A Prospective Randomized Trial of Biologic Mesh versus Synthetic Mesh for the Repair of Complex Ventral Hernias.

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SPECIFIC AIMS

Ventral hernia (VH) is a frequent sequela of abdominal surgery, occurring after up to 10-20% of laparotomy incisions.[1, 2] This results in almost 250,000 annual ventral hernia repairs in the United States, making it one of the most common procedures performed by general surgeons.[3] The annual health care expenditures associated with hernia repairs now exceeds $5 billion US dollars. [4] Since prosthetic mesh has been shown to reduce recurrence rates by over 50%, most surgeons agree that some form of prosthetic reinforcement should be added to all but the smallest ventral hernia repairs.[5] Traditionally, this has involved a permanent prosthetic material in clean cases (without bacterial contamination) at relatively low cost (approximately $150 for a 900cm$^2$ mesh). The management of more complex hernias where infection or contamination is present is not well defined. These patients are often severely disabled by the chronic infectious nidus and suffer a very poor quality of life until reconstruction of their abdominal wall anatomy and resolution of the infection. Historically, this has involved a two-stage approach for hernias where simultaneous gastrointestinal, biliary, and/or genitourinary procedures were performed or if there was an active infection of a prosthetic material. The first stage involved removing the infectious or contaminated source, and performing a temporary closure of the abdominal wall with an absorbable material. This approach of so-called “planned hernia” would almost uniformly require another major operation.
Approximately 6 months to one year later, the abdominal wall is repaired with a permanent synthetic material in a clean setting. This approach avoids placing the synthetic mesh in the field of contamination, but is associated with significant morbidity as it requires two operations and potential long-term disability during the recovery period. Recognizing the limitations of this two-stage approach, new biologic materials have been designed to offer a single-stage approach for infected and contaminated abdominal wall reconstruction. These materials are derived from various sources including human or porcine dermis, bovine pericardium, or intestinal submucosa. They are processed in such a way as to render an acellular collagen-rich graft that reportedly acts as a cellular scaffold to allow native tissue in-growth and regeneration of tissue. They are marketed as resistant to infection and therefore as a suitable hernia repair material for clean-contaminated and contaminated abdominal wall defects. However, these products are very expensive with a 400cm$^2$ porcine dermal graft costing over $10,000, and currently represent the most rapidly growing market in hernia repair, with estimates of almost $500 million dollars in annual revenue by the year 2013.[6] Despite the rapid acceptance of these materials, there is little preclinical or clinical evidence to support their claims of regeneration, or that they provide a durable repair to the abdominal wall in the setting of a clean-contaminated (Class 2) or a contaminated (Class 3) surgical procedure.

In our experience, the single-stage repair of contaminated abdominal wall defects with these biologic grafts has resulted in a 40%-80% recurrence rate with long term follow up.[7, 8] Given these disappointing results, other investigators
have evaluated the role of synthetic mesh in the repair of contaminated abdominal wall defects in small, retrospective non-randomized trials.[9-11] Importantly, several recent modifications to polypropylene mesh have yielded potential options for repairing contaminated defects. These modifications include reducing mesh weight, increasing mesh porosity, and utilizing monofilament unwoven material that resists bacterial colonization in animal studies.[12, 13] In fact, our lab has recently evaluated 9 commercially-available prosthetic materials and has found that certain meshes can clear bacterial contamination similar to biologic grafts in a rat hernia model. These materials have been used for years in clean repairs of large complex abdominal wall defects with significantly lower recurrence rates (<5%) than a biologic mesh.[14] Beyond doubt, a non-biased prospective randomized trial is long overdue to compare the safety and efficacy of a biologic graft versus a macroporous light-weight polypropylene synthetic material for the repair of complex ventral hernias. This study will compare the safety, efficacy, and cost-effectiveness of a permanent synthetic mesh versus a biologic prosthesis for the open repair of ventral hernias in the setting of clean-contaminated (Class 2) or contaminated (Class 3) surgical procedures.

This study will have a major impact on the field of hernia surgery, as the study findings will provide an objective guide to mesh selection, optimize surgical approaches for complex ventral hernia repair, and ultimately significantly improve patient outcomes. The lack of data as to the ideal mesh selection for complex ventral hernia repair has resulted in physicians relying on anecdotal experience, industry marketing, and personal bias. Presently, hundreds of thousands of
patients are affected by this condition and would significantly benefit from clear practice guidelines regarding the best approach and the most appropriate prosthetic selection for repairing these complex ventral hernias. In order to address this important need, the overall safety, efficacy, and cost-effectiveness of a biologic prosthetic as compared to a synthetic material for the open repair of complex defects should be subjected to a prospective randomized clinical trial.

We hypothesize that reinforcement of single-stage open repairs of complex abdominal wall defects with a macroporous light-weight polypropylene synthetic mesh will result in significantly lower rates of hernia recurrence (HR) and surgical site occurrences requiring procedural intervention (SSOPI) at 24 postoperative months, and greater cost-effectiveness compared to reinforcement with a biologic mesh. We further hypothesize that reinforcement with macroporous light-weight polypropylene synthetic mesh will be associated with a significantly greater change in preoperative to postoperative patient-reported quality of life (QOL) compared to reinforcement with biologic mesh for clean-contaminated or contaminated ventral hernias.

The specific aims of the proposal are:

1. To demonstrate that a single-stage repair of clean-contaminated (Class 2) or contaminated (Class 3) ventral hernias using a macroporous light-weight polypropylene synthetic mesh will result in superior clinical outcomes compared to a biologic mesh.
   a. Task 1- Demonstrate that repairs of clean-contaminated and contaminated ventral hernias performed with macroporous light-
weight polypropylene mesh will result in fewer recurrent hernias and fewer surgical site occurrences requiring procedural intervention at 24 postoperative months compared to repairs of clean-contaminated and contaminated ventral hernias performed with biologic mesh.

b. Task 2- Compare postoperative pain, and demonstrate greater change in preoperative to postoperative quality of life (QOL) at 1 month, 6 months, 12 months, and 24 months following clean-contaminated or contaminated ventral hernia repair with a macroporous light-weight polypropylene mesh versus a biologic prosthesis.

