METHOD AND DEVICE FOR USE IN ENDOSCOPIC ORGAN PROCEDURES

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ABSTRACT

Methods and devices for use in tissue approximation and fixation are described herein. The present invention provides, in part, methods and devices for acquiring tissue folds in a circumferential configuration within a hollow body organ, e.g., a stomach, positioning the tissue folds for affixing within a fixation zone of the stomach, preferably to create a pouch or partition below the esophagus, and fastening the tissue folds such that a tissue ring, or stoma, forms excluding the pouch from the greater stomach cavity. The present invention further provides for a liner or bypass conduit which is affixed at a proximal end either to the tissue ring or through some other fastening mechanism. The distal end of the conduit is left either unanchored or anchored within the intestinal tract. This bypass conduit also includes a fluid bypass conduit which allows the stomach and a portion of the intestinal tract to communicate.
METHOD AND DEVICE FOR USE IN ENDOSCOPIC ORGAN PROCEDURES

CROSS-REFERENCE TO RELATED APPLICATIONS


FIELD OF THE INVENTION

[0002] The present invention relates generally to medical apparatus and methods and more particularly to devices and methods for dividing a hollow body organ or otherwise restricting or partitioning a certain section of that organ, such as a stomach, intestine or gastrointestinal tract as well as devices and methods for placing a liner within or partially within the hollow body organ.

BACKGROUND OF THE INVENTION

[0003] In cases of severe obesity, patients may currently undergo several types of surgery either to tie off or staple portions of the large or small intestine or stomach, and/or to bypass portions of the same to reduce the amount of food desired by the patient, and the amount absorbed by the gastrointestinal tract. The procedures currently available include laparoscopic banding, where a device is used to “tie off” or constrict a portion of the stomach, vertical banded gastroplasty (VBG), or a more invasive surgical procedure known as a Roux-En-Y gastric bypass to effect permanent surgical reduction of the stomach’s volume and subsequent bypass of the intestine.

[0004] Typically, these stomach reduction procedures are performed surgically through an open incision and staples or sutures are applied externally to the stomach or hollow body organ. Such procedures can also be performed laparoscopically, through the use of smaller incisions, or ports, through trocars and other specialized devices. In the case of laparoscopic banding, an adjustable band is placed around the proximal section of the stomach reaching from the lesser curve (LC) of the stomach around to the greater curve (GC), thereby creating a constriction or “waist” in a vertical manner between the esophagus (ES) and the pylorus (PY) (See Prior Art FIG. 1). During a VBG (See Prior Art FIG. 2) a small pouch (P) (approximate 20 cc in volume) is constructed by forming a vertical partition from the gastroesophageal junction (GEJ) to midway down the lesser curvature of the stomach by externally applying staples, and optionally dividing or resecting a portion of the stomach, followed by creation of a stoma (ST) at the outlet of the partition to prevent dilation of the outlet channel and restrict intake. In a Roux-En-Y gastric bypass (see Prior Art FIG. 3), the stomach is surgically divided into a smaller upper pouch connected to the esophageal inflow, and a lower portion, detached from the upper pouch but still connected to the intestinal tract for purposes of secreting digestive juices. A resected portion of the small intestine is then anastomosed using an end-to-side anastomosis to the upper pouch, thereby bypassing the majority of the intestine and reducing absorption of caloric intake and causing rapid “dumping” of highly caloric or “junk foods”.

[0005] Although the outcome of these stomach reduction surgeries leads to patient weight loss because patients are physically forced to eat less due to the reduced size of their stomach, several limitations exist due to the invasiveness of the procedures, including time, general anesthesia, healing of the incisions and other complications attendant to major surgery. In addition, these procedures are only available to a small segment of the obese population (morbid obesity, Body Mass Index≥40) due to their complications, leaving patients who are considered obese or moderately obese with few, if any, interventional options.

[0006] In addition to surgical procedures, certain tools exist for approximating or otherwise securing tissue such as the stapling devices used in the above-described surgical procedures and others such as in the treatment of gastroesophageal reflux (GERD). These devices include the GIAR® device (Gastrointestinal Anastomosis device manufactured by Ethicon Endosurgery, Inc. and a similar product by USCC), and certain clamping and stapling devices as described in U.S. Pat. Nos. 5,897,562 and 5,571,116 and 5,676,674. Non-Invasive Apparatus for Treatment of Gastroesophageal Reflux Disease (Bolanos, et al) and 5,403,326 Method for Performing a Gastric Wrap of the Esophagus for Use in the Treatment of Esophageal Reflux (Harrison et al) for methods and devices for fundoplication of the stomach to the esophagus for treatment of gastro esophageal reflux (GERD). In addition, certain tools as described in U.S. Pat. No. 5,947,985 Tissue Cutting and Stitching Device and Method (Solar et al), detail an endoscopic suturing device (C.R.Bard, Inc., Billerica, Mass.) that is inserted through an endoscope and placed at the site where the esophagus and the stomach meet. Vacuum is then applied to acquire the adjacent tissue, and a series of stitches are placed to create a pleat in the sphenicter to reduce the backflow of acid from the stomach up through the esophagus. These devices can also be used transorally for the endoscopic treatment of esophageal varices (dilated blood vessels within the wall of the esophagus).

[0007] Further, certain devices are employed to approximate tissue such as in U.S. Pat. No. 5,355,897 (Petriflata) describing the use of a circular stapler to perform a pyloroplasty to create a narrowing at the pylorus. In addition, intraluminal anastomosis, such as bowel anastomosis, use suturing or stapling and employ tools such as the circular stapler, such as that described in U.S. Pat. Nos. 5,309,927 (Welch), 5,588,579 (Schnur et al), 5,639,008 (Gallagher et al), 5,697,943 (Sauer), 5,839,639 (Sauer), 5,860,581 (Roberson et al), and 6,119,913 (Adams et al). Such circular staplers are available from Ethicon Endosurgery, Cincinnati, Ohio (Proximate™ and EndoPath Stealth™ staplers, see www.surgicalstapling.com), Power Medical Interventions, New Hope, Pa., and United States Surgical, a unit of Tyco Healthcare Group LP, Norwalk, Conn.

[0008] There is a need for improved devices and procedures. In addition, because of the invasiveness of most of the surgeries used to treat obesity, and the limited success of others, there remains a need for improved devices and methods for more effective, less invasive hollow organ restriction procedures.

SUMMARY OF THE INVENTION

[0009] The present invention provides for improved methods and apparatus for the transoral, or endoscopic, restriction of a hollow body organ, such as the creation of a small stomach pouch. For purposes of the present invention, the hollow body organ shall include the entire gastrointestinal tract, including, but not limited to, the esophagus, stomach, portions of or the entire length of the intestinal tract, etc., unless specified otherwise. In the case of the present inven-
tion, the surgeon or endoscopist may insert devices as described below through the patient’s mouth, down the esophagus and into the stomach or intestine as appropriate. The procedure can be performed entirely from within the patient’s stomach or other organ, and does not require any external incision. The end result of the procedure is the formation of a variety of organ divisions or plications that serve as barriers or “partitions” or “pouches” that are substantially sealed off from the majority of the organ cavity. For example, in the case of dividing the stomach, the “pouch” or partitions that are created may seal a small portion of the stomach just below the esophagus to allow only small amounts of food or liquid to be consumed by the patient. This pouch or partition will mimic the section of stomach sealed off from the majority of the organ in a traditional obesity surgery heretofore described; however, it can be formed and secured entirely from inside the stomach endoscopically, obviating the need for a prolonged procedure, external incisions, minimizing the risk of infections, and in some cases, general anesthesia.

