Abstract: A roll-up wound protector has a distal ring, a tricuspidate proximal ring, and a flexible sleeve extending between the proximal and distal rings. The proximal ring may have varying geometries and is Tollable to gather the flexible sleeve around the proximal ring and shorten the length of the flexible sleeve.
ROLL-UP WOUND PROTECTOR WITH TRICUSPIDATE RING

BACKGROUND

The present invention relates in general to surgical devices and procedures, and more particularly to wound protectors and wound retractors.

Surgical procedures are often used to treat and cure a wide range of diseases, conditions, and injuries. Many surgical procedures require access to internal tissue through open surgical procedures or endoscopic surgical procedures. The term "endoscopic" refers to all types of minimally invasive surgical procedures including laparoscopic and arthroscopic procedures. Endoscopic surgery has numerous advantages compared to traditional open surgical procedures, including reduced trauma, faster recovery, reduced risk of infection, and reduced scarring. Endoscopic surgery is often performed with an insufflatory fluid present within the body cavity, such as carbon dioxide or saline, to provide adequate space to perform the intended surgical procedures. The insufflated cavity is generally under pressure and is sometimes referred to as being in a state of pneumoperitoneum. Trocars are often used to provide a port through which endoscopic surgical instruments are passed. Trocars generally have a sealing valve that prevent the insufflatory fluid from escaping while an instrument is positioned in the trocar. Sometimes hand access devices are also used during endoscopic surgery, often referred to as hand assisted laparoscopic surgery ("HALS"). A HALS device will typically seal around a surgeon's hand or arm to prevent the insufflatory fluid from escaping while allowing the surgeon to manipulate tissue within the patient's body.

While wound protectors and wound retractors are known, no one has previously made or used a wound protector or wound retractor in accordance with the present invention.

BRIEF DESCRIPTION OF DRAWINGS

While the specification concludes with claims which particularly point out and distinctly claim the invention, it is believed the invention will be better understood from the following description taken in conjunction with the accompanying drawings illustrating some non-limiting examples of the invention. Unless otherwise indicated, the figures are drawn to scale and like reference numerals identify the same elements.

Fig. 1 depicts an non-scaled schematic view of a roll-up wound protector;
Fig. 2 depicts a non-scaled schematic view of a roll-up wound protector deployed in an abdomen;
Fig. 3 depicts a cross-sectional view of a proximal ring;
Fig. 4 depicts a cross-sectional view of a proximal ring;
Fig. 5 depicts a cross-sectional view of a proximal ring;
Fig. 6 depicts a cross-sectional view of a proximal ring;
Fig. 7 depicts a cross-sectional view of a proximal ring; and
Fig. 8 depicts a cross-sectional view of a proximal ring;

DETAILED DESCRIPTION

Fig. 1 depicts a perspective view of a roll-up wound protector in an extended position. The wound protector comprises a distal ring (10), a proximal ring (30), and a flexible sleeve (20) having a length extending between the proximal and distal rings. The wound protector can be used as a stand-alone device or in combination with a cap having sealing valve for endoscopic instruments or a surgeon’s arm.

In this embodiment the distal ring (10) is circular with a circular cross-sectional geometry; however, non-circular rings and non-circular cross-sectional geometries are also possible. For instance, the distal ring could have a oval or elliptical in cross-sectional shape. The distal ring (10) can be made from a variety of different materials with different characteristics. In this example the distal ring is made from an elastomer such as polyurethane, polyethylene, silicone, and the like. The distal ring can also vary in size. For instance, the distal ring can have an inside diameter greater than 1 inches and less than 9 inches, and a thickness less than 1 inch, but dimensions outside these ranges are also possible. Optionally, the distal ring (10) will have a durometer between 4OA and 9OA or 7OD, but other material properties are also possible.