2. To demonstrate that a macroporous light-weight polypropylene mesh is the more cost-effective strategy than a biologic prosthetic in clean-contaminated and contaminated abdominal wall reconstruction.

   a. Task 1-Estimate direct and indirect economic costs associated with clean-contaminated or contaminated ventral hernia repair using either polypropylene mesh or biologic mesh from a limited societal perspective.

   b. Task 2 – Perform health utility valuation in patients undergoing repair of clean-contaminated or contaminated ventral hernias using either polypropylene mesh or biologic mesh.

   c. Task 3 – Calculate and compare incremental cost-effectiveness ratios for patients undergoing repair of clean-contaminated or
contaminated ventral hernias using polypropylene mesh versus biologic mesh.

Research Strategy

SIGNIFICANCE

The repair of ventral hernias in the presence of active infection or contamination is an extremely challenging problem for the reconstructive surgeon. Very little data exists as to the ideal surgical approach or the appropriate class of reconstructive material to affect a long-term durable repair without promoting chronic mesh contamination or infection. This lack of scientific evaluation of this very common clinical scenario has resulted in the development of one of the fastest growing markets in abdominal wall reconstruction: biologic mesh. Healthcare expenditures associated with hernia repair have topped $5 billion dollars, and a substantial and growing amount can be attributed to biologic prosthetics.

Based on well-designed prospective randomized trials, typical hernia repairs in a “clean” (Class 1 wound) field are almost always performed with a synthetic mesh, as long term results have demonstrated reduction in recurrence rates of up to 50%. [5, 15] Despite almost no strong clinical evidence, it has generally been accepted by the US surgical community that synthetic mesh should be avoided in the presence of contamination.[10, 12, 16-19] The exact basis of these concerns is not entirely clear. Based on historic data and evaluating microporous or heavy weight materials, several authors have reported
high rates of mesh sepsis requiring explanation.[16] This has created an overwhelming fear amongst surgeons that synthetic mesh will almost always become infected and will always require surgical excision. However, a closer inspection of the literature would suggest that some synthetic materials are in fact quite resistant to bacterial contamination. Furthermore, both in vitro and in vivo analyses have demonstrated that various prosthetic meshes can be salvaged if exposed to bacterial contamination.[11-13, 19-22]

The fear of placing a synthetic mesh in a clean-contaminated or contaminated field has resulted in two common approaches to this clinical problem. Historically, the standard of care for repairing these hernias occurring in compromised fields has involved two staged procedures. Initially, the contaminated portion of the procedure (takedown of fistula, bowel resection, removal of infected prosthetic, or clearance of abdominal wall infection) is performed and the abdominal wall is temporarily closed with sutures or an absorbable mesh. After 6 to 12 months and complete healing of the wound, the patient undergoes a definitive abdominal wall reconstruction, often with synthetic mesh. While this approach has resulted in acceptable long-term outcomes, the patients have a prolonged period of convalescence, extended hospitalizations, restricted activity, and poor quality of life. Recognizing these limitations, the potential alternative of a definitive single-stage reconstruction for clean-contaminated and contaminated abdominal wall defects is particularly appealing. [23] These single-stage repairs have routinely been performed with biologic prosthetics with variable results. If biologic mesh could result in healed wounds
and durable single-stage repairs with low recurrence rates, it could reduce patient recovery times and improve return to a normal functional status while eliminating a second procedure. Since these repairs are often occurring in clean-contaminated, contaminated, and dirty fields, the reported incidence of surgical site infections are as high as 8%, 15%, and 27%, respectively.[24]

Over the past decade, a multitude of biologic prosthetics have been released to improve the results of single-stage repairs for these challenging cases. These materials are derived from various sources including human or porcine dermis, bovine pericardium, or intestinal submucosa. They are processed to render an acellular collagen-rich graft designed to act as a cellular scaffold to allow native tissue in-growth and regeneration of tissue. They are marketed as resistant to infection and therefore as a suitable alternative to repairing clean-contaminated and contaminated abdominal wall defects. [8, 25-28] However, these products are very expensive, with a 400cm² porcine dermal graft costing over $10,000 dollars, and currently represent the most rapidly growing market in hernia repair, with estimates of almost $500 million dollars in annual revenue by the year 2013.[6] Despite the rapid acceptance of these materials, there is little preclinical or clinical evidence to support their claims of regeneration, or that they provide a durable repair to the abdominal wall in the setting of clean-contaminated (Class 2) or contaminated (Class 3) surgical procedures. In our experience, the single-stage repair of contaminated abdominal wall defects with these biologic grafts has resulted in a 40%-80% recurrence rate with long-term follow-up.[7] Given these disappointing results,
other investigators have evaluated the role of synthetic mesh in the repair of clean-contaminated and contaminated abdominal wall defects in small, retrospective non-randomized trials.[9-11] Importantly, several recent modifications to polypropylene mesh have yielded potential options for repairing contaminated defects. These modifications include reducing mesh weight, increasing mesh porosity, and using monofilament unwoven materials that resist bacterial colonization in animal studies.[12, 13] In fact, our lab has recently evaluated 9 commercially-available prosthetic materials and has found that certain meshes can clear bacterial contamination similar to biologic grafts in a rodent chronic infection model.[29] (Figure 1) These synthetic materials have been used for years in repairing large complex abdominal wall defects in clean settings with significantly lower recurrence rates (<5%) than biologic meshes.[14] A non-biased prospective randomized trial is long overdue to compare the safety and efficacy of a biologic graft versus a macroporous light weight polypropylene synthetic material for the open repair of complex ventral hernias to appropriately guide mesh selection and optimize patient outcomes. This study will compare the safety, efficacy, and cost-effectiveness of a permanent synthetic mesh versus a biologic prosthesis for the open repair of ventral hernias in the setting of clean-contaminated (Class 2) or contaminated (Class 3) surgical procedures.