[0010] The methods and tools of the present invention may also be used in treating GERD in that stomach folds just below the esophagus can be acquired and fastened to create a desired “pleat”, thereby effectively extending the length of the esophagus and preventing reflux. Preferably, multiple folds of tissue can be acquired to effect this end. Further, features of the present invention would assist in the longevity of the GE Junction (GEJ)/Esophageal pleat as compared to current devices and techniques as the plication would include a more significant amount of muscular tissue. In addition, the devices and methods of the present invention may be used to revise or repair failures seen in current surgical procedures, such as dilation of the pouch and/or stoma (stomata) formed in a traditional Roux-En-Y gastric bypass, or VBG. In these cases, when the stoma dilates or shifts, the tools of the present invention would be useful to circumferentially gather tissue at the site of dilation to narrow it, thereby making the stoma functional again, or by further reducing the volume of an existing pouch which has dilated.

[0011] The devices shown and described herein can be used to form a pouch or partition by the approximation and fixation of a circular section of tissue acquired circumferentially from the wall of the target organ. The tissue acquisition device and fastener may include an acquisition feature (utilizing, e.g., a vacuum, and/or some other mechanical method for acquiring a circumferential “bite” of tissue), a fixation element (such as a stapling mechanism) and possibly a cutting element. In addition, the device may be adapted to receive a standard endoscope to allow viewing of the target region at various points during the procedure. The devices may be articulatable through a variety of conventional methods; alternatively, they may be articulated by an endoscope or other articulation device inserted within.

[0012] The fastening assembly of the present invention may employ a similar design and function to those circular staplers heretofore referenced, taking advantage of their ability to deploy multiple rows of staples with one actuation, and their relative clinical efficacy in performing other types of fastening (e.g. anastomoses procedures, hemorrhoid plication, etc.). Such devices can be adapted to perform the novel procedures described herein. Such devices may be adapted to incorporate a tissue acquisition system within the stapler body to allow sufficient tissue to be acquired during a procedure, and other modifications may be done to enable use of the stapler in these novel procedures.

[0013] In the procedures of the present invention relating to treatment of gastric disorders such as gastroesophageal reflux disease (GERD), or in cases of treating obesity, a flexible circular stapler may be inserted transorally down the patient’s esophagus and into the stomach at the region of the GEJ. Tissue may then be acquired circumferentially about the stapler device, or at least partially about the circumference of the stapler device at some point less than 360 degrees (possibly in a 180 degree formation) relative to a longitudinal axis of the device such that the tissue acquisition creates a “waist” within the organ volume. Subsequently, the tissue fixation element may then be deployed to fix the tissue in a manner to promote healing.

[0014] As set forth in U.S. patent application Ser. No. 10/188,547 filed Jul. 2, 2002, which is fully incorporated herein by reference in its entirety, the layered tissue structure of, e.g., the stomach, and the amount of desirable tissue acquisition and approximation is described in further detail. The devices and procedures of the present invention would allow the operator to reliably acquire and secure the necessary type of tissue, such as the muscularis, in creating the circumferential or curved tissue plication desirable to ensure a lasting clinical result.

[0015] Any of the fastening devices described herein may employ, e.g., bioabsorbable or biofragmentable staples or fixation element. Such fastening devices would typically dissolve or otherwise degrade leaving only the fixation region once the desired tissue healing has occurred. The remaining healed tissue, now a tissue “ring” (TR), would be sufficiently adhered or healed together to maintain the integrity of the pouch and stoma. In addition, the fastening devices may include coatings or other secondary features to aid healing, such as resorbable meshes, sclerosing agents, surgical felt, or tissue grafts.

[0016] The pouch or partitions may be created by a procedure of the present invention to remain permanently within the stomach to restrict it indefinitely. Alternatively, the creation of the pouch or partitions may be reversible (e.g., once weight loss is achieved, or reflux minimized) or revised (in the event pouch side needs to be modified). Reversal can also be achieved via various methods such as dilation of the restricted section, or, e.g., using an electro-surgical device such as a Bowie to cut the restricted section to free the tissue folds. Further, if the physician so desires, techniques of the present invention may be augmented or assisted by the use of other techniques such as laparoscopy. Optionally, techniques of the present invention may be combined with other procedures such as for the treatment of GERD or the transoral placement of a bypass prosthesis or other type of liner in the intestine to bypass the hormonally active portion of the small intestine, typically between the stoma to just proximal of the jejunum. Such a liner may be placed within the orifice of a stoma created by devices described herein or within stomas created by various conventional procedures, as also described herein. For present purposes, a stoma refers simply to an artificial or “man made” narrowing within a body organ. The liner may be tubular in construction and made to match the diameter of the stoma created by the present invention such that they can be hooked together to achieve the desired clinical effect. Additionally, the distal end of the liner may also be anchored to tissue distally located from the stoma or it may be left unanchored relying on its resilient physical structure to avoid kinking or twisting.
Moreover, such a liner may vary in construction and in placement within the stomach. The liner, which acts as a bypass conduit, may also include fenestrations or openings that provide for fluid communication between the stomach cavity (for instance, following a bypass procedure the remaining stomach cavity is commonly referred to as the "gastric remnant") and/or common duct (e.g., the duct that enters the intestine at the duodenal ampulla), and certain parts of the intestinal tract to maintain alimentary flow of digestive secretions. Allowing such flow may facilitate in preventing adhesions from forming between the liner and regions of the intestines. Such adhesions may typically cause blockage of the common duct with potentially fatal consequences, such as bowel necrosis. The liner may also include a secondary fluid conduit adjacently positioned along the liner to provide for fluid communication. The fluid conduit may thus have a length which is less than, greater than, or equal to a length of the liner and sufficient to communicate from the inflow point (e.g., gastric remnant or duodenal ampulla) and a point in the lower intestine (e.g., near the jejunum). The liner and fluid conduit may also be configured to ensure that the liner and/or fluid conduit does not inhibit fluid communication from the common bile ducts, such as channels or fenestrations along their length. The fluid conduit may be attached to the liner as a parallel tube or in any number of configurations. Another variation may have the fluid conduit as a coaxial tube positioned along the liner.

In either case, the liner may define one or more fenestrations or channels on the portion of the liner in communication with the gastric remnant, and/or at or near the site of the common bile ducts so as to allow fluids to drain from the organ or ducts. The liner and the fluid conduit may be made separately and attached together or they may be made integrally from the same material. Also, the liner and/or the fluid conduit may be made of a braided design to inhibit kinking as the device reacts to the peristalsis motion of the intestines. Another alternative may utilize a singular liner having one or more channels defined longitudinally along the outer surface of the liner rather than as a separate fluid conduit. These channels may form spaces between the tissue and the liner itself to allow for the flow of fluids within the channels. In another variation of the singular liner, the liner may have fenestrations or openings positioned along its length near or at the zones of active secretion in the intestines to permit fluid flow from the organ or bile ducts into the lumen of the liner (so as to prevent blockage thereof), while still maintaining a barrier to the majority of the intestine to achieve malabsorption and to facilitate "dumping" syndrome upon ingestion of high fat or high caloric foods. An alternative variation of this singular liner may have multiple valved openings along its length to allow for the redirection of flow of secretions into the liner, but prohibiting contact between the intestines and the food contents within the liner.

BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 depicts the prior art procedure commonly known as laparoscopic banding;

FIG. 2 depicts the prior art procedure commonly known as the vertical banded gastroplasty or "VBG";

FIG. 3 depicts the prior art procedure commonly known as surgical Roux-En-Y procedure;

FIG. 4A-4B depicts one variation on a procedure of the present invention, showing a cut-away section of the tissue being acquired by the distal tip of the device of the present invention, and the resulting modification to the body organ (creation of a "pouch" within the stomach);

FIGS. 5A-5D depict one variation of procedural steps of performing the methods of the present invention, by showing a cross section of an organ (stomach) and the placement of the device to create a narrowing or "pouch" within the organ;

FIG. 5E depicts one variation of a result of the present invention, including a bypassing sleeve installed to bridge from the point of the stoma at the GEJ, to the pylorus, or further into the intestine;

FIGS. 6A-6I shows a schematic depiction of an organ (stomach) following completion of one variation on a procedure of the present invention and the resulting cross sectional view of the treated region in various configurations;

FIGS. 7A-7F show a variation on the circular tissue acquisition and fixation device of the present invention, including details on the inner working elements and flexible shaft thereof;

FIG. 8 depicts details of one variation on the distal portion of the circular tissue acquisition and fixation device of the present invention showing an angled annular acquisition space;

FIG. 9 depicts another variation of the tissue acquisition mechanism of the circular tissue acquisition and fixation device of the present invention;

FIGS. 10A-10B depict variations of the distal working end of the distal tip of the circular tissue acquisition and fixation device of the present invention, detailing an anvil designed to be intraprocedurally manipulated to assist in removal of the circular tissue acquisition and fixation device of the present invention once the desired tissue has been acquired and fixed according to the present invention.

FIG. 11A depicts a variation of a bypass conduit assembly.

FIGS. 11B-11E depict variations on possible cross sections of the bypass conduit assembly.

FIG. 11F depicts a variation of the bypass conduit assembly having an irregular cross section.

FIG. 12 depicts another variation of the bypass conduit assembly but with the addition of a fluid bypass conduit located adjacent the conduit wall.

FIGS. 13A-13B depict perspective and cross-sectional views, respectively, of another variation of the bypass conduit having a coaxial fluid bypass conduit.

FIG. 14 depicts a perspective view of a braided tubular structure which may be utilized for the bypass conduit.

FIGS. 15A-15B depict variations on anchoring devices for the bypass conduit.

FIGS. 16A-16B depict a bypass conduit with a fluid bypass conduit deployed within a stoma created by a laparoscopic banding procedure.

FIGS. 17A-17B depict a bypass conduit with a coaxial fluid bypass conduit deployed within a stoma created by a laparoscopic banding procedure.

FIGS. 18A-18B depict a bypass conduit with spaced apart fenestrations deployed within a stoma created by a vertical banded gastroplasty procedure.

FIGS. 19A-19B depict a bypass conduit having valved fenestrations deployed within a stoma created by laparoscopic banding to construct the stomach cavity and create a stoma.
FIGS. 19C-19F depict variations on maintaining fluid communication through or along the bypass conduit.

FIGS. 20A-20B depict a bypass conduit deployed within a stomach which has an intragastric staple line.

FIGS. 21A-21B depict a bypass conduit deployed within a stoma created by a horizontal gastroplasty procedure.

FIGS. 22A-22B depict a bypass conduit deployed within a stoma created by a biliopancreatic diversion procedure.

DEDICATED DESCRIPTION OF THE INVENTION

The present invention provides, in part, for methods and devices for hollow organ division and restriction, more particularly providing methods and devices to perform a transoral, endoscopically mediated stomach reduction for purposes of, e.g., treating obesity. For purposes of the present invention, the hollow body organ shall include the entire gastrointestinal tract, including, but not limited to, the esophagus, stomach, portions of the entire tract, etc., unless specified otherwise.

As previously discussed, the results of some clinical procedures of the prior art are shown in FIGS. 1-3, from a perspective external to the stomach. An example of a result of the procedure in one variation of the present invention is shown in FIG. 4A, which depicts an external anterior view of a stomach organ 100, having an esophagus 101 (cut away to reveal the esophageal lumen 102), and further depicting a circumferential orifice or stoma 103, configured from staple line 104, producing a pouch (P). Orifice 103 is preferably positioned close to and on the distal side of the gastroesophageal junction (GEJ) at the base of the esophagus, and angled toward the lesser curve of the stomach (LC), leaving a stoma or opening having a diameter of approximately 1 cm between the pouch (P) and the remaining stomach volume. A desirable pouch (P) volume is between 15-100 cc, preferably 15-20 cc. The orifice 103 operates to restrict food from entering from the pouch, while still allowing communication between the pouch and the greater stomach volume for purposes of passage of digestive fluids and secretions and absorption of nutrients. FIG. 4B depicts an example of a cross sectional view of the esophagus where it joins the stomach, and further depicts one variation of a tissue acquisition device of the present invention 105, actively engaging the tissue to be fastened in a circumferential fashion.

Method of Hollow Organ Volume Reduction

A clinical work-up, including a physical and mental assessment of the patient may be performed to determine whether a transoral stomach reduction clinically indicated. This assessment may include inspecting the esophagus and stomach of the patient to determine whether any contraindications exist for undertaking the procedure such as ulcers, obstructions, or other conditions that may preclude treatment. Once the assessment has been completed, either in an operating room with the patient under general anesthesia, or in an endoscopy suite with the patient under sedation, the operator can introduce a tissue acquisition and fixation device, as shown in FIGS. 5A-5D, down the patient’s esophagus and into the stomach to a location just beyond the GE Junction (GEJ). Once in place, an optional calibration device (not shown) such as a balloon or bougie can be inflated or deployed proximally or adjacently to the GE Junction (GEJ) to assist in correctly sizing the pouch to be created. Alternatively, the physician may opt to use direct vision and place an endoscope through the main lumen of the tissue acquisition device to view the site of entry and resultant treatment zone.

FIGS. 5A through 5D depict cross sectional schematic views of the procedure of the present invention showing tissue being manipulated within a hollow organ, the stomach. FIG. 5A depicts the esophagus (ES) the stomach cavity (SC), including the landmarks of the lesser curve of the stomach (LC), the gastroesophageal junction (GEJ), and the pylorus (PY). Tissue layers represented are the serosal layer (SL), the muscularis or fibrous muscular layer (ML), and the mucosal layer (MUC). Further, FIG. 5A shows the tissue acquisition device 105 positioned within the esophagus at a location within the stomach cavity (SC) between the lesser curve (LC) of the stomach and the GEJ.

The device 105 includes a main body 106 having at least one lumen therethrough (not shown), an outer portion 107, having a distal end 108 containing a fixation mechanism and a proximal end (not shown). The device 105 further comprises an inner portion 109, which has a distal portion 110 containing a fixation mechanism and a proximal portion (not shown) received therein. Once device 105 is positioned in the preferred anatomical location, outer portion distal end 108 and inner portion distal end 110 are separated by relative movement of inner portion 109 within outer portion 107, to expose opening 112. As described in further detail later below, opening 112 is operatedly connected to at least one lumen within the main body 106 and provides a force, e.g., a vacuum force, to facilitate tissue acquisition. Such a force may be provided by a vacuum or by a mechanical element.

As shown in FIG. 5B, in the case of vacuum, once the opening 112 is exposed to the surrounding tissue within the stomach cavity (SC), the vacuum may be activated and tissue 111 may be drawn into the opening 112 in an entirely circumferential manner or a substantially circumferential manner, i.e., at least partially about the circumference of the device at some point less than 360 degrees (possibly in a 180 degree formation) relative to a longitudinal axis of the device. The amount of tissue 111 acquired can vary, but the amount drawn is preferably sufficient enough to result in healing of the fastened sections, thereby creating a tissue ring (TR) around the circumference of the fastened tissue. Said tissue ring may be formed of various layers of the stomach and may include scar tissue and other elements of effective wound healing.

