Suturable Mesh Demonstrates Improved Outcomes over Standard Suture in a Porcine Laparotomy Closure Model

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INTRODUCTION

Ventral hernia after routine laparotomy is the most common major complication in general surgery with a 10%-23% occurrence rate. Incisional hernia formation and optimal closing of the laparotomy incision is the focus of much discussion and research. One cause of these failures is “suture pull-through.” Like a wire cutting through ice or cheese, the sharp suture filament either acutely or chronically cuts through abdominal wall tissues, leading to dehiscence in the acute situation and incisional hernia formation when the wound healing failure occurs over time.

A suturable mesh was developed to limit pull-through while approximating tissues. Suturable mesh is made of individual polypropylene filaments that are latticed together in a hollow cylindrical configuration with a macroporous outer mesh wall. Meshes are well established in their ability to maintain tissues in apposition after repair with improved outcomes over standard sutures. A suturable mesh prototype was tested in a rat hernia model. Eighty-five percent of the standard sutures pulled through the abdominal wall, but every suturable mesh prototype maintained its hold on the abdominal wall. A realistic porcine laparotomy model demonstrated a stronger earlier

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repair at 8 days with a suturable mesh prototype than with standard suture. Cadaveric abdominal wall studies with suturable mesh showed that even without fibrovascular incorporation, the suturable mesh showed increased resistance to pull-through in comparison to monofilament suture in an abdominal wall model.

A knowledge gap existed as to the efficacy of a suturable mesh to maintain tissue approximation for longer than 8 days. We hypothesized that a suturable mesh used for laparotomy closure would demonstrate improved macroscopic and microscopic outcomes in comparison to standard suture. To this end, a preclinical study was conducted to compare the outcomes of 0 suturable mesh and number 1 suturable mesh with standard number 1 polypropylene monofilament suture. Primary endpoints were hernia formation at a 13-week sacrifice and histological tissue response. Secondary endpoints included adhesions, surgical site occurrence, documentation of “loose sutures” (ability to distract the suture away from the tissues), adverse events, and change in animal weight.

**MATERIALS AND METHODS**

**Animals**

All animal procedures and housing were performed under protocols approved by the University of Saskatchewan Animal Research Ethics Board. Fifteen nulliparous and non-pregnant female pigs (Sus scrofa domesticus) weighing approximately 72 kg were used for the study (Prairie Swine Centre, Saskatoon, Canada). Fifteen animals were randomly divided among the three groups.

**Implants**

The suturable mesh (Duramesh Suturable Mesh, MSi) is manufactured from 12 polypropylene filaments for the 0 size and 18 filaments for the number 1. The implant is 36 inches long and has an attached swaged standard surgical needle. The 0 suturable mesh has an inner diameter of 2 mm, and the number 1 has an inner diameter of 3.4 mm. The Duramesh device is investigational and not yet allowed for use in the United States. For a comparison suture, a number 1 polypropylene monofilament (Surgipro, Covidien, Minneapolis, Minn.) was chosen as it is an accepted suture for fascial closure (Fig. 1).

**Surgical Procedure**

A ventral midline abdominal skin incision 17 cm in length was made sharply from the xyphoid to the umbilicus. A 2.5 cm wide area of linea alba was cleared of soft tissue with cautery. The linea alba was opened, and the peritoneum incised. Three centimeters of peritoneum and preperitoneal fat were excised from both hemiabdomens.

**Fig. 1.** Number 1 suturable mesh (top), 0 suturable mesh (middle), and number 1 polypropylene suture with attached needles used in protocol.

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**Takeaways**

**Question:** Can a novel tissue approximation device improve on the tendency of sutures to tear through tissues, leading to surgical failure and hernia formation?

**Findings:** Suturable mesh elicits an inflammatory response comparable to that of a standard suture in its quality, yet is larger in size to provide additional support at the suture/tissue interface. This novel device acts as a scar scaffold. Fibrous incorporation of its fine filaments occurs without capsule formation, leading to fewer loose sutures and numerically fewer hernias as compared to a standard suture.

**Meaning:** Suturable mesh is a more reliable tissue approximation device than a standard suture.
for the length of the laparotomy incision. One-centimeter hash marks were drawn with a surgical marking pen to ensure 1-cm bites and 1-cm travels. The linea alba was randomly closed with either suturable mesh or polypropylene monofilament in a simple continuous (running) suture pattern. A knot was created at each of the two opposing ends of the incision, and the sutures were run toward each other then tied to each other in a third knot (Fig. 2). A video of the implantation is available here: https://youtu.be/glCxHgDg0PK. The subcutaneous tissues were closed using interrupted and running 2-0 polydiaxanone sutures.

Animal Care

Each pig was weighed before surgery and then weekly, before necropsy. Incision observations were recorded daily. Physical examination of the pigs was performed at 1 month, 2 months, and before necropsy.

