Management of patients diagnosed with lumbar spinal stenosis and disc degeneration undergoing transforaminal lumbar interbody fusion using a novel ceramic implant with one year follow up.

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Abstract

This case report focuses on the first clinical and radiographic outcomes of 2 patients with 1-year follow-up with a novel ceramic interbody implant which allows bony growth to occur onto the corresponding vertebral bodies as well as through the implant. Historical reports of transforaminal lumbar interbody fusion (TLIF) reflect good clinical and radiographic outcomes. During the procedure, a structural support is placed within the middle or anterior aspect of the disc space accompanied by pedicle screw fixation. Two patients underwent a TLIF procedure with posterior spinal fusion instrumentation and received the Valeo® TL (Valeo® TL Lumbar Interbody Fusion Device, Amedica® Corporation, Salt Lake City, UT) implant and were evaluated at 1-year for clinical and radiographic outcomes using CT scans and dynamic x-rays.

Length of hospital stay was 2 days in the hospital without complications. At 1-year, a CT scan and dynamic x-ray demonstrated solid interbody fusion and solid posterolateral fusion. Bone appeared to be well formed to the Valeo® TL implant as well as through and behind the implant in the interbody space; radiographic imaging was not distorted by the implant. These findings suggest that the Valeo® TL implant with its proprietary ceramic promotes bony growth on the implant; making the implant less likely to migrate as fusion occurs to the host vertebral bodies at the same rate as fusion occurs through the implant itself.

PEEK and carbon fiber implants may still migrate, despite concomitant posterior fixation, as there is no ability for bony ingrowth to the native vertebral bodies. These preliminary results suggest that the Valeo® TL implant may provide an attractive alternative for use in a TLIF procedure as compared to standard titanium, PEEK or carbon fiber grafts. The need for a larger prospective study to determine the efficacy of the use of Valeo® TL implant is warranted.

Introduction

Historically, reports of transforaminal lumbar interbody fusion (TLIF), first described by Blume2 and popularized by Harms and Rolinger8, reflect good clinical and radiographic outcomes. In the TLIF procedure, a structural support (allograft bone or assorted cage designs) is placed within the middle or anterior aspect of the disc space via a posterolateral transforaminal route accompanied by pedicle screw fixation. Many interbody grafts have been created that not only focus on restoring disc height but also maintain lordosis through the vertebral segment, create distraction and restore the normal weight distribution within the anterior column. Aside from the mechanical properties, the graft must also have osteoconductive properties in order to promote fusion. Although, most grafts do allow bony growth through the implant, this case report represents the first clinical experience with a novel ceramic graft that allows growth onto the implant as well as through the implant. The proprietary ceramic, composed of medical-grade silicon nitride with a special textured surface, allows such bony ingrowth to occur to the corresponding vertebral bodies.

Historically, imaging of these implants has been a challenge due to metallic scatter and distortion with sophisticated imaging such as MRI and CT. An abstract presented at the 8th Annual Spine Arthroplasty Society Global Symposium on Motion Preservation Technology in May 2008 compared the clinical visibility of cylindrical shaped specimens composed of cobalt chromium, PEEK, titanium and silicon nitride ceramic. As compared to the other commercially available specimens, the study concluded that silicon nitride ceramic implants may be easier to follow postoperatively from lack of distortion under magnetic resonance and lack of scattering under computed tomography (Anderson M, Bernero J, Brodke D: Medical imaging characteristics of silicon nitride ceramic: a new material for spinal arthroplasty implants. Abstract presented at the 8th Annual Spine Arthroplasty Society Global Symposium on Motion Preservation Technology, Miami, FL, May 2008).