In this embodiment the sleeve (20) is a single layered tube of material; however, a discontinuous sleeve or multi-layered sleeves are also possible. The sleeve (20) can be made from a variety of materials with different characteristics. In one example, the sleeve (20) is made from an elastomer such as polyisoprene, silicone, polyurethane, silicone, and the like; however, inelastic materials such as mylar could also be used. The sleeve (20) may be clear, transparent, translucent, or opaque. As shown here, the sleeve (20) is fastened at its ends directly to the proximal and distal rings using an adhesive or heat sealing techniques; however,
alternative techniques may also be employed. The sleeve (20) could also be attached to the rings at locations other than the sleeve ends. For instance, the sleeve (20) can wrapped around the distal ring (10) and adhesively attached or sealed to itself. The length of the sleeve (20) can also vary. For instance, the sleeve may be between 2 cm and 20 cm in length; however, other lengths are also possible. The thickness of the sleeve (20) can also vary. For instance, the sleeve thickness in this embodiment is between 0.010 and 0.020 inches; however, other thicknesses are also possible.

In this embodiment the proximal ring (30) is circular; however, non-circular rings are also possible. The proximal ring (30) can also vary in size. For instance, the proximal ring (30) can have an inside diameter between 1 and 9 inches, but other dimensions are also possible. Optionally, the ratio of the distal ring (I) and proximal ring (30) diameters is greater than 0.4. The proximal ring (30) in this example has a tricuspidate cross-sectional geometry. In this embodiment, the cross-sectional geometry is substantially constant around the circumference of the proximal ring (30). A geometry is substantially constant if any variations are insignificant. For example, geometric variations resulting only from molding or other manufacturing factors would be considered substantially constant. Also in this embodiment the cross-sectional geometry is substantially solid; however, holes or cavities may also be present.

Fig. 2 depicts an example of the wound protector in a deployed position in a patient. In this example the wound protector in positioned in a patient's abdominal wall (40) through an incision (46). The distal ring (10) is held in a collapsed position (e.g., in an oblong shape like an oval, a peanut, a figure eight, and the like) to reduce its size and then inserted through the incision (46). After insertion, the distal ring (10) is released and then expands to its ring-like shape. As shown here, the expanded distal ring (10) is larger than the incision (46) and sits against the peritoneal surface of the abdominal wall (40). The proximal ring (30) is rollable to gather the flexible sleeve (20) around the proximal ring (30), and the wound proximal ring (30) sits on the cutaneous surface of the abdominal wall (40). The proximal ring (30) is rollable in the outward directions (as shown by the arrows) to shorten the sleeve (20) and in the inward direction to lengthen the sleeve (20), or vice versa. The shortening of the sleeve (20) pulls the sleeve (20) taut against the incised wound (42, 44). As one with ordinary skill in the art will recognize, surgical procedures can be performed through the incision (46) and the sleeve (20) protects the incised wound (42, 44) from infection and contamination. In addition, the taut sleeve
(20) tends to pull the incised wound (42,44) open thus functioning as a wound retractor. As demonstrated in this example, more retraction is possible by rolling the proximal ring (30) outward, while less retraction is possible by rolling the proximal ring (30) inward.

In this example the proximal ring (30) rolls in resting increments of 360 degrees. In other words, when the ring rolls it "snaps" between resting positions. Optionally, the flip force for the proximal ring (30) can be less 10 in*lbs / 180 degrees of rotation, and can be less than 3 in*lbs. Flip force is a way of measuring the force required to roll the ring about itself. The flip force is measured at room temperature on a stand-alone proximal ring without the sleeve attached. An equal and opposite torque is applied simultaneously to a ring at two diametrically opposite points along the circumference of the ring. The peak measured torque to roll the ring is used to calculate the flip force. By compiling 100 peak measured torques for a given ring, the statistical median value is the flip force. Preferably, the flip force is substantially the same for each sequential resting incremental rotation. Optionally, the proximal ring (30) may have substantially no residual hoop stress. One way to achieve this is through a molding process where the proximal ring (30) is injection molded and transfer molded using a thermoplastic or thermoset elastomer such as polyisoprene, silicone, polyurethane, silicone, and the like. In one embodiment, the proximal ring is molded from Desmopan 9370. The proximal ring (30) may have a durometer between 5OA and 5OD, but other material properties are also possible.