APPROACH

We hypothesize that reinforcement of single-stage open repairs of complex abdominal wall defects with a macroporous light-weight polypropylene
synthetic mesh will result in significantly lower rates of hernia recurrence (HR) and surgical site occurrences requiring procedural intervention (SSOPI) at 24 postoperative months, and greater cost-effectiveness compared to reinforcement with a biologic mesh. We further hypothesize that reinforcement with macroporous light-weight polypropylene synthetic mesh will be associated with a significantly greater change in preoperative to postoperative patient-reported quality of life (QOL) compared to reinforcement with biologic mesh for clean-contaminated or contaminated ventral hernias.

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the more cost-effective strategy than a biologic prosthetic in clean-
contaminated and contaminated abdominal wall reconstruction.**
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      clean-contaminated or contaminated ventral hernia repair using
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**STUDY DESIGN**

We propose a multi-center prospective double-blinded randomized
controlled trial comparing 253 patients with clean-contaminated (Class 2) or
contaminated (Class 3) abdominal wall ventral hernias undergoing single-stage
open repair. Soft Mesh™ by CR Bard™, a macroporous monofilament
polypropylene permanent mesh, will be compared to Strattice™ mesh by LifeCell™, a non-crosslinked porcine dermal biologic graft, for the single-stage open reconstruction of clean-contaminated and contaminated abdominal wall defects. The primary outcome variables will be the absence of surgical site occurrence requiring procedural intervention and the absence of a hernia recurrence from the time of surgery up to 24 months of postoperative follow-up.

**Study Procedures**

Patients undergoing open ventral hernia repair for clean-contaminated and contaminated abdominal wall hernias meeting inclusion criteria will be randomized to receive a synthetic mesh or a biologic mesh. Randomization will be carried out using computer-generated randomization blocks at the time of enrollment. Stratified randomized will be used with the strata formulated by medical center then by clean-contaminated or contaminated surgical site class. The Investigator will be blinded to patient randomization assignment until the point of intra-operative device use following final CDC wound classification, whereas patients and co-investigators responsible for data analysis will remain blinded to patient randomization until the conclusion of the study period. As such, a double-blinded study protocol will be maintained. Patients randomized to synthetic mesh will receive Soft Mesh™ (CR Bard™, Murray Hill, NJ), and those patients randomized to biologic mesh will receive Strattice™ (LifeCell™, Branchburg NJ). The use of biologic and synthetic meshes in clean-contaminated and contaminated fields is considered experimental; however, the
selection of these prosthetics was based on a careful review of the multiple animal models, preclinical data, and our own clinical experience with each of these materials placed in both clean and contaminated abdominal wall reconstructions.[26, 30-35]

Soft Mesh™ is a light-weight (44 g/m²) monofilament macroporous polypropylene synthetic mesh. It does not have an anti-adhesive barrier and must be placed in an extraperitoneal position. Our lab has evaluated this material in a chronic rat infection model and has shown clearance rates of bacterial contamination comparable to biologic grafts (Figure 1). This prosthetic has become our material of choice for routine abdominal wall reconstruction and ventral hernia repairs, given its chemical and structural properties. In addition, we have also utilized Soft Mesh™ in the retrorectus position in several cases of elective bowel surgery, parastomal hernia repair, and inadvertent enterotomies with excellent results including no mesh infections and no long-term hernia recurrences. Other authors have reported excellent outcomes with polypropylene-based synthetic material placed in the extraperitoneal position.[14] Interestingly, reports of placing several forms of polypropylene-based meshes with anti-adhesive barrier coatings into the peritoneal cavity for elective hernia repair with concomitant bowel surgery has shown very high rates of mesh sepsis and subsequent mesh excision.[36] Mesh placed in the intraperitoneal position comes in contact with the viscera and therefore requires some form of an anti-adhesive barrier. We noted that these anti-adhesive barriers prevent bacterial clearance in an experimental study (Figure 1). Based on our findings, it is likely
that both the specific type of synthetic mesh and the compartment of the abdominal wall in which it is placed will have significant effects on bacterial clearance and success when placed in a clean-contaminated or contaminated field. Potential extraperitoneal compartments for mesh deployment include the onlay (placed above the fascia in the subcutaneous space) or a retro-rectus (below the muscles but above the peritoneum) position. The onlay position can result in early mesh exposure if wound infection or breakdown occurs and has been shown in other prospective randomized trials to result in a high rate of mesh infections.[37] Alternatively, the retro-rectus repair with synthetic mesh reinforcement has been demonstrated by multiple authors as a durable repair with very low rates of mesh infection and hernia recurrence.[14, 38] For these reasons, we feel it is particularly important to design this experiment to use an unprotected macroporous light-weight polypropylene mesh placed in the retro-rectus position.

The Strattice™ biologic mesh is derived from porcine dermis and processed to remove the cells but maximally preserve the dermal matrix. The processing avoids the use of collagen cross linking agents in a reported effort to minimize immunogenic response, improve biocompatibility, and ultimately promote rapid revascularization and tissue regeneration.[39] Multiple biologic grafts have been developed to repair contaminated abdominal wall defects, however little comparative data exists to definitively guide selection of these grafts. Our lab has performed several preclinical evaluations of these materials and has chosen Strattice™ based on our findings. Strattice™ mesh showed
excellent biocompatibility from an immunologic perspective when compared to other human and porcine derived biologic products.[40] This selection should limit the potential for immunologic responses to the biologic material confounding our results. We also evaluated the ability of various biologic grafts to clear bacterial contamination in a chronic infection rodent model.[29] Strattice™ mesh had the highest rates of bacterial clearance when compared to other porcine derived materials. Interestingly, it appears that based on our findings, Soft Mesh™ and Strattice™ result in similar rates of bacterial clearance, which is the primary driving force of our renewed interest in evaluating the usage of these inexpensive synthetic meshes in clean-contaminated and contaminated fields.

