PLLA cages compared with titanium cages, indicating that there is stress shielding associated with those devices. The rate of bone formation was not constant; it increased when the PLLA cages began to degrade. The cages disintegrated at 1 year postsurgery, and were replaced by both bone and fibrous tissue. Of all the animals followed for 1 year or longer, 15 (88%) of 17 goats treated with PLLA cages had a bone fusion, whereas only 2 (66%) of three of those that underwent treatment with titanium cages attained adequate fusion. The difference was significant. All cages had disintegrated at 12 months, and were completely gone at 24 months. A mild inflammatory response was seen, which had disappeared by 36 months.

Van Dijk, et al. studied PLLA cages used to treat goat lumbar spine, and followed up for 3 years. They found that 19 (86%) of the 22 spines treated with the PLLA cages appeared to be fused on radiographic studies, which was a significantly higher rate than the two (33%) of six spines that attained fusion with titanium cages. The PLLA cages disintegrated at 12 months postsurgery, and the disintegrating cages were found to be partially replaced by trabecular bone.

This study is, besides our initial pilot study, the only one to address the use of biodegradable cages in the goat cervical spine. In the pilot study, we evaluated 85:15 PLLA/PGA cages packed with autograft bone, and found that at 12 weeks postsurgery, a primarily fibrous, yet stable union had formed. We concluded that the use of bioabsorbable materials held promise, but that a cage producing less of an inflammatory reaction might be more suited for internal spinal fixation. In this study, we used cages composed of 70:30 PLDLLA/PGA, which has been shown to elicit a smaller immune response. We found that the fusions were initially fibrous, but at the 6-month follow-up review the fibrous tissue was replaced by spicules of bone. Fusion was occurring both within and anterior to the cage. There was no sign of cage degradation at 6 months, and a minimal inflammatory response was present. There were no cage extrusions, and no signs of infection or other sequelae as a result of the use of these cages.

Future studies should follow the animals for a longer period to determine when the cages degrade. Because the 70:30 PLDLLA/PGA cages are still intact 12 months after placement, future studies should be continued beyond that point. Histological and radiographic studies should examine where bone union occurs: inside the cage, outside the cage, or both. Further investigations should examine whether the spaces left by cage degradation are filled in by bone or fibrous tissue, or if permanent voids are left between fused vertebrae. Biomechanical testing performed at various stages after cage implantation would be used to evaluate how strong the treated spine is over time.

Our findings indicate that the short-term union results of bioabsorbable cages in the goat cervical spine are similar to those seen when tricortical iliac crest autograft is used. These cages supplied sufficient stability to the spine during the load-bearing period immediately postsurgery; in all treated cohorts the results of histological, radiographic, and manual strength testing were similar to or

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higher than in the control animals, and there were similar or higher rates of final interbody fusion.

In this pilot study we demonstrated the feasibility of using bioabsorbable cages in the cervical spine for anterior arthrodesis and interbody fusion after discectomy.

SUMMARY OF FINDINGS

The 85:15 PLDLLA/PGA cages resorb too quickly to be functionally useful, whereas the 70:30 cages are still functionally intact and unresorbed at 6 and 12 months postimplantation. There is relatively little inflammatory response to the intact cages. There is some inflammatory response to the degradation products of disintegrating cages, but it appears to be clinically insignificant. Bone will grow in the presence of the cages and their breakdown products. The rhBMP-2 appears to be functional in the presence of the cages and their breakdown products, and may even lead to superior arthrodesis than autologous bone.

Future studies must address when and how 70:30 PLDLLA/PGA cages resorb. Biomechanical studies will address the stability and mechanics of the cages at implantation, and at various times thereafter. Results of histological and microradiographic analysis will determine whether and when the voids left by cage resorption are filled in with bone or inflammatory tissue. Finally, the 70:30 bioabsorbable cage must be compared with more recently reported cervical polymeric devices. Ultimately, future studies in this field must address whether bioabsorbable devices offer any clinically significant advantages over nonresorbing polymeric devices and traditional metal devices for interbody fusion, and if so, which polymer in what configuration is best.

CONCLUSIONS

There are numerous theoretical advantages in using a bioabsorbable cage in the cervical spine after discectomy. Radiolucency allows for better evaluation of arthrodesis on follow-up images, and does not interfere with or preclude the use of computerized tomography or magnetic resonance imaging studies to evaluate the spine in future investigations. By adjusting the composition of such devices, it should be possible to develop an implant that would provide immediate stability to the spine after surgery, but would slowly dissolve, allowing the spine to take on more weight, and preventing stress shielding and subsequent fusion failure. These devices would reduce the long-term risks of hardware extrusion, and thus reduce risks to adjacent organs and the need for surgery to remove hardware.

These devices must be subjected to the same controlled comprehensive evaluations given to titanium devices. They should be tested biomechanically, radiographically, microradiographically, and histologically in animal and cadaveric spines at various intervals after implantation to evaluate their strength, the adequacy of bone fusion, and the extent of inflammatory reactions. Only after these devices are better understood quantitatively based on investigations conducted in animals in vivo and in cadaveric models in vitro, should clinical studies begin.

Dedication

This paper is dedicated to the memory of Dr. David W. Cahill, who was a colleague, mentor, and friend.

References


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