Fig. 3 illustrates an example of a cross-sectional geometry of the proximal ring (30). The proximal ring (30) comprises a central axis (50). The cross-sectional shape shown in this example comprises three cusps (31, 33, 35) directed radially outward and spaced about 120 degrees from one another. The proximal cusp (31) points proximally parallel the central axis (50). The proximal cusp (31) has a flat tip normal the central axis (50), which may be used as a surface to which the sleeve (20) may be fastened. The distal cusps (33, 35) are aligned with each other along a plane normal the central axis (50). The distal cusps (33, 35) have convex arcuate tips. Concave arcuate recesses (32, 34, 36) extend between the cusps (31, 33, 35). The recesses are substantially constant about the circumference of the proximal ring (30). In the present embodiment the various surfaces transition smoothly, but geometric abrupt transitions are also possible.

Fig. 4 illustrates another example of a cross-sectional geometry of a proximal ring (60). The cross-sectional geometry is tricuspidate and is very similar to the proximal ring (30);
however, the proximal cusp (61) has a convex arcuate tip. In this example the cusps are all symmetric one another.

Fig. 5 illustrates yet another example of a cross-sectional geometry of a proximal ring (70). The cross-sectional geometry is tricuspidate, comprising three symmetric cusps (71, 73, 75) directed radially outward and spaced about 120 degrees from one another. The proximal cusp (71) points proximally and parallel the central axis. The distal cusps (73, 75) are aligned with each other along a plane normal the central axis. The cusps (71, 73, 75) each have a bulbous end portion, shown in this example as a generally circular cross-sectional shape. Concave recesses (72, 74, 76) extend radially inward relative the bulbous end portions, shown in this example as flats that intersect at about 120 degree angles. The surfaces are substantially constant about the circumference of the proximal ring (70). In the present embodiment the various surfaces transition smoothly, but geometric abrupt transitions are also possible. The proximal ring (70) rolls in increments of 360 degrees. Optionally, the proximal ring (70) is made with a molding process and may have substantially no residual hoop stress.

Fig. 6 illustrates still another example of a cross-sectional geometry of a proximal ring (80). The cross-sectional shape approximates an equilateral triangle, with three symmetric cusps (81, 83, 85) directed radially outward and spaced about 120 degrees from one another. The proximal cusp (81) points proximally and parallel the central axis. The distal cusps (83, 85) are aligned with each other along a plane normal the central axis. The cusps (81, 83, 85) each have an arcuate rounded tip. Flat surfaces (82, 84, 86) extend between the cusps (81, 83, 85). The surfaces are substantially constant about the circumference of the proximal ring (80). In the present embodiment the various surfaces transition smoothly, but geometric abrupt transitions are also possible. The proximal ring (80) rolls in increments of 360 degrees. Optionally, the proximal ring (80) is made with a molding process and may have substantially no residual hoop stress.

Fig. 7 illustrates another example of a cross-sectional geometry of a proximal ring (90). The cross-sectional shape approximates an equilateral triangle, with three symmetric cusps (91, 93, 95) directed radially outward and spaced about 120 degrees from one another. The lateral cusp (93) points laterally and in alignment with a plane normal the central axis. In this example, the plane bisects the proximal ring (90). The proximal and distal cusps (91, 95) point at oblique angles relative the central axis and are aligned with each other parallel the central axis. The cusps
(91, 93, 95) each have an arcuate rounded tip. Outer flat surfaces (92, 94, 96) extend between the cusps (91, 93, 95). The surfaces are substantially constant about the circumference of the proximal ring (90). In the present embodiment the various surfaces transition smoothly, but geometric abrupt transitions are also possible. The proximal ring (90) rolls in increments of 360 degrees. Optionally, the proximal ring (90) is made with a molding process and may have substantially no residual hoop stress.