FIG. 5C further depicts the device 105 after the desired amount of tissue 111 has been acquired, outer portion distal end 108 and inner portion distal end 110 may be moved towards one another such that the acquired tissue 111 is clamped therebetween. Device 105 is then actuated to engage at least one fastening element (not shown) through the acquired tissue 111 thereby fastening it in place in a circumferential fashion. This fastening step may also include a cutting step to score or otherwise abrade the acquired tissue 111 after it is fastened to enhance the healing response of the tissue 111 to increase the durability of the tissue ring. In addition, bulking agents, such as collagen, may be injected at the time the stoma is formed, or thereafter, to aid in healing and durability of the tissue. Once the tissue 111 has been fastened or fixed, the tissue acquisition device 105 is then removed. In doing so, the inner portion distal end 110 of the device may be carefully pulled through the newly-created tissue ring or stoma created by the procedure so as to mini-
mize stretching of the ring or stoma. Finally, FIG. 5D depicts
the stomach showing the final result and placement of a
circumferential tissue ring (TR) or stoma (ST).

As depicted in FIG. 5E, it is also contemplated that
the procedural steps described above may be followed by
the placement of an optional bypass conduit 113 to create a
bypass from the newly created pouch (P) directly to the
pylorus (PY) or beyond into the small intestine. Such a bypass
would channel food directly from the pouch (P) into the small
intestines to achieve a malabsorptive effect in cases where
such an effect may enhance weight loss. Such a bypass con-
duit 113 may be formed of any suitable biocompatible graft
material such as polyester or PTFE, and may be secured to
the newly created tissue ring (TR) or stoma (ST) endoscopically
using a clip or stent like structure at the anchored end to
produce an interference fit within the stoma. Alternatively,
the bypass conduit could be placed over the acquisition device
of the present invention, and secured by the same fastening
elements, and at the same time as the formation of the stoma.
In doing so, the end of the bypass graft to be anchored may be
placed over the tissue acquisition device such that the end
of the graft coincided with the tissue acquisition device opening
112, allowing it to be acquired into the device and fastened
along with the surrounding tissue. Similarly, the bypass con-
duit may be anchored in the pylorus (PY) or intestine by similar
methods, or may just be left unanchored in the intes-
tine to allow for movement due to peristalsis of the intestinal
difficulties.

FIGS. 6A-6D depict variations of the tissue rings and
pouches created using the method, and variations thereof,
located herein. FIGS. 6A and 6B depict the results of uti-
lizing the procedure described above, showing a complete
circumferential ring, in this variation, created just distal
from the where the esophagus (ES) and the stomach join each
other. FIG. 6B shows a cross section of the stomach and
organ (TR) and further depicts the resulting tissue folds
acquired by the device 105 and the fixation elements 115
deployed to fix the acquired tissue. This cross section further
depicts a cut zone or abraded zone 116 as described above.
FIGS. 6C and 6D depict another variation in which fixation of
the acquired tissue in a position centered between the lesser
curve of the stomach (LC) and the greater curve (GC) in such
a manner that multiple lumens 117, 118 result as shown in
FIG. 6D. Although only two additional lumens 117, 118 are
shown in this variation, a number of lumens may be created in
other variations depending upon the number of times and
positions the tissue is affixed.

One method of the present invention is to use the
device 105, or a variation thereof, to modify or otherwise
assist in other procedures that utilize stomach or organ plica-
tion such as those described in co-pending U.S. patent appli-
cation Ser. No. 10/188,547 earlier incorporated herein by
reference, which describes, in part, in further detail methods
and devices for stapling regions of the stomach in a linear
fashion. In cases where a zone of the stomach is linearly
stapled, the device 105 may be employed to create circular
stomata at either end of the linear staple line so as to enhance
the efficacy of a volume reduction procedure or to enhance
durability of the staple line. It may also be advantageous to
place semi-circular or partially circumferential fixation zones
at various locations within the target hollow organ. The
devices and methods described herein are particularly well-
suited for this because of their ability to “gather” the tissue
and create a circumferential restriction that acts to limit the
flow of matter, such as food, through the organ.

Devices

FIG. 7 depicts a cross-sectioned view of one varia-
tion of tissue acquisition device 120. As shown, device 120
has a main body portion 123 which has a proximal end, a
distal end, and a main lumen 121 defined therethrough.
Device 120 also has a grip portion 122 and an opposing
handle portion 122 which may be pivotally attached to main
body portion 123 such that handle portion 122 is angularly
positionable relative to grip portion 122. Main body portion
123 may further define one or more circumferentially defined
lumens along its length such that these lumens terminate at
the distal end of body portion 123 at outer distal portion 124.
Main body portion 123 further houses main body inner
portion 125, which may be an elongate tubular member config-
ured to be slidable positioned within main body lumen (121
defined through the length of main body portion 123. At the
distal end of inner portion 125, an inner body distal portion
126 may be attached thereto. This distal portion 126 may be
integrated formed onto inner portion 125 or attached separ-
ately and may be used as a clamping member to facilitate the
mechanical retention of tissue invaginated into the device
120. Distal portion 126 may also function as an anvil for re-
configuring fastening members inserted into the tissue, as
further described below. The proximal end of inner portion
125 may terminate proximally of main body portion 123 in a
fluid port 127, which may be utilized for fluid connection to,
e.g., a vacuum pump (not shown). Alternatively, distal portion
126 may function as the staple housing and outer distal portion
124 may function as the opposing anvil. In this variation,
the fasteners, as positioned within distal portion 126, may be
deployed through inner face 128 into the tissue using an
actuation device, as known in the art.

Inner body distal portion 126 may further comprises
an inner face 128 which may define an anvil or fastener
element detent 129. Where inner portion 125 joins with distal
portion 126, one or more distal ports 132 may be defined
which are in fluid communication through inner portion
125 with fluid port 127. To actuate device 120, handle portion 122
may be urged to pivot relative to grip portion 122. Slider pins
130 may be fixedly attached to main body inner portion 125
and configured to extend perpendicularly relative to inner
portion 125, as shown in FIG. 7B. Pins 130 may be opera-
tively connected with handle 122 such that rotation of move-
ment of handle 122 is translated into the linear motion of inner
portion 125. Pins 130 may be positioned within slot 131
which are defined longitudinally within main body portion
123. Slots 131 may be configured to allow limited transla-
tional movement of pins 130 thereby limiting the overall
translational distance traveled by inner portion 125.

Actuation of handle 122 in a first direction may urge
pins 130 to slide within slots 131 a first direction, e.g., distally,
thereby moving inner portion 125 distally, and actuation of
handle 122 in a second direction may urge pins 130 to slide in
a second direction, e.g., proximally, thereby moving inner
portion 125 proximally. Main body inner portion 125 may be
acted to linearly move inner body distal portion 126 relative
to outer distal portion 124 to a desired distance between the
two. When the two portions 124, 126 are moved into
position to one another, a circumferential tissue acquisition
chamber or space 130 may be created about or defined
between the outer surface of inner portion 125, inner distal
portion 126, and outer distal portion 124. Space 200 may be in fluid communication with distal port 132 and/or optionally through main body lumen 121. In operation, a vacuum force may be applied through distal port 132 and/or main body lumen 121 to invaginate or draw tissue into space 200 such that the tissue is held or configured to then receive at least one fastening element to affix the tissue configuration.