Table 1. Scoring System for Visceral Adhesions at Necropsy

| Adhesion Scoring, Extent | Score | Percent of material area involved with adhesions |
|--------------------------|-------|-----------------------------------------------|
| 0                        | No adhesions |
| 1                        | 1% to 25% |
| 2                        | 26% to 50% |
| 3                        | 51% to 75% |
| 4                        | Greater than 75% |

| Adhesion Scoring, Severity | Score | Percent of material area involved with adhesions |
|-----------------------------|-------|-----------------------------------------------|
| 0                           | No resistance to separation |
| 1                           | Mild resistance to separation |
| 2                           | Some resistance (moderate force is required) |
| 3                           | Sharp dissection needed |

| Adhesion Scoring, Degree/Intensity | Grade | Description |
|-----------------------------------|-------|-------------|
| 0                                 | No adhesions present |
| 0.5                               | Thin, translucent adhesions composed of few fibrinous strands |
| 1                                 | Continuous fibrous, avascular adhesions; disrupted by gentle blunt dissection |
| 2                                 | Fibrous adhesions, some vascularity; disruption of adhesions requires sharp dissection; identifiable tissue planes |
| 3                                 | Dense scar with obliteration of tissue planes |

Necropsy

At necropsy, the suture line for each animal was graded for adhesion formation between the peritoneal side of the suture line and adjacent abdominal organs using the scoring table (Table 1). The abdominal wall was then examined carefully for evidence of loose sutures, suture pull-through, and abdominal wall defects scored as per Table 2.

Pathology Evaluation and Analysis

The sutured incision sites were excised and the excised blocks of tissues were trimmed to allow a sufficient area around the implant for proper histological preparation. Three cross sections were obtained from each sutured incision site and identified as cranial, middle and caudal. A section was then obtained to the left of the incision line from the cranial end, a section to the right of the incision line from the middle portion, and a section to the left of the incision line from the caudal end, each at approximately 1–3 mm away from the incision line (Fig. 3). These sections were embedded in methyl methacrylate (MMA) and subsequently polymerized. Thin, approximately 8 µm, sections from each block were stained with hematoxylin and eosin (H&E). A semiquantitative descriptive microscopic evaluation of histology sections (sutured incision sites) was performed by a certified veterinary pathologist (AccelLAB, Montreal, Canada). The sections were analyzed and graded according to cell type and tissue response. The quality of the tissue response to the implant was evaluated following the irritancy/reactivity grading scheme adapted from the ISO 10993-6 Annex E. Assessment of the size of the soft-tissue response surrounding the three implants using measurements taken using an optical micrometer (Fig. 4) was made to illustrate differences in histological response. A semiquantitative microscopic evaluation of tissue ingrowth was performed as per Melman. As the suturable mesh and the comparison standard suture have different appearances under the microscope, the pathologist could not be blinded as to implant type.

Statistical Analysis

No statistics were used to assess the primary endpoint of hernia formation. For histology, equal variance and
Fig. 3. Diagram of tissue blocks taken for histologic analysis of midline closures.

Fig. 4. Grade of tissue response quantitated histologically.
normality tests were performed. When both were successful, one-way analysis of variance (ANOVA) means tables with Dunnett tests were used to calculate the significance of differences between continuous variables of the study implant groups. When either equal variance test or normality test failed, a Kruskal–Wallis one-way ANOVA on ranks with Dunn’s method was conducted. A P value of less than 0.05 was considered statistically significant.

RESULTS

Hernia Formation
For the primary endpoints, there were numerically fewer hernias with the number 1 suturable mesh than the 0 suturable mesh or the number 1 polypropylene sutures. All of the animals with intact abdominal walls were in the suturable mesh groups (Figs. 5 and 6). All of the polypropylene monofilament closures demonstrated small abdominal wall defects (Fig. 7).

Histology
Neovascularization was significantly higher for 0 suturable mesh and number 1 suturable mesh as compared to number 1 polypropylene (P = 0.005). The 0 suturable mesh had a score of 2.53 ± 0.51 for tissue ingrowth on a three-point scale. The number 1 suturable mesh had a score of 2.33 ± 0.63 for tissue ingrowth on the same scale. The polypropylene suture is solid standard suture and, therefore, did not permit any tissue ingrowth.

Both sizes of suturable mesh at 3 months were regarded as nonirritants to the tissues in comparison to polypropylene monofilament using the semi-quantitative histiologic scoring system. Capsule formation was not noted in any of the samples, and therefore capsular thickness was not measured. The dimensions of the soft-tissue response were graded as 3.93 ± 0.15 for number 1 suturable mesh, 3.40 ± 0.15 for the 0 suturable mesh, and 2.13 ± 0.99 for number 1 polypropylene monofilament (Fig. 8). (See figure, Supplemental Digital Content 1, which displays histologic cross-section of number 1 polypropylene suture (top), 0 suturable mesh (middle) and number 1 suturable mesh (bottom) after staining with hematoxylen and eosin (left), and Masson’s trichrome (right). Implantation of all devices is associated with mild peridevice inflammatory response characterized by a presence of macrophages and multinucleated giant cells as well as lymphocytes, plasma cells, and polymorphonuclear cells. The number 1 polypropylene is characterized by a region of fibroplasia surrounding the suture monofilament and neovascularization. Suturable meshes are characterized by presence of fibroplasia (organizing fibrous connective tissue) and neovascularization surrounding, and occasionally intercalating between strands of the multifilament suture material (C–F), http://links.lww.com/PRSGO/B811.) There was a statistical difference between the groups (P < 0.0001).