This paper presents the clinical and radiographic outcomes of two patients that received the Valeo® TL (Valeo® TL Lumbar Interbody Fusion Device Amedica Corporation, Salt Lake City, UT) implant with one year follow up. This implant offers a dense, hydrophilic construct for load bearing, easy insertion and can be packed with autograft. As a non-oxide ceramic implant, it is strong, exhibits high fracture toughness, is hydrophilic and allows bony growth through the implant as well as onto the implant.
Case Report One

History and Physical Examination
A 47-year old woman reported back pain and weakness in her legs. The patient was diagnosed with lumbar spinal stenosis at L4-L5 with neurogenic claudication and axial low back pain at L4-L5, which failed nonoperative efforts. The patient was offered operative intervention in the form of a posterior decompressive laminectomy and posterior spinal fusion at L4-L5 using pedicle screw instrumentation and TLIF technique with application of interbody device, bone marrow aspirate, allograft bone and bone morphogenetic protein, as well as local bone.

Radiographic Imaging
A magnetic resonance imaging (MRI) scan of the lumbar spine was obtained. The MRI showed progressive disc herniation at L4-L5, moderate lateral recess stenosis, severe facet arthropathy, Modic changes within the vertebral bodies of L4 and L5 and a new central extrusion superimposed on a previously central disc protrusion (Figure 1).

Anterior-posterior (AP), lateral and flexion-extension x-ray views of the lumbar spine were also obtained. The x-rays showed increased angulation across the disc space but no evidence of any significant listhesis (Figure 2).

Preoperative Management
The patient was treated conservatively with physical therapy and oral medications. Due to the lack of improvement, treatment proceeded with diagnostic and therapeutic bilateral L5 transforaminal epidural injections. The injections were repeated in 2 months with no resolution of pain.

Operative Management
A standard decompression, including laminectomy, facetectomy, posterior spinal fusion (PSF) with instrumentation and TLIF was performed. Bone morphogenetic protein (Medtronic INFUSE® Bone Graft, Minneapolis, MN) was placed in the implant. Local autograft harvested from lamina and spinous processes was placed posterior to the implant and in the disc space.

Postoperative Course
Length of stay was 2 days in the hospital without complications. Postoperative x-rays were obtained demonstrating excellent placement of the implants, excellent restoration of the disk space height and adequate placement of the pedicle screws (Figure 3).

At two-weeks postoperatively, all preoperative symptoms were resolved and the patient was walking over 2 miles a day. The patient was treated with a brace for 6-weeks postoperatively and started physical therapy shortly after being weaned off the brace. At one year follow-up, a CT scan and dynamic x-rays demonstrated solid interbody fusion and solid posterolateral fusion screws in expected position contained within the pedicles (Figure 3, 4). Bone appears to be well formed to the Valeo® TL implant as well as through and behind the implant in the interbody space.
Case Report Two

History and Physical Examination
A 77 year old female with low back pain and bilateral leg pain, left greater than right. The patient was diagnosed with lumbar spinal stenosis L4-L5 and disc space collapse L4-L5, which failed nonoperative efforts. The patient was offered operative intervention in the form of posterior decompressive laminectomy and posterior spinal fusion at L4-L5 with TLIF and pedicle screw instrumentation at L4-L5.

Radiographic Imaging
An MRI scan showed 2 levels of stenosis, most severe on the right at L4-L5 with disc bulging, disc osteophyte complex, asymmetric disc collapse and severe stenosis at the L4-L5 level on the right. At L2-L3, the patient had some moderate central stenosis due to diffuse disc bulging (Figure 5).

Preoperative Management
The patient was treated conservatively with physical therapy and oral medications. Due to the lack of improvement, treatment proceeded with diagnostic and therapeutic bilateral L5 transforaminal epidural injection. The patient failed non-operative efforts and was offered operative intervention.

Operative Management
Similar to the first case report, the patient underwent a standard decompression, including laminectomy, facetectomy, posterior spinal fusion (PSF) with instrumentation and TLIF was performed. Bone morphogenetic protein (Medtronic INFUSE® Bone Graft, Minneapolis, MN) was placed in the ceramic implant. Local autograft harvested from lamina and spinous processes placed posterior to the implant and in the disc space.

Postoperative Course
Length of stay was 2 days in the hospital without complications. At two-weeks postoperatively, the patient had some mild left leg pain, which decreased gradually. The patient was treated with a brace for 6-weeks postoperatively and started physical therapy shortly after being weaned off the brace.