Main body portion 123 may further house driver element 133 within circumferentially-shaped fastener lumen 134. Driver element 133 may be a tubularly shaped member which is configured to traverse longitudinally within fastener lumen 134. Disposed distally of driver element 133 within fastener lumen 134 are fasteners 135 and fastener pusher mechanism 136. Fasteners 135 may comprise any variety of staples or mechanical fasteners which are made from a bio-compatible material, e.g., stainless steel, platinum, titanium, etc., and fastener retention mechanism 136 may also comprise any variety of staple retainer which is configured to hold fasteners 135 within fastener lumen 134 until fasteners 135 have been pushed or urged out of the lumen 134 and into the tissue. The proximal end of driver element 133 abuts driver actuator 137 in handle portion 122. Handle portion 122 may define a threaded cavity 138 at its proximal end which is configured to correspondingly receive and is in operative communication with driver actuator 137, which may also define a threaded insertion surface for mating with threaded cavity 138. In operation, upon tissue acquisition within circumferential space 200 and approximation of main body inner distal portion 126 and main body outer distal portion 124, driver actuator 137 may be rotated in a first direction so as to engage the threads of handle portion threaded cavity 138 and thereby engage the proximal end of driver element 133 to cause driver element 133 to move distally. As driver element 133 is advanced longitudinally in a corresponding manner as driver actuator 137 is rotated, the distal end of driver element 133 may contact fastener pusher mechanism 136 and actuating fastener 135 to distally advance and deploy fastener 135 into any acquired tissue.

Main body portion 123 may be bendable as depicted in FIG. 7C. As shown, the device 201 may be seen in one configuration in which main body portion 123 may be configured in an infinite number of different configurations for negotiating pathways within a body. This particular variation 201 shows handle grip 202 having an opposing actuation handle 203 for actuating movement of inner body distal portion 126. Also shown is an optional scope lumen 204 in the handle 202 which may be used for visualizing the tissue region being treated during deployment or retrieval. The flexibility of the main body portion 123 may be imparted, in part, by the use of, e.g., linking multiple rings 211, as shown in the isometric view in FIG. 7D. A portion 210 of the main body 123 is shown with the covering, control mechanisms, etc., omitted for clarity. Although this variation shows the use of stacked multiple rings, other variations may also be used as known in the art for flexible and/or articulatable elongate devices, e.g., endoscopes, etc. A plurality of individual rings 211 may be aligned with one another to create a length of the main body portion 123. Any number of rings 211 may be used depending upon the overall desired length of the device or the desired length of a flexible portion of the device. Each of the rings 211 may have at least one main channel or lumen 212, which when individual rings 211 are aligned as a whole, create a main channel throughout the length of the device. Each of the rings 213 may also have a number of spacers or protrusions 213 defined on or around the circumference of the device for creating pivotable sections for facilitating relative motion between adjacent rings 211, as shown in the art. Although the rings 211 are shown with two oppositely positioned protrusions 213, any number of protrusions 213 may be used as practicable depending upon the degree of relative motion desired between adjacent rings 211. Alternatively, device main body 123 may be constructed in part, e.g., a coil spring, to achieve a similar functional result. Coil springs may be made of superelastic materials, e.g., nitinol, or spring steels made, e.g., from stainless steels. The main body 123 or main body segments may be constructed of various bio-compatible materials, such as stainless steel, Delrin or other engineering thermoplastics, etc.

FIG. 7E depicts a single ring 211 having the main lumen 212 defined therethrough. Main lumen 212 may be modified and enlarged to provide a channel having a large enough diameter to receive a conventional endoscope for possible use with the present device. One example of such a device may have a lumen diameter of, e.g., 10 mm, with an outer diameter of, e.g., 18 mm. One or more of such lumens may be created within the annular section 211 to enable linkage of each section 211 to another by one or several cables or flexible wires (not shown) adapted to be positioned through the lumens. These wires or cables may be routed through the length of the device and fixed at the proximal end of the main body portion 123.

As shown in FIG. 7E, an optional sheath or thin film 221 may be placed over the device or at least along a portion 210 of the device to encapsulate the linkages and create a smooth shaft surface, while still maintaining its flexibility. The sheath or thin film 221 may be made of a variety of bio-compatible materials, e.g., heatshrink polymers, plastics, etc.

FIG. 8 depicts another variation on the distal end of a tissue acquisition device. The inner distal portion 230 is shown defining an inner face 231 and device outer distal portion 232 having an inner face 233. The inner distal portion inner face 231 and the outer distal portion inner face 233 may be formed to face one another in apposition and both faces 231, 233 may each be formed at an angle (A) relative to a longitudinal axis of the device main body 123. The angle (A) may range anywhere from 0-90 degrees, but is preferably in the range of 15-45 degrees, depending on the desired angle of the resulting tissue fixation zone. This variation may be used to allow the operator to position the tissue acquisition device perpendicularly to a surface of the organ to be treated (for ease of use) while acquiring and fixing the tissue at an angle relative to the tissue surface. In doing so, the operator may fashion the resulting fixation zone to more closely approximate a curvature of the organ, such as the curvature between the GEJ and the LC of the stomach. FIG. 9 depicts a further variation 240 of the tissue acquisition device in which fenestrations or ports 241 may be defined over the surface of the device inner distal portion 242. Additional fenestrations or ports 243 may be defined over a portion of the device outer distal portion 232, and additional fenestrations or ports 244 may also be defined over a surface of body inner portion 245. These additional ports may allow this variation 240 to acquire tissue along a length of the distal end of the tissue acquisition device 240 at multiple locations therealong. In practice, this method of tissue acquisition may allow the operator some freedom to manipulate the acquired tissue by the relative movement of device inner distal portion 242 and the device
outer distal portion 232. This technique can also assist in positioning the tissue to be fixed, and/or assuring that the required amount of tissue (e.g. some muscular layers of the organ wall), have been uniformly acquired prior to fixation.

Following fixation, the tissue acquisition device of the present invention is withdrawn from the organ. In doing so, care should be used not to over-dilate or stretch the newly created tissue ring or stoma. To mitigate any dilation or stretching, the inner distal portion may also be modified. FIGS. 10A and 10B depict variations 250, 260 of the tissue acquisition inner distal portion that are adaptable to effectively reduce in cross sectional area to allow for easier removal of the tissue acquisition device from the organ once the circumferential fixation zone has been created. FIG. 10A depicts tissue acquisition device inner distal end 251 which is pivotally mounted on main body inner portion 253 about pin 252. Activation of the pivoting action may be controlled by release of an interface between pin 252 and a stay (not shown) housed within main body inner portion to activate rotation of inner distal end 251, e.g., in a direction 256. The inner distal end 251 may be rotated by any angle such the inner face 254 is angled or parallel relative to the longitudinal axis 255 of the device.

FIG. 10B depicts another variation 260 on tissue acquisition inner portion distal end which may have a segmented configuration. In this variation 260, the inner portion distal end may be made of a plurality of individual segments 262 which when collapsed, reduces the diameter of inner portion distal end to facilitate removal. Thus, during tissue acquisition and/or fixation, the expanded inner distal portion 264 may be utilized and after the procedure, it may then be compressed radially 265 about a pivot 263 to reduce the cross-sectional profile for removal from the area.

