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(19) **United States**(12) **Patent Application Publication****Deem et al.**(10) **Pub. No.: US 2007/0213748 A1**(43) **Pub. Date: Sep. 13, 2007**(54) **OBESITY TREATMENT TOOLS AND METHODS****Publication Classification**(51) **Int. Cl.****A61B 17/00** (2006.01)(52) **U.S. Cl.** **606/151**

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Correspondence Address:

FULWIDER PATTON LLP**HOWARD HUGHES CENTER****6060 CENTER DRIVE, TENTH FLOOR****LOS ANGELES, CA 90045 (US)**

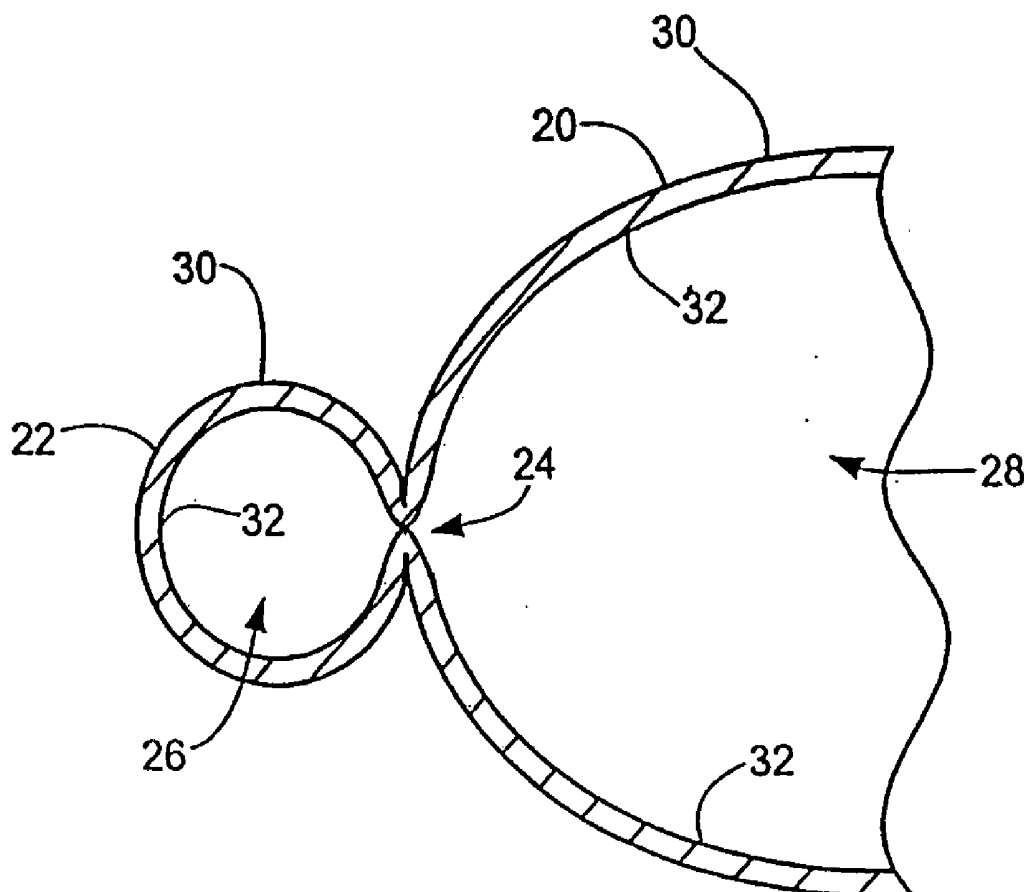
(57)

ABSTRACT

Various obesity treatment tools and methods are described herein, as well as treatments for other gastric-related diseases, e.g., GERD. Treatment includes reducing the size of the stomach pouch to limit the caloric intake as well as to provide an earlier feeling of satiety. This may be done by creating a smaller gastric pouch within the stomach directly from the interior of the stomach itself. The smaller pouches may be made through the use of individual anchoring devices, rotating probes, or volume reduction devices. A pyloroplasty procedure may also be performed to render the pyloric sphincter incompetent. A gastric bypass procedure may additionally be performed using atraumatic magnetic anastomoses devices so that sugars and fats are passed directly to the bowel while bypassing the stomach. Many of these procedures may be done in a variety of combinations. Treatment may create enforced behavioral modifications by discouraging the ingestion of high-caloric foods

(21) Appl. No.: **11/801,450**(22) Filed: **May 10, 2007****Related U.S. Application Data**

(63) Continuation of application No. 10/402,061, filed on Mar. 27, 2003, which is a continuation of application No. 09/871,297, filed on May 30, 2001, now Pat. No. 6,558,400.