Fig. 8 illustrates another example of a cross-sectional geometry of a proximal ring (100). The cross-sectional shape approximates an equilateral triangle, with three symmetric cusps (101, 103, 105) directed radially outward and spaced about 120 degrees from one another. The medial cusp (105) points medially and in alignment with a plane normal the central axis. In this example, the plane bisects the proximal ring (100). The proximal and distal cusps (101, 103) point at oblique angles relative the central axis and are aligned with each other parallel the central axis. The cusps (101, 103, 105) each have an arcuate rounded tip. Outer flat surfaces (102, 104, 106) extend between the cusps (101, 103, 105). The surfaces are substantially constant about the circumference of the proximal ring (100). In the present embodiment the various surfaces transition smoothly, but geometric abrupt transitions are also possible.

The proximal ring (100) includes a circumferential cavity (107), shown in this example having three crooks aligned with each cusp (101, 103, 105). A biasing member (108), such as a full or partial ring made from an elastomer or metal, is positioned in medial crook of the cavity (107). As the proximal ring (100) is rolled, the biasing member (108) will leave the medial crook and track the surface of the cavity (107). The biasing member will also circumferentially expand, thus inducing a hoop stress on the biasing member (108). After the proximal ring (100) is rolled more than 60 degrees, the biasing member (108) will tend to relieve the hoop stress and “snap” to the next crook and rotate that crook to the medial position. Thus, proximal ring (100) rolls in resting increments of 120 degrees.

One way to make the proximal ring (100) involves extruding a length of material with the desired cross-sectional geometry, bending the length into a ring and inserting the biasing member (108) into the cavity (107), and then fastening the ends of the length together using a coupling and/or fastening techniques (e.g., adhesives, heat welding, ultrasonic welding, and the like). Optionally, the assembled proximal ring (100) may be heat cured to reduce hoop stresses induced during the bending step.
Preferably, the wound protectors described above will be processed before surgery. First, a new or used wound protector is obtained and if necessary cleaned. The wound protector can then be sterilized. In one sterilization technique the wound protector is placed in a closed and sealed container, such as a plastic or TYVEK bag. Optionally, the wound protector can be bundled in the container as a kit with other components, including one or more of the following: a sealing cap to maintain pneumoperitoneum, a sealing cap with a valve to allow passage of surgical instruments or a surgeon's arm while maintaining pneumoperitoneum (e.g., iris valve, gel seal, cuff, and the like), a tube of lubricant, a mounting ring in which the proximal ring may be seated and to which a cap can be attached, a marker, an incision template or scale, an instruction sheet, and the like. The container and wound protector, as well as any other components, are then placed in a field of radiation that can penetrate the container, such as gamma radiation, x-rays, or high-energy electrons. The radiation kills bacteria on the wound protector and in the container. The sterilized wound protector can then be stored in the sterile container. The sealed container keeps the wound protector sterile until it is opened in the medical facility.

The wound protectors described above can be used as a stand-alone device, for instance in open surgical procedures, or in combination with a cap having sealing valve for endoscopic instruments or a surgeon's arm. Among other advantages, the foregoing examples provide effective wound protection to prevent infection and facilitate wound retraction. Because the sleeve rolls-up, its length can be adjusted by the surgeon for any given anatomy and patient. Further, the surgeon can select the amount of retraction desired for a given procedure. The cross-sectional shapes of the proximal ring are easy to grip thus facilitating ease of use. Furthermore, the flip forces are relatively low and constant, further facilitating ease of use.