Additional Bypass Conduit Devices

As mentioned above for FIG. 5E, an optional bypass conduit 113 may be placed within the stomach cavity (SC) at the site of the narrowing or stoma (ST). Such a conduit may be placed not only in conjunction with the intragastric staple line described herein, but with various other conventional procedures to create a bypass from the pouch (P) directly to the pylorus (PY), or beyond into the small intestines to effect the rate at which food is metabolized. It may also further enhance the efficacy of a bariatric procedure by facilitating “dumping syndrome”. One variation on the bypass conduit is seen in FIG. 11A in bypass conduit assembly 270. In this variation, assembly 270 comprises a conduit wall 272, which may be tubular in shape. Bypass lumen 274 may be defined throughout the length of conduit wall 272. The conduit wall 272 may extend between a proximal end 271 and a distal end 273 and may be made from a variety of biocompatible materials. For instance, conduit wall 272 may be made from a rubber material or from a polymeric material which may be configured to be lubricious, e.g., Teflon, Nylon, Dacron, PTFE, polyethylene, polyurethane, polyethylene terephthalate, etc.

To further increase the structural resiliency of the conduit wall 272, an optional reinforcing member 278 may be utilized within the structure. Reinforcing member 278 may include any number of structural enhancements such as a coil member as shown in the FIG. 11A. The coil member may be wound in a helical manner along the body of conduit wall 272. Another variation may have wires positioned longitudinally along conduit wall 272 rather than a coiled member. Alternatively, a wire-framed structure may be utilized along the conduit wall 272.

In any of these structural enhancements, the reinforcing member 278 may be disposed in a laminate structure between layers of conduit wall 272 material. Alternatively, the reinforcing member 278 may be formed integrally into the conduit wall 272 by forming the conduit material about the member 278. Another variation may have reinforcing member 278 adhered onto the outer and/or inner surface of the conduit wall 272 through the use of adhesives, sutures, clamps, or any other number of conventional attachment methods. Moreover, these optional structural enhancements may be utilized not only in the variation shown in FIG. 11A, but in any of the other variations described herein depending upon the desired structural characteristics.

The proximal end 271 may be affixed or secured to the stomach tissue within stomach cavity (SC), to the tissue adjacent to pouch (P), or to other tissue, as further described below. In the present variation, conduit assembly 270 may have a first gasket 275 and a second gasket 276 positioned distal of the first gasket 275 along wall 272. Gaskets 275, 276 may be made of a rubberized material or a polymeric material configured to be flexible during the deployment of assembly 270. Such a gasketed assembly 270 may be used in conjunction with the tissue ring (TR) or stoma (ST), as described in detail above. Upon deployment and positioning of assembly 270 within the stomach cavity (SC), these gaskets 275, 276 may be allowed to expand such that first gasket 275 is located proximally of the stoma (ST) and second gasket 276 is located distally of the stoma (ST). A portion of the conduit wall 277 located inbetween the gaskets 275, 276 may be in contact with the stoma (ST) and may be sufficiently flexible to form around the stoma (ST).

When the bypass conduit is properly positioned to extend from the narrowing, e.g., the stoma, to within the intestinal tract, e.g., to the jejunum or further, the distal end of the liner may be positioned to extend distally of the duodenal ampulla 323. As explained in further detail below, the duodenal ampulla is a duct which connects the common bile duct and the pancreatic duct to the duodenum for discharging digestive fluids into the duodenum. These fluids (alimentary flow) normally intern mix with partially digested food from the stomach cavity (SC). To facilitate such fluid exchange and to prevent the duct from being blocked by the liner, the liner may include communications to the inside of the liner, such as fenestrations, or channels alongside the liner wherein the cross section of the liner may be varied to allow such an exchange.

FIGS. 11B-11E show variations of cross sections of the bypass conduit from FIG. 11A which may allow for fluid exchange to occur along the outer surface of the liner. FIG. 11B shows one variation in which the conduit wall 272 defines one or more longitudinal channels 279 along the outer surface of the wall 272. FIGS. 11C and 11D show variations in which the conduit walls 272”, 272”, respectively, are angled such that the contact between the outer surface and the tissue is non-continuous, thereby allowing fluids to seep within or along these spaces or channels created between the angled outer surface and the tissue. FIG. 11E shows yet another variation in which the conduit wall 272” defines an undulating outer surface forming at least one or more longitudinal channels 279. These examples of possible irregularly defined cross sections are merely illustrative and are not
intended to be limited only to these examples. Other variations, as should known to those in the art, are intended to be included herewith.

As shown in FIG. 11F, these irregular cross sectional areas may extend along the entire length of the conduit wall 272 or just partially along the conduit wall 272, as shown. The length of the irregular cross section may extend just from within the body organ to distal of the body organ, or along any desired length of the conduit wall 272, depending upon the desired results.

Another variation on the bypass conduit is shown in FIG. 12 in conduit assembly 280. This variation is similar to that shown in FIG. 11A but with the addition of a fluid bypass conduit 281 located adjacent to conduit wall 272. The fluid conduit 281 has a proximal end 282 for positioning within the stomach cavity (SC) and a distal end 283 for positioning within the intestines distal to the stomach cavity (SC), as described in further detail below. Fluid conduit 281 may be made in a variety of ways; for instance, conduit 281 may be manufactured separately from conduit wall 272 and attached to the outer surface of the conduit wall 272 using any variety of methods, e.g., adhesives, clamping, etc., in which case conduit 281 may be made from a similar or same material as conduit wall 272. For instance, fluid conduit 281 may be made of a braided material, as described above, to inhibit kinking of the conduit. Alternatively, conduit 281 may be formed integrally with the conduit wall 272 as a uniform assembly.

In either case, conduit 281 has a length which is typically co-extensive with the length of the main conduit wall 272 but may be less than or greater than the length of the main conduit. The conduit 281 may also be configured such that the conduit 281 doesn’t block the alimentary flow from the ducts. The distal end 283 of the conduit 281 may thus terminate proximally or distally of the distal end 273 of the conduit wall 272, or it may optionally terminate at or distally of the distal end 273, depending upon the desired structure and use. Although a single fluid conduit 281 is shown in the figure, any number of additional fluid conduits may be incorporated into the assembly. These additional fluid conduits may be aligned in parallel with conduit 281 or positioned about the circumference of the conduit wall 272. Moreover, the additional conduits may be made of various lengths depending upon the desired results. Furthermore, although fluid conduit 281 is shown as being parallel with main conduit wall 272, fluid conduit 281 may be positioned about conduit wall 272 in a helical or spiral manner, or it may be positioned in a variety of ways, e.g., such as a bent or hooked proximal end, etc.

Yet another variation is shown in FIGS. 13A and 13B, which show conduit assembly variation 290. This variation incorporates a fluid bypass conduit 291 which is coaxially positioned about a portion of conduit wall 272. Fluid conduit 291 has a proximal end 292 for positioning within the stomach cavity (SC) and a distal end 294 for positioning distally of the stomach cavity (SC). To maintain the coaxially adjacent lumen 295, support struts 293 may be positioned between fluid conduit 291 and conduit wall 272, as seen in FIGS. 13A and 13B, which is a cross-sectional view taken from FIG. 13A. Support struts 293 may be positioned circumferentially between fluid conduit 291 and conduit wall 272 in a variety of configurations so long as coaxial lumen 295 is substantially unobstructed. Support struts 293 may be fabricated separately or integrally with conduit wall 272 and/or fluid conduit 291. Alternatively, struts 293 may be extensions of a laminated wireframe making up the tubular structure for conduit wall 272 and/or fluid conduit 291.