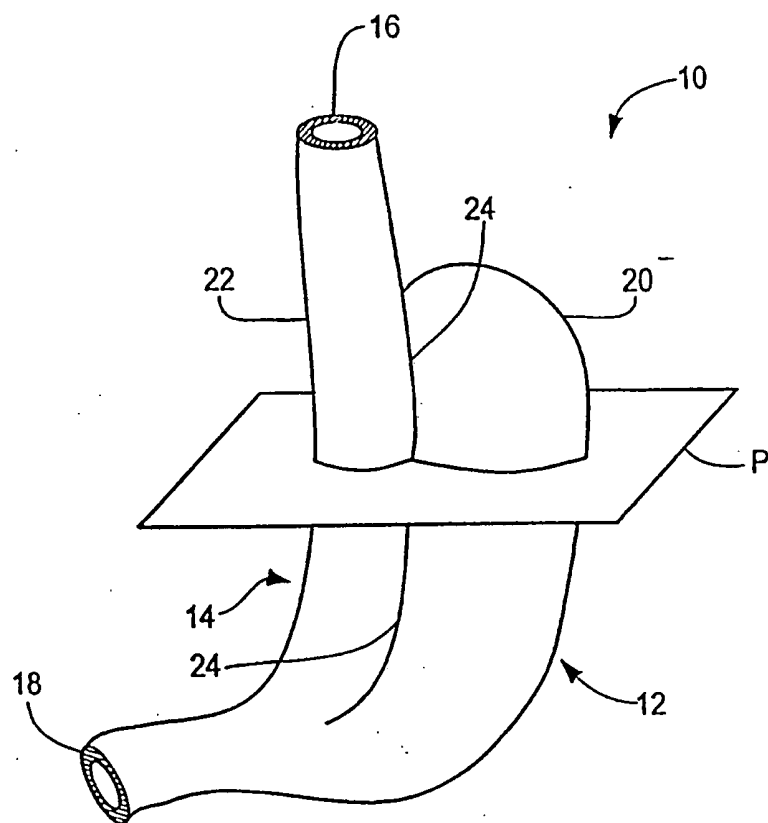


FIG. 1A

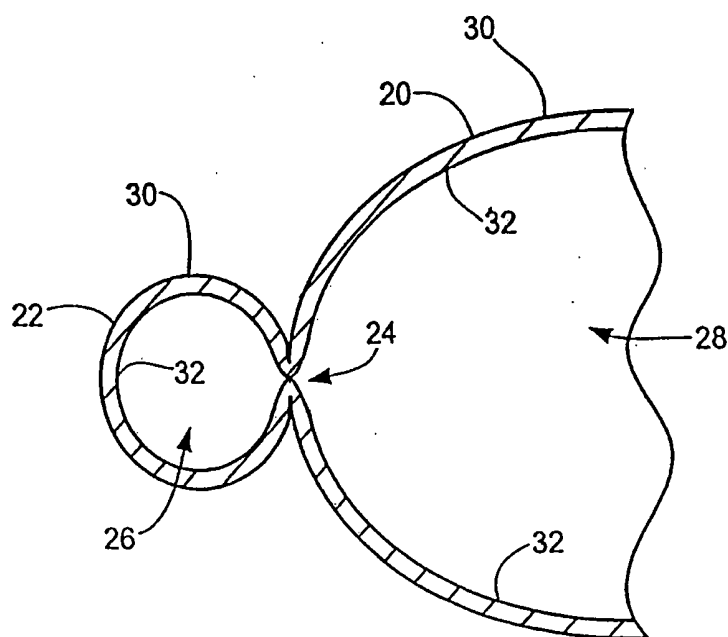


FIG. 1B

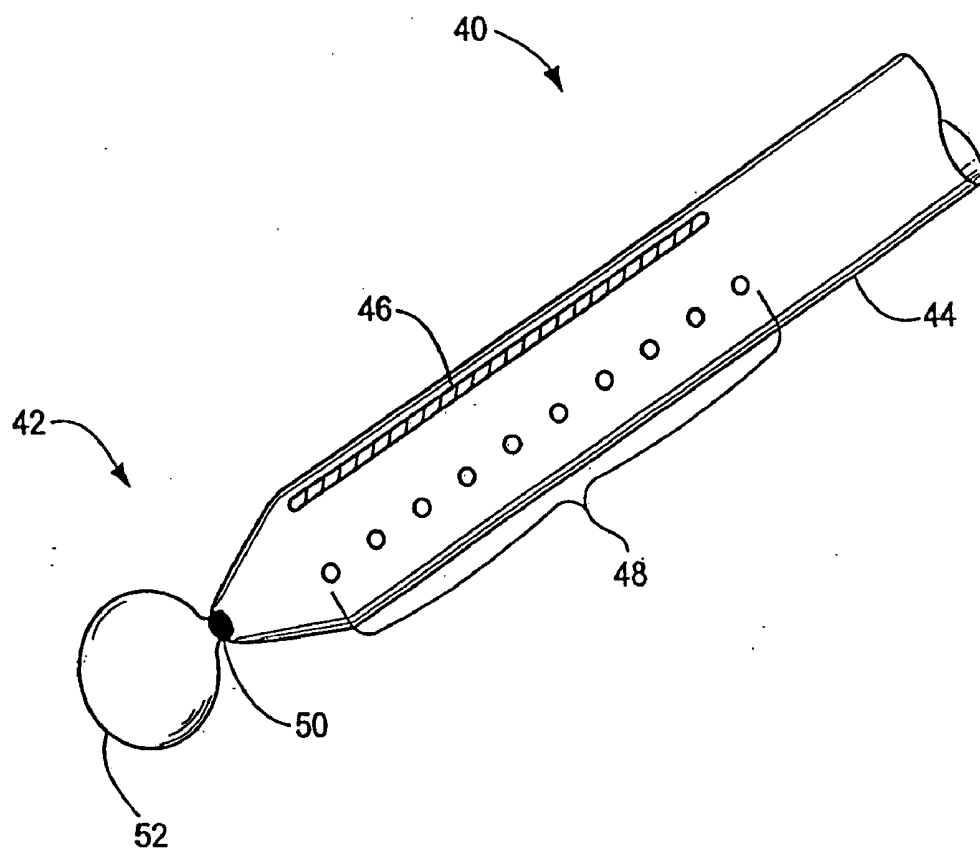


FIG. 2