Patients will be evaluated at initial preoperative visit for meeting inclusion and exclusion criteria. Patients will be included in this study if they are 21 years of age or older (including women of childbearing age), undergoing a planned single-stage open reconstruction of a contaminated (CDC wound class 2 or 3) abdominal wall defect (including concomitant procedures: creation of a stoma, bowel resection, panniculectomy, removing uninfected mesh, and gastrointestinal, genitourinary, or gynecologic procedure) under general anesthesia, can achieve midline fascial closure, and have a parastomal hernia or midline defect at least 9 cm². Patients will be excluded from the study if they meet any of the following criteria: are undergoing a laparoscopic or robotic repair of the abdominal wall defect, have a CDC class 1 or 4 wound (see CDC guidelines below), have a defect that the surgeon cannot achieve primary fascial apposition and requires a bridge of mesh, body mass index (BMI) >45 kg/m²,
chronic immunosuppression including medically-induced with >10 mg of prednisone/day, collagen vascular disorder, severe malnourishment (albumin <2.0 g/dl), ascites refractory to medical management, end stage renal disease (indwelling hemodialysis or peritoneal dialysis), pre-existing liver disease (hepatitis B or C or total bilirubin >3.0 mg/dl), smoking history within 1 month of surgery, current pregnancy, require removal of a prior surgical mesh during a planned ventral hernia repair due to active mesh infection (as defined by a synthetic mesh that is not incorporated into the tissue, is extracorporeally exposed, or has a chronic draining sinus with clear fluid around the material; but not including synthetic mesh that is incorporated into the abdominal wall and not infected), are unable to undergo successful retro-rectus preperitoneal mesh placement, if they object to the implantation of porcine products, or if they are participating in other clinical trials.

Upon enrollment into the study, patients will be randomized to either synthetic or biologic mesh for their repair. All patients will undergo our standard pre-operative evaluation. Briefly, it will include a complete set of laboratory studies including complete blood count, comprehensive metabolic panel, albumin, prealbumin, HbA1C (for diabetic patients), and urinalysis. Pregnancy test will be performed for those patients of child bearing potential. Photos may be taken of the anterior abdominal wall. An abdominal pelvic CT scan will be obtained preoperatively in all patients based on our standard approach, (abdominal pelvic CT scan within the past twelve months is sufficient) and any issues postoperatively (including suspected recurrence) will be evaluated with a
CT scan as clinically indicated. Preoperative demographics and clinical data including sex, race, age, body mass index (BMI), location of the hernia, length and width of the hernia defect, wound classification (per CDC guidelines, table 1), smoking status (active within 3 month of surgery), medical history, surgical history of prior abdominal surgical procedures and prior ventral/incisional hernia repairs will be documented. Intraoperative details will include patient ASA score, patient temperature, use of epidural catheters, size of fascial defect (measured as maximal width and length), fascial layers released (external oblique, posterior rectus sheath, or transversus abdominis muscle), adhesions, concomitant procedures, wound characterization, mesh type, mesh placement, operative time, estimated blood loss, blood transfusion requirement, perioperative antibiotic administration (including type, dose, frequency, and times of initiation and discontinuation), and intraoperative fluid administration. Postoperatively, patients will be evaluated for signs and symptoms of complications along with presence or absence of surgical site infections (SSIs) per CDC definitions as categorized below (Appendix 4), presence or absence of surgical site occurrences (SSOs) and any procedural interventions required to treat these SSOs, presence or absence of hernia recurrence and any reoperations, length of hospital stay, discharge date, time to return of bowel function and any readmission. Wound erythema treated with antibiotics will be considered a wound cellulitis. Wounds that are opened and cultured will be appropriately categorized as postoperative SSIs based on CDC definitions. Type, dose, frequency, and duration of antibiotics will be recorded. Patients will also fill out HerQLes (Appendix 1) and
EQ-5D (Appendix 2) quality of life tools preoperatively and during each postoperative visit, at 4 weeks (+ 2 weeks), 6 months (+2 months), 12 months (+ 3 months), and 24 months (+ 4 months). See Appendix 7 for Plan for Continuation of Study Follow-up Visits for Patients Previously Enrolled at University Hospital.

There is substantial evidence that the majority of hernia recurrences occur within the first 24 months after repair.[5, 15, 41]

Preoperative antibiotic usage will be standardized (as per SCIP protocol) as follows. All patients will receive a second generation cephalosporin within 60 minutes prior to the surgical incisions. Patients with a prior history of penicillin allergy or MRSA wound/mesh infections will instead receive a preoperative dose of intravenous vancomycin. The exact drug, along with the dose, frequency, and time administered will be recorded. All antibiotics will be discontinued after 24 hours of surgery unless otherwise indicated. Prolonged antibiotic usage will be clearly documented as to indication, type, dose, frequency, and duration.

The surgical approach to repairing these defects will be standardized, as previously described. All patients will receive a chlorhexidine skin preparation, an iodine-impregnated skin barrier, and all stoma sites will be over sewn at the muco-cutaneous junction to limit bacterial contamination prior to skin preparation. If patients have an allergy to chlorhexidine, then Duraprep™ will be utilized. For any patient who has an allergy to iodine, then a Steri-Drape may be used as a skin barrier. Hair will be removed at the time of surgery with electric clippers.

The midline fascia will be opened and complete adhesiolysis performed to free up the entire abdominal wall. All concomitant procedures will be performed prior
to beginning the abdominal wall reconstructive phase, and documented.

Intraoperative concomitant procedures will be allowed unless they change the wound classification to a class 4. Acceptable concomitant procedures include: the creation of a stoma, bowel resection, gastrointestinal surgery, genitourinary surgery, gynecological surgery, panniculectomy, and removing uninfected mesh.

The abdominal wall is reconstructed by initially incising the posterior rectus sheath just lateral to the linea alba. The release is performed at least 5 centimeters above and below the fascial defect. The posterior rectus sheath is then separated off the rectus muscle to the linea semilunaris. If additional release is necessary to achieve fascial closure, the transversus abdominis muscle or the external oblique muscle may be released at the discretion of the surgeon and documented. The posterior components are then reapproximated to exclude the abdominal viscera from the mesh. If the mesh cannot be placed in the retro-rectus or preperitoneal position, then the patient will be excluded from the study. Unless contraindicated due to drug allergies, a pulse lavage antibiotic irrigation using a 3 liter bag with Gentamycin (240 mg), Ancef (3gm), and Bacitracin (50,000 units) will be applied to the posterior rectus sheath and subcutaneous tissues after the components are reapproximated and prior to mesh placement. If there is a drug allergy, then pulse lavage irrigation with sterile saline only will be applied. Final wound classification will occur just prior to mesh placement per CDC criteria. Surgical wounds will be classified based on CDC criteria and only Class 2 and 3 wounds will be included in this study (see Appendix 3):
Class 2: Clean-Contaminated

Operative wounds in which the respiratory, alimentary, genital*, or urinary tract are entered under controlled conditions, and without unusual contamination. Specifically, operations involving the biliary tract, appendix, vagina, and oropharynx are included in this category, provided no evidence of infection or major break in technique is encountered.

