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Minimal Access Expandable Mesh Device for Transforaminal Lumbar Interbody Fusion: A Case Series

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ABSTRACT
Bone-sparing techniques for transforaminal lumbar interbody fusion (TLIF) are promoted to help maintain spinal stability and to minimize operative time. We present a series of seven patients who underwent TLIF with use of an interbody expandable mesh device and with supplemental instrumentation. This device is deployed and filled with bone through a small cannula. All patients experienced pain relief and suffered no complications. Our results support the data from other centers which have performed similar procedures with this device.

Keywords: transforaminal lumbar interbody fusion, OptiMesh, deployable mesh, minimal access

INTRODUCTION
Lumbar interbody devices can be difficult to insert from a posterior approach due to their sizes and shapes, especially when the disc space is significantly collapsed (Figure 5). In addition, stabilizing posterior elements are removed in order to create a clear corridor to the disc space. A less invasive approach to the disc space involves the placement of a collapsed, expandable mesh pouch (OptiMesh by Spineology, St. Paul, MN). The pouch is gradually filled with increasing amounts of allograft bone chips through the small portal. (Figure 1) The mesh itself is made out of polyethylene terephthalate, a non-absorbable and pliable material commonly used in vascular grafts. We describe a series of patients who underwent this procedure with supplemental instrumentation.

Spineology OptiMesh graft containment system has been approved for the treatment of stable vertebral body defects. The device has also been used in the disc space for interbody fusion.2,5,6,7,8,9 The OptiMesh portal, despite its low profile, allows for an extensive discectomy and with preparation of the endplates, especially when the cannulas are introduced bilaterally. The bone chips are packed into the mesh until good filling of the disc space is noted on fluoroscopic imaging.

CASE PRESENTATIONS
From August 2016 through January 2019, seven patients underwent lumbar transforaminal interbody fusions by the senior author using the mesh containment device with supplemental instrumentation, either interspinous clamps (Patients 2, 3 and 5) or bilateral pedicle screws/rods (Patients 1, 4, 6 and 7). Five patients had the interbody device placed with some minimal bony removal. Two patients had the mesh inserted via a purely percutaneous, bone-sparing approach via Kambin’s triangle (Figures 2 and 3).4 We utilized neuromonitoring for the percutaneous approaches.

Patient 1 was a 66-year-old female who presented with five years of low back pain and one year of right lower extremity pain. An MRI revealed retrolisthesis of L2 on L3, which had progressed slightly over seven years. There were stable findings of anterolisthesis at L4-5 and at L5-S1.

Patient 2 was a 49-year-old male with many years of back pain with one year of paresthesias and numbness in the bilateral lower extremities. An MRI showed...
Patient 3 was a 73-year-old female who had undergone an emergent laminectomy at an outside institution for sudden-onset bilateral lower extremity weakness three months earlier. She regained her strength and sensation but her preoperative, chronic low back pain and left lower extremity pain had worsened after the procedure. An MRI revealed very severe degenerative disc disease at L1-2 with Modic changes. Also noted were postoperative changes at L4 to S1.

Patient 4 was a 66-year-old male with chronic lower back pain which had worsened over six months. He also complained of bilateral lower extremity weakness when going up or down stairs. Physical exam revealed no objective weakness. An MRI revealed a grade I spondylolisthesis at L4-5 with Modic changes and moderate central stenosis.

Patient 5 was a 42-year-old male with a history of end-stage renal disease and had presented to the hospital with severe, intractable low back pain radiating into the left lower extremity. An MRI showed evidence of a discitis at L5-S1 and cultures from a needle biopsy by interventional radiology grew Staphylococcus epidermidis. Despite initial antibiotic treatment and narcotic medications, he remained in severe intractable pain. He did have a history of chronic low back pain.

Patient 6 was a 57-year-old female who had undergone a right hemilaminectomy and bilateral sublaminar decompression at L4-5. She continued to have bilateral lower extremity pain two months after surgery, and dynamic radiographs had demonstrated abnormal motion at L4-5.

Patient 7 was a 46-year-old male who had undergone an L5-S1 instrumented interbody fusion nine years ago. He had developed progressive low back and bilateral lower extremity pain, left greater than right. An MRI showed some mild degenerative disc disease at L4-5 but dynamic radiographs revealed instability at this level with a solid fusion at L5-S1. This particular patient required revision of his existing hardware in order to couple it to the new pedicle screws which were inserted into L4. However, the interbody mesh was delivered via an entirely percutaneous approach.

All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperatively with no complications (nerve root injuries, durotomies, infections, or hardware failures). All patients experienced pain relief compared to preoperative...
the shape of the actual disc space. The PEEK or metal used in most interbody devices also occupy significant space, whereas the mesh allows for more surface area contact for bone to remodel.

Various expandable interbody cages are available, but most devices expand only in the sagittal plane, not in the axial and coronal ones. The OptiMesh expands in all planes and, as mentioned, contours to the shape of the actual disc space. The PEEK or metal used in most interbody devices also occupy significant space, whereas the mesh allows for more surface area contact for bone to remodel.
and potentially increase the likelihood of a robust bony fusion.

Based on our limited experience, the mesh device is a safe, effective option for TLIF. Its main advantages appear to be decreased anatomical disruption during delivery and deployment, the ability to expand in all planes with conformity to the endplates, and greater surface area contact of bone for remodeling and fusion. A study involving a larger number of patients and further long-term follow up is warranted.

REFERENCES