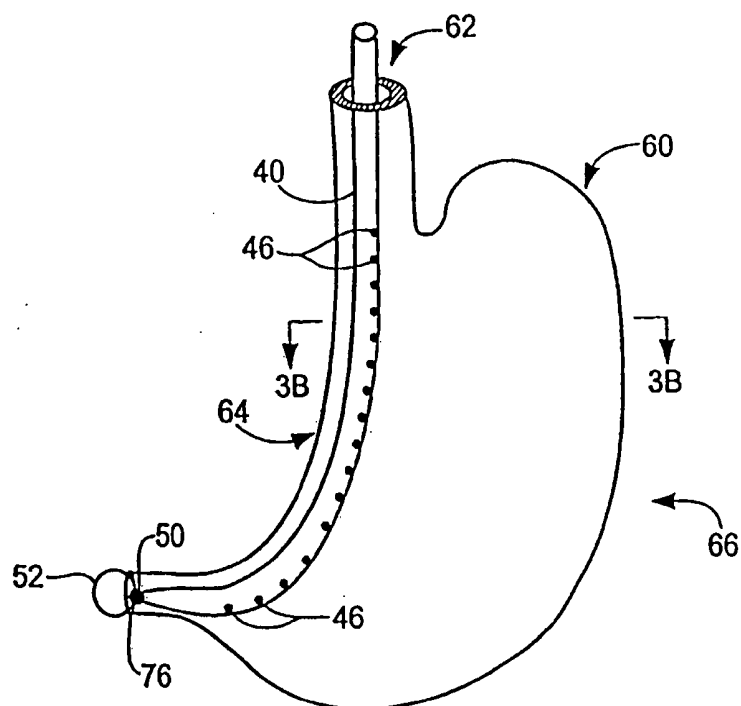


FIG. 3A

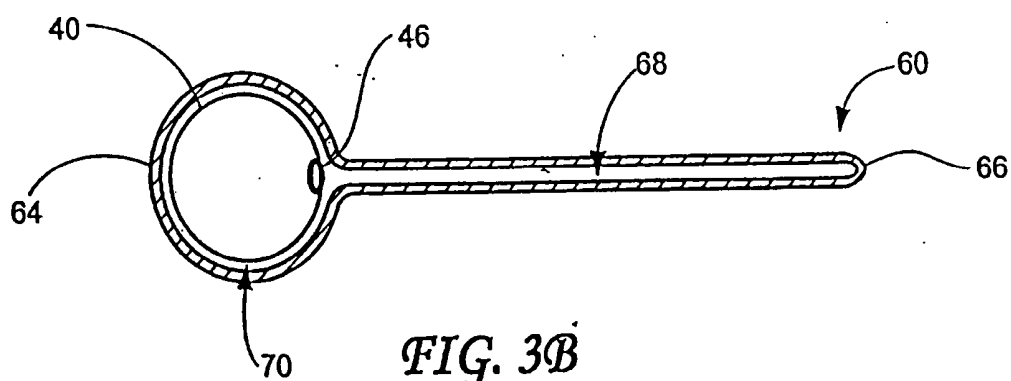


FIG. 3B

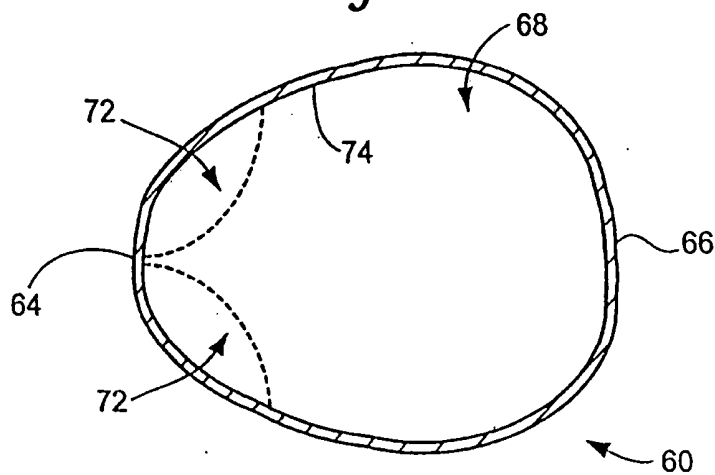


FIG. 3C