At 14 months, dynamic x-rays and a CT scan both demonstrated an appropriately aligned L4-L5 fusion site with no hardware failure or other complication from the surgery. The interbody implant was correctly positioned at this level and showed good incorporation with osseous bridging through the center of the graft (Figure 6,7). Bone appears to be well formed to the Valeo® TL implant as well as through and behind the implant in the interbody space. There is no distortion of the imaging due to the implant.
Discussion

Transforaminal lumbar interbody fusion has emerged as a successful procedure for treating lumbar spinal stenosis and disc degeneration. Compared to posterior lumbar interbody fusion, historic literature presents findings that suggest successful outcomes after TLIF, low morbidity rates, decreased operative time, decreased operative blood loss, decreased incidence of dural tears and fusion rates as high as 92%. The TLIF technique can also provide immediate postoperative stability and correction of anatomical deformities. The TLIF procedure also leads to increased surface area for fusion since it spares the contralateral lamina, facet and pars.

Considering the advantageous features of the TLIF technique, the design of synthetic interbody graft chosen for the procedure may potentially affect the clinical success of the procedure. Originally, titanium cages were placed in the interbody space during a TLIF procedure. These cages lead to subsidence through the vertebral body endplates, especially in osteoporotic patients. Post-operative imaging was also challenging with the titanium interbody implants as the titanium results in significant image distortion making fusion determination challenging. With the problems associated with the use of metal cages, nonresorbable polymers, such as PEEK or carbon fiber reinforced PEEK (CFRP), were developed.

A large case series found that the rates of collapse, slippage and graft migration associated with the use of CFRP and PEEK cages were found to occur at rates of 3-10%. Shortly after CFRP implants were popularized for use in TLIF procedures, porous ceramics were being developed for use in the TLIF procedure. These ceramics are known to be safe, allergy free and are associated with a high bone-bonding capacity. Most ceramics are attributed with osteointegration and osteoconductive properties. However, they do lack osteoinductive properties that are associated with allograft or autograft bone. The ceramic grafts are dependent on the remaining bone for their successful outcome. Literature has tried to focus on providing this capacity to ceramic implants by adding bone marrow cells or in addition to osteoinductive proteins to improve their osteoinductive capabilities.

The findings in this paper suggest that the Valeo® TL implant promotes bony growth onto the implant and through the implant. Due to the unique qualities, and better imaging compatibility, both patients demonstrated solid bony fusion on dynamic x-rays and CT scans at 1-year follow up. Our findings suggest that the Valeo® TL implant may provide an attractive alternative for use in a TLIF procedure as compared to standard titanium and CFRP implants.

Many studies have found that PEEK and CFRP implants may still migrate despite appropriate placement within the interbody space, as there is no ability for bony ingrowth to the native vertebral bodies. Aoki et al. presented a report of three patients with PEEK cage migration after a TLIF procedure. In this study, the authors suggested that surgeons use as much bone graft as possible, along with the PEEK cage, when packing the disc space; thereby, facilitating bony fusion prior to an event of cage migration. Ceramic implants, however, may not have this issue as there is bone adherence to the implant occurring at the same rate as the interbody fusion itself; therefore, the implant is less likely to migrate.

These preliminary results suggest that the Valeo® TL implant may provide an attractive alternative for use in a TLIF procedure as compared to standard titanium, PEEK or CFRP implants. The need for a larger prospective study to determine the efficacy of the use of Valeo® TL implant is warranted.
References


19. Valeo ceramic interbody implant, excellent restoration of the disk space but no evidence of any significant listhesis.

20. Valeo ceramic interbody implant, as well as through the implant and behind the implant in sagittal (C, D) show bone appearing to be well formed to the implant, with disc bulging, disc osteophyte complex, asymmetric disc collapse and severe stenosis at the L4-L5 level on the right. At L2-L3, the patient had some moderate central stenosis due to diffuse disc bulging.

21. Valeo ceramic interbody implant, excellent restoration of the disk space height and appropriate placement of the pedicle screws.

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Disclosure
The following author acknowledges a potential conflict of interest: JAY, royalties, stock options from Amedica® Corporation, Salt Lake City, UT.