*Includes female and male reproductive tracts

Class 3- Contaminated

A surgical field with any of the following: open, fresh, accidental wounds; operations with major breaks in sterile technique or gross spillage from the gastrointestinal tract; and incisions in which acute, nonpurulent inflammation is encountered.

Following wound classification, the allocation of the patient to either the biologic mesh cohort or the permanent synthetic mesh cohort will be revealed to the operating surgeon, according to the previously-described computer-generated block randomization scheme stratified by wound classification and medical center. The corresponding prosthetic material will then be placed with at least 5
cm of fascial coverage on all sides of the defect. The mesh will be fixated with trans-abdominal #1 Maxon or PDS sutures at 5-10 cm intervals. Number of sutures to secure the mesh will be documented. All patients will have closed suction drains placed above the mesh and below the fascial closure. These drains will be removed postoperatively when the collected fluid is < 30 cc/day for 48 hours and documented. The fascia will be closed with a running or interrupted Maxon or PDS #1 suture. The skin will be closed loosely with staples. Dry sterile dressings will be placed at the conclusion of the procedure and will be removed on postoperative day 2. No further dressings will be applied to the wound.

Postoperative complications will be defined and recorded on the CRF's based on CDC standardized terms and definitions for Surgical Site Infections as follow: [42, 43]-See appendix 4.

**Superficial Incisional Surgical Site infection (SSI)**

A **superficial incisional SSI** must meet the following criteria:

Infection occurs within 30 days after the operative procedure AND involves only skin and subcutaneous tissue of the incision AND patient has at least ONE of the following:

a. purulent drainage from the superficial incision.

b. organisms isolated from an aseptically-obtained culture of fluid or tissue from the superficial incision.
c. superficial incision that is deliberately opened by a surgeon and is culture-positive or not cultured AND the patient has at least one of the following signs or symptoms: pain or tenderness, localized swelling, redness, or heat. A culture-negative finding does not meet this criterion.

d. diagnosis of superficial incisional SSI by the surgeon or attending physician.

NOTE:

a. Do NOT report stitch abscess (minimal inflammation and discharge confined to suture penetration site) as an infection.

b. Do NOT report a localized stab wound or pin site infection. Instead, report these as skin or soft tissue infections, depending on their depth.

c. “Cellulitis” by itself does NOT meet criteria for superficial incisional SSI

d. If infection involves or extends into the fascial and muscle layers report as a deep incisional SSI.

Deep Incisional SSI

A deep incisional SSI must meet the following criteria:

Infection occurs within 30 or 90 days after the operative procedure AND the infection involves deep soft tissues (e.g., fascial and muscle layers) of the incision AND patient has at least ONE of the following:
a. purulent drainage from the deep incision but not from the organ/space component of the surgical site

b. a deep incision spontaneously dehisces or is deliberately opened by a surgeon AND is culture-positive or not cultured AND the patient has at least one of the following signs or symptoms: fever (>38°C), or localized pain, or tenderness. A culture-negative finding does not meet this criterion.

c. an abscess or other evidence of infection involving the deep incision is found on direct examination, during invasive procedure, or by histopathologic examination or imaging test.

d. diagnosis of a deep incisional SSI by a surgeon or attending physician.

NOTE:

a. Classify an infection that involves both superficial and deep incision sites as a deep incisional SSI.

b. Classify infection that involves superficial incisional, deep incisional, and organ/space sites as deep incisional SSI. This is considered a complication of the incision.

Organ/Space SSI

An organ/space SSI must meet the following criteria:

Infection occurs within 30 or 90 days after the operative procedure AND infection involves any part of the body, excluding the skin incision, fascia, or muscle layers
that is opened or manipulated during the operative procedure AND the patient has at least ONE of the following:

a. purulent drainage from a drain that is placed into the organ/space
b. organisms isolated from an aseptically obtained culture of fluid or tissue in the organ/space
c. an abscess or other evidence of infection involving the organ/space that is found on direct examination, during invasive procedure, or by histopathologic examination or imaging test.
d. diagnosis of an organ/space SSI by a surgeon or attending physician and meets at least one criterion for a specific organ/space infection site listed in NHSN.

All study co-investigators agree to follow these CDC definitions of SSIs for study subjects enrolled in this trial to maximize objectivity of this study measure.

**Surgical Site Occurrence**

A surgical site occurrence (SSO) will be defined as a complication or adverse event occurring at the surgical site, including but not limited to, superficial, deep incisional, and organ/space surgical site infections. Consensus definitions and treatment plans for common SSOs following open complex ventral hernia repair were developed *a priori* by the co-investigators for the purposes of this study protocol (Appendix 6). All study co-investigators agree to follow these consensus definitions and treatment plans for study subjects enrolled in this trial to maximize objectivity of this outcome measure.
Surgical Site Occurrences Requiring Procedural Intervention

A surgical site occurrence requiring procedural intervention (SSOPI) will be defined as a complication or adverse event occurring at a surgical site that is managed or treated with an invasive procedure. Consensus definitions and treatment plans for common SSOs following open complex ventral hernia repair were developed a priori by the co-investigators for the purposes of this study protocol (Appendix 6). All study co-investigators agree to follow these consensus definitions and treatment plans for study subjects enrolled in this trial to maximize objectivity of this study measure.