Conduit wall 272 and/or any of its auxiliary fluid conduits may be directly fabricated from various materials, as described above. Alternatively, they may be fabricated from an underlying braided tubular structure such as that shown in bypass conduit variation 300, as seen in FIG. 14. The walls of the conduit may be made of a braided material to form a braided tubular structure 304 defining a bypass lumen 302. The braided structure 304 may be made to make the assembly 300 more resistant to kinking, as is generally known in the art. The tubular structure 304 may be made, for instance, from superelastic materials like Nickel-Titanium alloys (nitinol) or from a metal such as stainless steel. Such construction may allow for the tubular structure 304 to be bent and twisted 308 in an infinite manner so as to allow the structure 304 to flex and move with the stomach without kinking or obstructing flow through the conduit. The braided structure 304 may be coated, covered, or laminated with a biocompatible material to aid in its lubricity; any variety of materials may be used, e.g., polymeric materials such as Teflon, Nylon, Dacron, PTFE, polyethylene, polystyrene, polyurethane, polyethylene terephthalate, etc.

To aid in the secure placement of the bypass conduit proximal to or within the stomach cavity (SC), the proximal end 301 of the conduit 304 may optionally be radially flared 305 such that the flared portion 305 securely contacts the tissue. The flared portion 305 may optionally be reinforced, either by additional braiding or an additional structural ring or band, to create a reinforced region 307 for further ensuring adequate structural support. Moreover, the distal end 303 may also be optionally flared 306 to assist in anchoring the distal end of the bypass conduit within the intestinal tract or distal to the stomach cavity (SC).

To further facilitate anchoring of a bypass conduit, a number of alternative anchors may be utilized aside from the gasketed configuration described above. Another variation is shown in FIG. 15A in conduit anchoring variation 310. As seen, conduit wall 311 may have a first gasket 312 and an optional second gasket 313 in which each gasket 312, 313 may comprise a coil which is biased to extend radially outward. As above, first and second gaskets 312, 313 may be separated by a conduit portion 314 and a partial length or the entire length of the conduit wall 311 may be reinforced with a reinforcing member 319, as described above.

Another alternative variation to facilitate the anchoring of the bypass conduit may be seen in variation 315 in FIG. 15B. Conduit anchoring variation 315 may have a reinforced portion or section 317 located near or at the proximal did of conduit wall 316. This reinforced section 317 may comprise a radially expanding portion, much like a self-expanding stent made of a shape memory alloy such as nitinol; alternatively, section 317 may also comprise a prosthetic ring or gasket made of a polymer material. Attachment points 318 may be optionally included to project from the proximal end of conduit wall 316 or from the reinforced section 317. These attachment points 318 may be configured to pierce into the tissue and aid in affixing the conduit 315 by helping to hold the conduit 315 securely in place along the tissue. The attachment points 318 may be positioned around the circumference of the conduit wall 316 or in any number of configurations as is known in the art. Although the figure shows attachments points 318 as hooks, any number of different
configurations may be utilized, e.g., barbs, clamps, sutures, staples, stents, bands, adhesives, etc., may also be used.

**Bypass Conduit Placement**

[0079] The bypass conduit may be positioned between the stomach cavity (SC) and the intestines in a variety of ways aside from that shown in FIG. 5E above. The bypass conduit assembly 280 may be used in conjunction with various gastric procedures. As seen in FIG. 16A, bypass conduit assembly 280 may be used with a stomach (SC) which has undergone a laparoscopic banding procedure. FIG. 16B shows a view of a lap band 321 which has been positioned around a portion of the stomach cavity (SC) below the esophagus (ES) prior to having a bypass conduit deployed. FIG. 16A shows a view of assembly 320 in which conduit assembly 280 has been positioned to extend from the stoma (ST) created by the banding, to a point past the pylorus (PY). As shown, the proximal end 271 of the conduit assembly 280 may be secured within the stoma created by the lap band 321 using any of the methods described above. The conduit wall 272 is appropriately sized such that it extends through the stomach cavity (SC) from, in this variation, the stoma (ST) into the intestines, e.g., the duodenum 322, although the distal end 273 may extend farther into the intestinal tract, e.g., to the jejunum. The distal end 273 of the conduit wall 272 may be left unanchored in the intestinal tract or it may be optionally anchored to the tissue. Anchoring of the distal end 273 may be achieved using any of the anchoring methods as described above for anchoring of the proximal end 271.

[0080] The fluid conduit 281 may be seen in this variation as being positioned along the conduit wall 272 and within the stomach cavity (SC) such that its proximal end 282 is placed within the stomach cavity (SC) at the stoma (ST) and its distal end 283 extends past the pylorus (PY) and partway into the duodenum. Although fluid conduit 281 may be sized to have a length that is shorter than the conduit wall 272, it may typically be sized to have a length which is longer than or coterminous with that of conduit wall 272, and further adapted to facilitate fluid communication between the stomach cavity (SC), or gastric remnant, and the intestines, or the duodenal ampulla 323 and the intestines. As positioned, fluid conduit 281 allows for the gastric fluids produced within the stomach cavity (SC) and the digestive fluids discharged through the duodenal ampulla (or duct) 323 to intermix and to be transported through the conduit 281 between the stomach cavity (SC) and the intestine distal of the duodenal ampulla 323. The fluid conduit 281 also allows for the fluids to intermix and for the fluids produced within the stomach cavity (SC) to drain without contacting any ingested foods transported through the bypass conduit 272. If the distal end 283 of the fluid conduit 281 extends past the duodenal ampulla 323, the region of the conduit 281 near or at the entrance to the duct 323 may define one or more fenestrations or openings 324 along its length. These fenestrations 324 may be positioned and sized appropriately such that they allow for the fluid communication between the duct 323 and the lumen of the fluid conduit 281.

[0081] FIG. 17A shows another variation 330 utilizing the lap band 321 with the coaxial fluid bypass conduit 290. FIG. 17B shows a view of the stomach prior to having the conduit assembly 290 deployed. In this variation, fluid conduit 291 may be positioned such that its proximal end is within the stomach cavity (SC) and its distal end 294 is positioned within the duodenum 322 to the jejunum, either proximally or at the duodenal ampulla 323. If the distal end 294 is positioned distally of the ampulla 323, one or more fenestrations 331 may be defined along the length of the fluid conduit 291 to facilitate the fluid exchange and to maintain the fluid communication, as described above, between the ampulla 323 and local intestine and the fluid conduit 291. The use of this coaxially adjacent conduit variation allows for the free rotation of the conduit wall 272 and/or fluid conduit 291 about its longitudinal axis within the stomach cavity (SC) without the problems of kinking or improper placement of the fluid conduit relative to the stomach cavity (SC). The proximal 271 and distal 273 ends of the conduit wall 272 may be anchored in much the same manner as described above.

[0082] In the case of a stomach which has undergone a vertical banded gastroplasty (VBG) procedure, the conduit may also be utilized to facilitate patient treatment. FIG. 18A is a view of the stomach which has had the VBG procedure prior to deployment of the bypass conduit. As shown, a vertical staple line 341 has been deployed along a portion of the stomach extending from the circular defect 342 within the stomach to the gastroesophageal junction (GEJ). A silastic band 343 has also been positioned to create a narrowing or stoma at the end of the staple line 341. As shown in the variation 340 of FIG. 18A, the bypass conduit 272 may be deployed such that its proximal end 271 is secured within the stoma (ST) created by placement of the silastic band 343 to bypass the stomach cavity (SC) and extend distally through the pylorus (PY), as described above. The conduit may thus extend from within the stomach cavity (SC) to within the intestinal tract. Moreover, one or more fenestrations 331 may be defined along certain portions of the length of the conduit wall 272 positioned at active secretory zones (such as within the stomach cavity (SC) and/or the duodenal ampulla) to allow fluid exchange through the walls of the bypass conduit 272 at the point of those anatomic structures. By spacing fenestrations 331, and limiting them to communication with only specified active zones, a single conduit construction can function both as a sufficient barrier between ingested food and the intestine (malabsorption), and a selected flow path for digestive fluids.