Having shown and described various embodiments and examples of the present invention, further adaptations of the methods and devices described herein can be accomplished by appropriate modifications by one of ordinary skill in the art without departing from the scope of the present invention. Several of such potential modifications have been mentioned, and others will be apparent to those skilled in the art. For instance, the specific materials, dimensions, and the scale of drawings will be understood to be non-limiting examples. Accordingly, the scope of the present invention should be considered in terms of the following claims and is
understood not to be limited to the details of structure, materials, or acts shown and described in the specification and drawings.
CLAIMS

1. A surgical wound protector, comprising:
   
   a) a distal ring;

   b) a proximal ring having a cross-sectional geometry, the cross-sectional geometry comprising three cusps directed radially outward and spaced about 120 degrees from one another and an outer surface with concave recesses extending between the cusps; and

   c) a flexible sleeve having a length extending between the proximal and distal rings;

   wherein the proximal ring isrollable to gather the flexible sleeve around the proximal ring and shorten the length of the flexible sleeve.

2. The surgical wound protector of claim 1, wherein the cross-sectional geometry is solid.

3. The surgical wound protector of claim 1, wherein the cross-sectional geometry is substantially constant around the proximal ring.

4. A sterile surgical kit, comprising a sealed container containing therein the surgical wound protector of claim 1 and a sealing cap with a valve.

5. The surgical wound protector of claim 1, wherein the proximal ring has substantially no residual hoop stress.

6. The surgical wound protector of claim 1, wherein the proximal ring rolls in resting increments of 360 degrees.

7. The surgical wound protector of claim 1, wherein the concave recesses are arcuate.
8. A method of processing a wound protector for surgery, comprising:
   a) obtaining the surgical wound protector of claim 1;
   b) sterilizing the wound protector; and
   c) storing the wound protector in a sterile container.

9. A surgical wound protector, comprising:
   a) a distal ring;
   b) a proximal ring having three circumferential cusps spaced about 120 degrees from one another and having substantially no residual hoop stress; and
   c) a flexible sleeve extending between the proximal and distal rings;

   wherein the proximal ring is rollable to gather the flexible sleeve around the proximal ring.

10. The surgical wound protector of claim 9, wherein the proximal ring rolls in resting increments of 360 degrees.

11. The surgical wound protector of claim 9, wherein the proximal ring defines a plane and one of the cusps points in alignment with the plane.

12. The surgical wound protector of claim 11, wherein the proximal ring points medially.

13. A sterile surgical kit, comprising a sealed container containing therein the surgical wound protector of claim 9 and a sealing cap with a valve.
14. A surgical wound protector, comprising:

   a) a distal ring;

   b) a rollable proximal ring defining a central axis, the proximal ring comprising three cusps, one of the cusps pointing in alignment with a plane normal the central axis, and the other cusps pointing at oblique angles relative the central axis; and

   c) a flexible sleeve extending between the proximal and distal rings.

15. The surgical wound protector of claim 14, wherein the proximal ring has substantially no residual hoop stress.

16. The surgical wound protector of claim 14, wherein the proximal ring rolls in resting increments of 120 degrees.

17. The surgical wound protector of claim 14, wherein the cusps are spaced about 120 degrees from each other.

18. The surgical wound protector of claim 14, wherein the proximal ring comprises a circumferential cavity with three crooks.

19. The surgical wound protector of claim 18, further comprising a biasing member positioned in the cavity.

20. A sterile surgical kit, comprising a sealed container containing therein the surgical wound protector of claim 14 and a sealing cap with a valve.
# International Search Report

## A. Classification of Subject Matter

### INV.

- A61B17/34 A61B17/02

### ADD.

- A61B17/00

According to International Patent Classification (IPC) or to both national classification and IPC

## B. Fields Searched

Minimum documentation searched (classification system followed by classification symbols)

- A61B

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

## C. Documents Considered to be Relevant

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X Further documents are listed in the continuation of Box C  

X See patent family annex

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Date of the actual completion of the international search

- 27 November 2007

Date of mailing of the international search report

- 05/12/2007

Name and mailing address of the ISA/

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