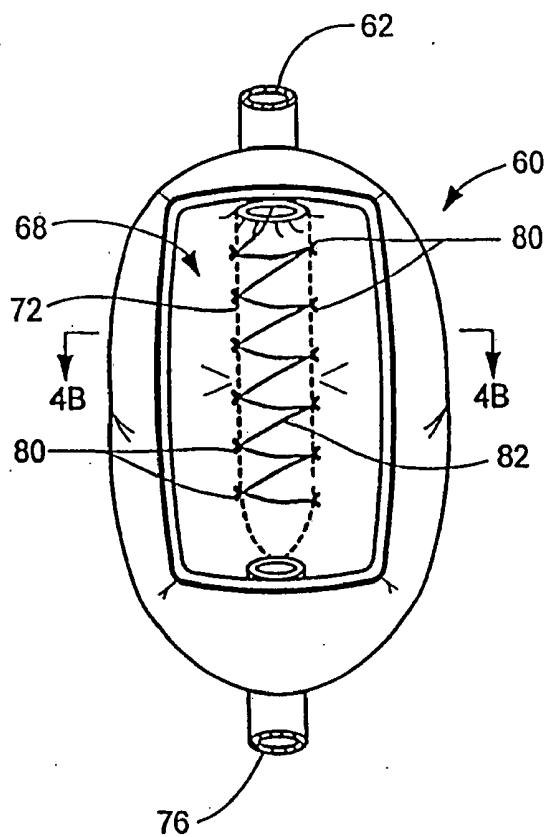


FIG. 4A

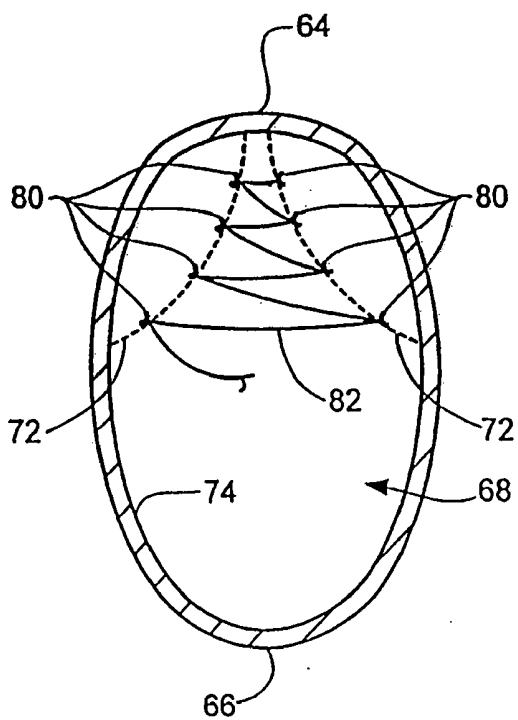


FIG. 4B

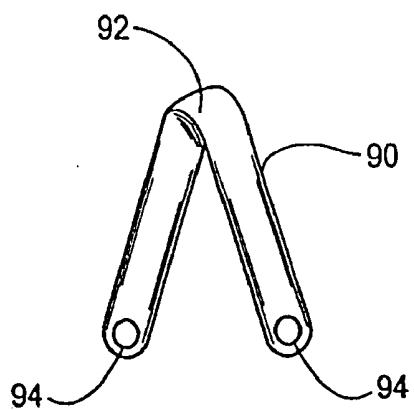


FIG. 5A