Secondary Endpoints
The pigs gained weight equally during their 13-week observation and were approximately 150 kg at necropsy. All incisions remained well healed with incidental skin lesions found equally in all three groups. Surgical site occurrence was noted in two animals with seromas that resolved spontaneously (one seroma in suturable mesh group, one in polypropylene group). Nine of the 10 suturable mesh
devices were well-incorporated within the tissues and could not be pulled away (Fig. 9), whereas four of the five polypropylene sutures were loose (Fig. 7). Visceral adhesions to the suture line were observed in two pigs closed with 0 suturable mesh and two pigs closed with number 1 polypropylene. Use of the preoperatively defined grading scale for adhesions showed two number 1 polypropylene monofilament suture animals to have a score of 1 for severity and degree, and scores of 1 and 2 for extent. The 0 suturable mesh animals had similar adhesion scores in two animals as the number 1 polypropylene group. There were no adhesions noted to the closure line with number 1 suturable mesh. There were 21 adverse events (two soft-tissue swellings thought to be seromas that resolved, lameness, skin lesions found to be unremarkable by pathology, and one febrile illness with weight loss), and none were thought related to the implant used for closure.

**DISCUSSION**

The attributes of sutures and their intrinsic failure patterns have been accepted by the surgical community and have not been questioned since the dawn of modern surgery. We believe, as others do others, that suture pull-through due to the action of an implant under tension on the encircled tissue is the primary cause of failed tissue approximations.10,11 Suturable mesh is a tissue approximation device that increases the implant surface area contacting the tissues in comparison to a standard suture. With fibrovascular ingrowth, the individual filaments act as a scar scaffold and become microencapsulated over time. Although the surface area of number 1 suturable mesh is over 500% times greater than the surface area of a number 1 monofilament, the total volume of foreign material is only 35% increased. The total volumes must be similar for both implant types to fit into the end of a standard swaged surgical needle.

Number 1 suturable mesh had three of the five closures with completely intact abdominal walls. One of the five closures with 0 suturable mesh was completely intact. None of the polypropylene closures were perfectly closed.
A “small-bites” closure technique may have demonstrated improved hernia outcomes. A “large-bites” closure with 1-cm bites and 1-cm travels was performed to put more stress at the suture/tissue interface and perhaps demonstrate differences between the closure devices. The data point to the suturable mesh having superior outcomes for tissue approximation, but also to be noted that this is a highly stressed model with human-sized pigs that grew to over 300 lbs during the study. No maneuvers were taken to limit the animals’ activity after the procedure to lessen the stress on the repairs. The animals are quadripped ambulators with all of their torso weight on their closures. Although only four of 15 closures were completely intact, it is almost surprising that any of these closures survived until necropsy.

Polypropylene was chosen for the implant material based on its nonreactivity to tissues and its long-standing use in surgery. The foreign body reaction and microencapsulation will produce a permanent support to the suture line. This may be important in abdominal wall surgery, where scar without a permanent implant is only 70% as strong as uninjured tissue. Some surgeons may prefer an absorbable product, and this is in development.

The optically measured size of the soft-tissue response was larger with number 1 suturable mesh than for number 1 polypropylene, though the intensity or quality of the tissue response measured using the ISO 10993-6 Annex E grading scale was judged to be equivalent. The latter makes sense, as both implants are manufactured from polypropylene. This data illustrate the mechanism by which small filament meshes provide support to tissues and to newly opposed suture lines.

Four of the five polypropylene monofilament running sutures were loose at necropsy. Loose permanent sutures are the hallmark of pull-through. As the animals doubled in size during the study, the only means for sutures to become loose is for the tissue encircled within the suture loop to become narrower in diameter. The one suturable mesh recorded as being loose was at the site of a small hernia defect. For this animal, the remainder of the laparotomy closure line demonstrated a well-incorporated implant.

The histological outcomes demonstrate a controlled inflammatory tissue reaction. Suturable mesh was graded as a “nonirritant” to the tissues in comparison with monofilament polypropylene, but there is a multiplication of this “normal” tissue response due to the multiple filaments of the suturable mesh implant. The soft-tissue response may elucidate the mechanism by which meshes reduce pull-through in current methods of abdominal wall tissue approximation. Adhesions were least in the number 1 suturable mesh group and greatest in the other two groups. Although each group had essentially no foreign material exposed to the viscera, perhaps it is the inflammation of slow tearing from suture pull-through that causes the bowel to adhere to a newly closed suture line. The animal group with the best “hold” and least pull-through (number 1 suturable mesh) therefore also had the least adhesions, but this is only a theory.

Suturable mesh demonstrated an ability to approximate tissues under tension in this realistic porcine laparotomy closure model in human-sized pigs. This preclinical study is an extension of prior work that showed a suturable mesh porcine laparotomy repair required double the force to fail as compared to polypropylene monofilament suture. Further studies are underway to better define the utility of suturable mesh for internal closures.

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