Ventral Hernia Recurrence

A ventral hernia recurrence (HR) will be defined as any fascial defect of the anterior abdominal wall located within 7 cm of the index ventral hernia repair site detected by physical exam or abdominal computed tomography (CT) examination. These defects will be categorized as to whether they occur at the midline or parastomal hernia site, or both. Alternatively, for patients that are not amenable to come to the hospital for an in-person visit, recurrence will be assessed using a validated patient-reported outcome tool denominated the Ventral Hernia Recurrence Inventory (VHRI).[32] The VHRI is a short 3-question survey that can be applied in person or through a telephone contact, and was shown to have higher sensitivity and specificity to diagnose hernia recurrence than physical examination. Also, it was shown to have the ability to rule out
hernia recurrence when patients answer “no” to its questions. All study co-investigators agree to follow this consensus definition of ventral hernia recurrence for study subjects enrolled in this trial to maximize objectivity of this study measure.

**RISKS AND DISCOMFORTS**

As with any surgical procedure, there are some risks that are associated and they will be discussed in a separate surgical consent form. The subjects may experience some pain, bleeding and discomfort; however this is with any surgical operation. Common occurrences following hernia repair include seroma or hematoma around the hernia repair, inflammation, opening of the wound, or infection. Subjects may also experience additional therapies or treatments, including the removal of the mesh to treat any of these events.

**BENEFITS**

There are no direct benefits to subjects for participating in this study. Subject participation will help us better understand the comparative safety and effectiveness of synthetic versus biologic meshes in clean-contaminated and contaminated open ventral hernia repair.

**COSTS TO THE SUBJECTS**
There are no extra costs to the subjects associated with the research.

Procedures related to the hernia surgery are considered standard of care and will be the responsibility of the subject and the subject’s insurance company.

**ALTERNATIVES TO PARTICIPATION**

The subjects are under no obligation to participate in this study. The subjects may decide not to have mesh used for their hernia repairs. The PI or surgeon co-investigator at each site will discuss all available options. Those subjects not willing to participate in this study may be considered for the alternative treatment of primary defect closure/hernia repair with or without a similar biologic or synthetic product.

**PAYMENTS TO THE SUBJECTS**

Patients will not be paid for participation in this study.

**PLAN FOR OBTAINING INFORMED CONSENT**

For each subject, written informed consent will be obtained prior to any protocol-related activities. As part of this procedure, the principal investigator, surgeon co-investigator, or one of the approved study coordinators must explain orally and in writing the nature, duration, and purpose of the study in such a manner that the subject is aware of the potential risks, inconveniences, or adverse effects that may occur. The subjects will be informed that they may withdraw from the study at any time. Subjects will receive all information that is required by federal regulations.
After a potential study patient is identified, the investigator or the study coordinator listed in this protocol as a person who will obtain consent will be responsible for instituting the informed consent process in a face-to-face manner. Before starting any study procedures, the investigator will discuss the proposed research study in detail with the potential subject during the office visit to discuss treatment options. The subject will be allowed ample time to read and review the informed consent document, and ask questions. The informed consent document will be reviewed with the subject in depth by the participating investigator or designated member of the research team to ensure that the potential participant has a good understanding of the study protocol; what is required of the study participants; the potential risks and benefits of study participation; and his or her rights as a study participant. The investigators will be available by phone or office visit to answer any questions that the participant may have. After consideration, the subject may return if necessary for another visit with the investigator to discuss the study, ask questions, and sign the informed consent document to participate in this study.

After the subject has read and reviewed the informed consent document and has agreed to participate, he/she will be asked to sign and date the document. The study member obtaining consent will also sign and date the form, and documentation of the informed consent process will be included in the research file (i.e., the person who obtained consent, where and when consent was
obtained, and who was present during the process). A copy of the consent form will be given to the subject.

**PROVISIONS FOR SUBJECTS FROM VULNERABLE POPULATIONS**

The population to be studied includes adults 21 years of age or over, so children are therefore excluded. Decisionally-impaired and cognitively-impaired persons will not be approached to participate in this study as we are seeking subjects who have the capacity to understand and actively consent to the procedure independently. Pregnant women will be excluded from participating in this study.

Staff and employees of the participating sites (Cleveland Clinic, Greenville Health Systems Vanderbilt University Medical Center, and Washington University) are considered vulnerable populations. Staff and employees of any of the participating sites may be eligible to participate in this study. Since subjects may or may not benefit from this study, we do not want to exclude this population. If an employee is a potential candidate for this study, the subject will be informed during the consent process that his/her participation or refusal to will in no way influence grades, employment, or subsequent recommendations. Every effort will be made to prevent coercion during this initial process and throughout study participation. According to IRB policy, students and house staff cannot be asked to participate in research conducted while under the direct supervision of the investigator, so those subjects will not be enrolled.
In those instances where potential participants cannot read the consent form because they do not speak English, we will work with the IRB to develop a language-appropriate consent form. In addition, a qualified translator will be present to assist with obtaining the informed consent of the participant.

In addition, in the unusual situation where a subject cannot read a consent form due to illiteracy or blindness, a member of the research study staff will read and explain the consent form to the participant or to the participant’s legally-authorized representative. A witness, who will sign and date the consent form, must also be present during this oral presentation.

**SUBJECT PRIVACY AND DATA CONFIDENTIALITY**

Anonymity and confidentiality of subjects participating in this study will be maintained. The only potential identifiers on any study documents submitted to the sponsor or designee will be subject study numbers, dates of birth, and dates of procedures. Every effort will be made to maintain the confidentiality of documents that identify the subject by name (e.g., signed informed consent documents, clinic charts), except to the extent necessary to allow monitoring by the Center for Clinical Research at Cleveland Clinic, internal monitoring by any of the participating sites, or auditing by the FDA or other regulatory authorities. Should the name and/or address of a subject participating in this trial be on a document submitted to the FDA or other regulatory authority (e.g., laboratory
report), the name and/or address will be completely blocked out and replaced with the subject study number.