[0083] FIG. 19A shows another variation 350 in which a bypass conduit may be used a stomach which has undergone laparoscopic banding to constrict the stomach cavity (SC) and create a stoma. The lap band 343 may be used to constrict the stomach such that the original stomach, as indicated by the outline 351, is constricted by the band 343 to create a constricted stomach, as indicated by the constricted outline 352. The bypass conduit proximal end 271 may then be secured within the stoma created by the lap band 343, as described above. Furthermore, fenestrations 331, which may be valved, may be placed along the length of the bypass conduit 272 to allow a single conduit to perform the dual functions of malabsorption and the maintenance of digestive fluid flow. Such fenestrations may include one-way valves that open to receive fluids from outside the bypass conduit. As the valves may be configured to selectively open at regions along the conduit length where the pressure from such flow overcomes the force which maintains the valve closed; adequate pressure from the flow may be generated by the fluids such as within the gastric remnant or at the inflow of the ducts (duodenal ampulla). Such a design would not require specific alignment at flow inlets. FIG. 19C depicts one variation of a one-way valve 354 having a door or flap 355 hinged or partially secured at 356 to the inside of bypass conduit wall.
272. Flap 355 may be biased to urge the valve shut in the absence of the fluid flow. FIGS. 19D and 19E are illustrative examples which show variations on the flap 355. FIG. 19D shows a flap 355 which may be attached to the conduit wall and hinged via notched section 356 about which flap 355 may rotate. FIG. 19E shows another example in which flap 355 may be attached about a biased hinge 356. In either case, these examples are merely intended to be illustrative and other methods of flap actuation are intended to be included herein. In addition, such selective communication between bypass conduit 272 and related organs or intestine can be established by varying the porosity or permeability of certain segments 357, 358 along the length of bypass conduit wall 272, as shown in FIG. 19F. FIG. 19G shows a cross-sectional view of the bypass conduit wall 272 secured to the stomach wall 353 by the lap band 343.

[0084] Yet another variation on conduit placement may include the use of conventional devices such as those described in U.S. Pat. No. 4,458,681 (Hopkins) and in U.S. Pat. No. 4,558,699 (Bashour), which are both incorporated herein by reference in their entirety. Both patents describe variations on clamps which may be placed across a stomach (externally) to create a stoma therewithin for the passage of food through the stomach. The clamps may be placed over the stomach, e.g., through conventional laparoscopic procedures, and a bypass conduit may be placed endoscopically within the stomach such that the proximal end of the conduit is supported by the clamp within the created stoma using any of the methods described above.

[0088] The steps of performing the method of organ division or reduction (transoral stomach reduction) are used to illustrate in detail the method and devices of the present invention, however the present invention is not limited thereby. Use of these steps and the tools deployed therein may be varied to achieve a similar result in other hollow body organs and it is anticipated that such techniques can be employed to divide or restrict other hollow body organs such as organs of the gastrointestinal tract such as bowel, stomach or intestine, or in procedures in the bladder (treatment for incontinence by reinforcing the bladder sphincter) or uterus, etc. In addition, as previously mentioned, other procedures such as the treatment of GERD may also benefit from the methods and devices disclosed herein. While certain embodiments have been illustrated and described in detail, those having ordinary skill in the art will appreciate that various alternatives, modifications, and equivalents may be used and that the invention is not intended to be limited to the specifics of these variations.

We claim:
1. A surgical method, comprising:
   - transorally advancing a proximal end of a tubular member through an esophagus of a patient to position the proximal end of the tubular member adjacent an attachment site near a gastroesophageal junction;
   - transorally advancing a distal end of the tubular member through the esophagus and through a stomach of the patient to position the distal end adjacent an intestine of the patient; and
   - attaching the proximal end of the tubular member at the attachment site such that the tubular member is configured to channel food from the esophagus directly into the intestine by allowing the food to pass from the esophagus into an inner lumen of the tubular member through the proximal end of the tubular member and exit the inner lumen of the tubular member into the intestine through the distal end of the tubular member.

2. The method of claim 1, further comprising anchoring the distal end of the tubular member to the intestine.
3. The method of claim 1, further comprising anchoring the distal end of the tubular member in a pylorus of the patient.
4. The method of claim 1, further comprising implanting the tubular member within the patient without anchoring the distal end of the tubular member to tissue such that the distal end of the tubular member is allowed to freely move within the patient.
5. The method of claim 1, wherein attaching the proximal end of the tubular member at the attachment site comprises attaching the proximal end of the tubular member distal of the esophagus.
6. The method of claim 1, further comprising transorally advancing an endoscopic device through the esophagus, wherein transorally advancing the proximal and distal ends of the tubular member through the esophagus comprises advancing the tubular member over the endoscopic device with the endoscopic device passing through the inner lumen of the tubular member.

7. The method of claim 1, wherein attaching the proximal end of the tubular member at the attachment site comprises attaching the proximal end of the tubular member at the attachment site using at least one fastening element.

8. The method of claim 1, wherein attaching the proximal end of the tubular member at the attachment site comprises attaching the proximal end of the tubular member at the attachment site using at least one fastening element without creating a tissue fold.

9. The method of claim 1, further comprising forming a tissue fold at the attachment site, wherein attaching the proximal end of the tubular member at the attachment site comprises attaching the proximal end of the tubular member to the tissue fold.

10. The method of claim 9, wherein the tissue fold is formed at the attachment site simultaneously with attachment of the proximal end of the tubular member at the attachment site.

11. The method of claim 9, wherein the proximal end of the tubular member is attached at the attachment site after the tissue fold is formed.

12. The method of claim 9, further comprising transorally advancing an endoscopic device through the esophagus, wherein the tissue fold is formed using the endoscopic device, and transorally advancing the proximal and distal ends of the tubular member through the esophagus comprises advancing the tubular member over the endoscopic device with the endoscopic device passing through the inner lumen of the tubular member.

13. The method of claim 9, wherein forming the tissue fold comprises forming a tissue ring from tissue adjacent the gastroesophageal junction, and attaching the proximal end of the tubular member at the attachment site comprises attaching the proximal end of the tubular member to a proximal side of the tissue ring such that the tubular member passes through an inner opening defined by a perimeter of the tissue ring.

14. The method of claim 9, wherein forming the tissue fold comprises forming a circumferential tissue fold around a partial circumference of the esophagus from tissue adjacent the gastroesophageal junction, and attaching the proximal end of the tubular member at the attachment site comprises attaching the proximal end of the tubular member to the tissue fold.

15. The method of claim 1, wherein the tubular member is formed from a flexible material that allows the tubular member to move within the intestine due to peristalsis of a wall of the intestine.

16. The method of claim 1, wherein the inner lumen of the tubular member is free of obstructions such that the food is allowed to freely pass through the inner lumen of the tubular member.

17. A surgical method, comprising: transorally advancing into a patient a conduit having an unobstructed inner lumen extending between proximal and distal ends thereof; positioning the proximal end of the conduit adjacent a gastroesophageal junction of the patient; positioning the distal end of the conduit within an intestine of the patient such that an intermediate portion of the conduit extending between the proximal and distal ends thereof is positioned within the stomach; and attaching the proximal end of the conduit adjacent the gastroesophageal junction using at least one fastening element such that food can pass directly from the esophagus into the inner lumen of the conduit through the proximal end of the conduit, advance through the inner lumen of the conduit, and exit the inner lumen through the distal end of the conduit to pass directly into the intestine.