Additionally, patient charts and clinical records will be requested and reviewed so that protocol adherence and source documentation can be verified. There is a possibility that the Institutional Review Boards of any of the participating sites, the Food and Drug Administration, and possibly foreign regulatory agencies may review the de-identified study records. All information collected, such as name or medical record number, will be stored utilizing a customized Research Electronic Data Capture (REDCap®) database program for multi-institutional data collection. This is in a secure network/firewall protected electronic database to which only the investigator and the designated members of the study team will have access using an individual assigned login and password. Only approved study members listed on the IRB protocol will have access to the separately-stored master list. User rights will be assigned such that the designated research coordinator at each site may only enter and review data from that site. Only the Principal Investigator, Lead Research Coordinators, and Biostatisticians will be granted access to retrieve patient data from all sites for routine data quality assessments and data analyses. All electronic records pertaining to the clinical study will be password-protected, and only approved study members listed on the IRB protocol will have password access.
Any information about the subject collected on paper, as well as the subject enrollment log linking the subjects to their identifiers, will be kept under lock and key in the Department of Surgery at the corresponding participating site.

**Anticipated results and limitations**

We expect that the synthetic mesh may be associated with an initial higher rate of mesh infections than a biologic mesh. However, we also believe that these infections will be treatable with local measures and not require surgical excision of the mesh. We believe that light-weight macroporous materials will granulate and heal without long-term adverse outcomes. This should result in a significant reduction in long-term hernia recurrences with the synthetic material. Another concern with this trial is the ability to recruit enough patients meeting inclusion criteria. The multi-center nature of this study with well-established expert hernia surgeons with extensive experience in complex abdominal wall reconstruction should significantly reduce the risk of poor enrollment. Each program is a high-volume referral center for complex abdominal reconstructions performing on average of 4 cases per week. Finally, the ability to generalize our results to all available biologic mesh and synthetic mesh will not be possible. However, if the synthetic mesh is proven safe and effective in the single-stage repair of clean-contaminated and contaminated defects, it is likely that no surgeon will utilize any biologic graft costing over 100 times the price.

**Power Calculation**
We are investigating two primary outcomes: surgical site occurrences requiring procedural intervention (SSOPI) and hernia recurrence (HR). Surgical site occurrences requiring procedural intervention (SSOPI) is a repeated binary outcome measurement. Hernia recurrence (HR) is a single binary outcome measurement. In a multi-institutional retrospective review of clean-contaminated and contaminated ventral hernia repair, hernia recurrence rates were 29.21% versus 9.01%, and surgical site occurrences requiring procedural intervention were 39.02% versus 9.01% for the biological mesh and permanent synthetic mesh cohorts, respectively.[44] Based on review of these multi-institutional data, an estimated 253 patients will be enrolled in the proposed trial. Assuming a maximum 20% loss to follow-up, 202 patients will remain in the study sample for a 1:1 randomized allocation to each treatment arm (101 subjects per cohort). At the two-tailed overall (two hypotheses) type I error rate of 0.05, the study will have 92% power to detect a significant difference in the rates of hernia recurrence (29.21% vs. 9.01%), and a 100% power to detect a significant difference in surgical site occurrences requiring surgical intervention for the primary hypothesis (39.02% vs. 9.01%; with four repeated measurements and autoregressive correlation of rho=0.5) With 4 centers each performing approximately 1 eligible procedures per week, there will be 192 total procedures performed per year, from which it would be feasible to achieve the total enrollment goal of 253 subjects within 2 years.

Analysis

**Specific Aim 1** - To demonstrate that a single-stage repair of clean-contaminated (Class 2) or contaminated (Class 3) ventral hernias using a
macroporous light-weight polypropylene synthetic mesh will result in superior clinical outcomes compared to a biologic mesh.

**Primary Analyses of Primary Outcomes:**

The primary outcomes are rates of surgical site occurrences requiring procedural intervention (SSOPI) and hernia recurrence (HR) assessed postoperatively at 1 month, 6 months, 12 months, and 24 months. For SSOPI with 4 repeated binary measurements, a generalized mixed model analysis with repeated measures will be performed to determine if a significant difference exists between the biologic mesh and the permanent synthetic mesh cohorts in the rate of SSOPI. For HR, which is a single binary outcome, simple chi-square tests will be used for unadjusted analyses and a logistic regression model will be used for adjusted analyses. As this is a randomized trial, differences in baseline demographic and clinical characteristics between the biologic mesh cohort and the permanent synthetic mesh cohort are expected to occur at random. Any significant differences found among the demographic or preoperative clinical characteristics between the two treatment groups will be controlled for in the final analysis to limit potential confounding of results.

**Secondary Analyses of Primary Outcomes:**

These primary outcomes will also be assessed as time-to-event analyses (time-to-a-healed-wound and time-to-hernia-recurrence, respectively. In these analyses, the time-to-recurrence in the synthetic mesh group will be compared to
the time-to-recurrence in the biologic mesh group using a two-sided log-rank test. The null hypothesis is that the time-to-recurrence is the same between the two groups (H0: Hazard Ratio = 1), and the alternative hypothesis is that the time-to-recurrence is significantly greater for the permanent synthetic mesh cohort compared to the biologic mesh cohort (H1: Hazard Ratio not= 1). Since this is a randomized clinical trial, observed differences in other covariates between the two study groups are assumed to arise by chance. Therefore, the simple unadjusted analysis is sufficient to answer the research question, assuming adequate power (see above). However, since the association between other covariates and time-to-recurrence may be of interest and because group imbalances may occur due to chance, it is of interest to consider further analyses that adjust for covariates in the relationship of time-to-recurrence and mesh type.

To that end, the following multivariable models are pre-specified: 1) Demographics: age, race, gender, and mesh type; and 2) Known linkages to hernia recurrence: BMI, history of smoking, size of defect, number of previous hernia repairs, and mesh type. These models will be fit with Cox proportional hazards models. If other covariates are found to be highly imbalanced between groups or found to be associated with time-to-recurrence, they can also be included in multivariable models; however, this type of post-hoc model selection is to be considered inferior to the pre-specified models and is speculative in nature. A similar analysis will ensue for the time-to-a-healed-wound outcome. The null hypothesis is that the time-to-a-healed-wound is the same between the two groups (H0: Hazard Ratio = 1), and the alternative hypothesis is that the
time-to-a-healed-wound is significantly greater for the permanent synthetic mesh cohort compared to the biologic mesh cohort (H1: Hazard Ratio not= 1).

**Analyses of Secondary Outcomes:**
The secondary outcomes are pain and quality of life (QOL). The assessment of the association between these outcomes and mesh type will parallel what was described above for the primary outcome. The differences are as follows. First, the primary analysis will utilize an ANCOVA model adjusting for baseline pain (QOL) and correlations among repeated measurements. The null hypothesis is that the pain (QOL) is the same, on average, between the synthetic and biologic mesh groups (H0: Beta-mesh-type = 0). The alternative hypothesis is that the pain (QOL) is different, on average, between the two groups (H1: Beta-mesh-type not= 0). Second, the multivariable modeling will utilize multiple linear regressions. A mixed model analysis with repeated measures will be performed to determine if a significant difference exists between the biologic mesh and the permanent synthetic mesh cohorts in the mean change in score from the preoperative assessment to the postoperative assessments at 1 month, 6 months, 12 months, and 24 months for the EQ-5D and HerQLes quality of life instruments. The null hypothesis is that there is no significant difference between the biologic mesh and the permanent synthetic mesh cohorts in the mean change in score from the preoperative to the postoperative assessments at any of the time points for either the EQ-5D or the HerQLes quality of life instruments. The alternate hypothesis is that there is a significantly greater change in EQ-5D
and HerQLes scores from the preoperative assessment to the postoperative assessment at each of the time points for the permanent synthetic mesh cohort compared to the biologic mesh cohort.

In these data analyses, we will explore the possibility of medical center effect and treatment by medical center interaction. In particular, in the regression analyses, we will use indicator variables for medical center effect and indicator variable*treatment for possible treatment by medical center interaction. If treatment by medical center interaction is significant, then it is more appropriate that the treatment effects are summarized by medical center.

**Specific Aim 2** - Demonstrate that a macroporous light-weight polypropylene mesh is the more cost-effective strategy than a biologic prosthetic in clean-contaminated and contaminated abdominal wall reconstruction.

The first task to address this specific aim will be to estimate direct and indirect economic costs associated with clean-contaminated and contaminated open ventral hernia repair. A micro-costing approach will be used assuming a limited societal perspective. Costs associated with the preoperative, operative, and postoperative phases of care will be considered as it pertains to the management of the ventral hernia. In most cases, one to two preoperative clinic visits will be necessary. Final operative costs will be based on each patient’s actual inpatient encounter cost obtained from the participating institutions. Postoperative costs
will account for routine outpatient visits, costs incurred for complications, and
time lost from work due to ventral hernia management. All costs will be reported
in U.S. dollars, adjusted for the years in which the data were obtained. Should
institutional data not be available for a particular cost, the best available evidence
from currently published analyses will be used. The second task for this specific
aim is to perform health utility valuation in patients undergoing open repairs of
clean-contaminated and contaminated ventral hernias. Patients will be
administered the EQ-5D and the HerQLes instruments at one preoperative visit
and postoperatively at scheduled visits at 4 weeks (± 2 weeks), 6 months (± 2
month), and 12 months (± 3 months), and 24 months (± 4 months). HerQLes
and EQ-5D valuations will be converted to health utility estimates based on
published norms from the Agency for Healthcare Research and Quality (AHRQ).
Quality-adjusted life years (QALYs) will be calculated based on these health
utilities obtained in the postoperative phase. For a given strategy (light-weight
macroporous polypropylene mesh or biologic mesh), it is anticipated that health
utility variation will be affected more than survival. As such, calculations of
QALYs will assume that all patients survive during the 24 month follow-up period,
but differ in their health utility valuation.

**Cost-Effectiveness Evaluation**

Once costs and QALYs have been calculated for each strategy, a decision
analysis model will be constructed incorporating these values and the associated
outcomes and complications that are thought to have an impact on which
strategy is the better choice for patients. One-way and two-way sensitivity analyses will be performed to help determine the degree to which each variable impacts the decision to choose either a light-weight macroporous polypropylene mesh or biologic mesh for repair of clean-contaminated or contaminated ventral hernias. If appropriate, a multi-way probabilistic sensitivity analysis will be performed incorporating the uncertainty associated with known values for certain variables. Costs, effectiveness (as measured by QALYs), cost-effectiveness ratios, and incremental cost-effectiveness ratios will be calculated to help determine the best strategy for mesh selection based on the results from this study and published data.

INNOVATION

This study would represent the first prospective randomized trial comparing synthetic and biologic meshes in the single-stage open repair of complex ventral hernias. Currently, there is no level 1 clinical evidence to guide surgeons to the most appropriate prosthetic choice for these challenging cases. Given the lack of FDA regulations for clinical evidence to support the usage of these materials or provide comparative evaluations of these materials, it is unlikely that any company will perform this trial and potentially risk a negative outcome. These materials have been available for almost 20 years, and some products still have no published data supporting their usage, even in an animal model. This vacuum of evidence has left the surgeon to make decisions based largely on anecdotal experiences that are heavily influenced by strategic industry
marketing campaigns. These marketing campaigns have included generalized claims of synthetic mesh being easily infected, and requiring major surgical resection for removal leaving the patient with a very morbid operation and high re-herniation rate. In practice, newer lighter-weight meshes have actually been shown to be quite resistant to infection and often tolerate exposure and/or direct bacterial contamination. It is paramount to the thousands of patients that are affected by this condition, and the hundreds of millions of dollars spent on these expensive products by hospitals and insurance companies that their indications and contraindications are clearly defined in appropriately-designed and conducted clinical trials. Proving the potential advantages of utilizing a permanent synthetic mesh material in clean-contaminated and contaminated abdominal wall reconstruction could not only reduce healthcare expenditure by hundreds of millions of dollars, but also result in improved long-term patient outcomes with healed wounds and reduced recurrence rates. Our results will establish the standard of care for complex ventral hernia repair and guide the reconstructive surgeon on appropriate mesh selection.

References:


Figure 1:

A rodent model of incisional hernia and chronic infection after exposure to $10^4$ colony forming units of MRSA. Bacterial clearance as measured by total
resolution of bacterial growth. Strattice versus Soft Mesh p=0.32.