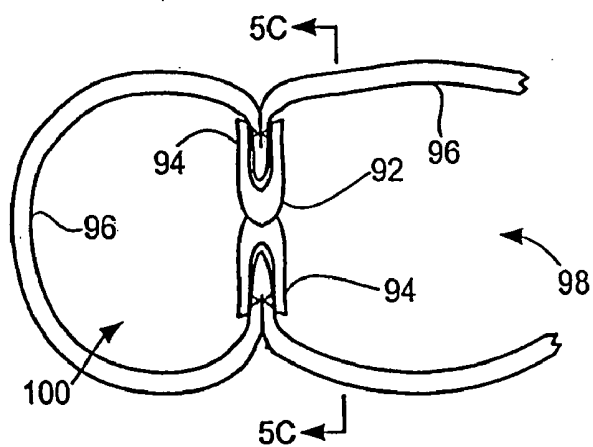


FIG. 5B

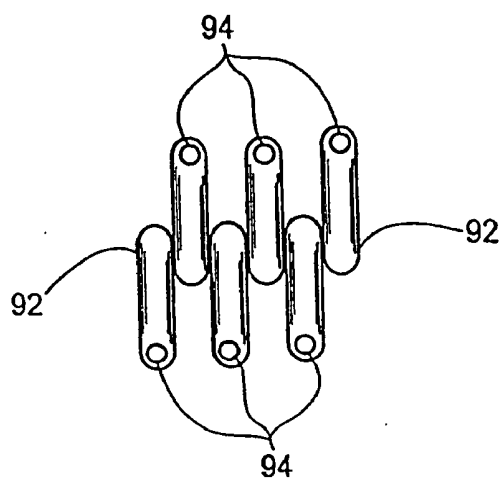


FIG. 5C

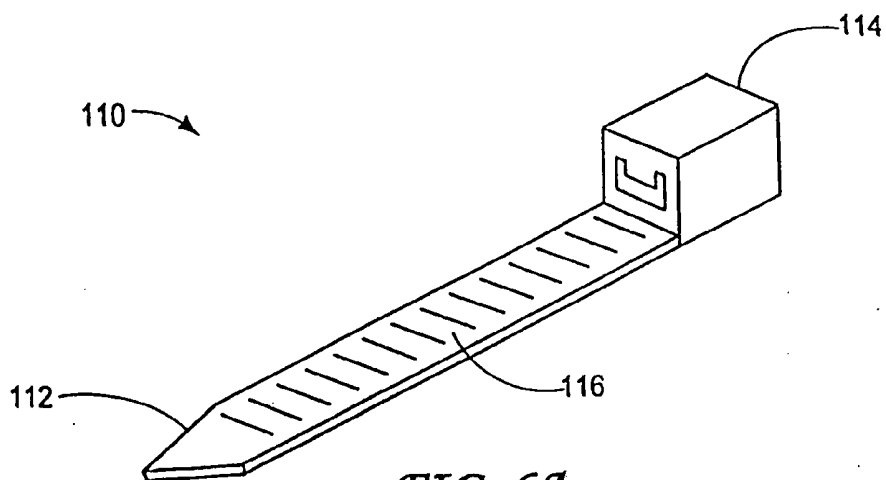


FIG. 6A

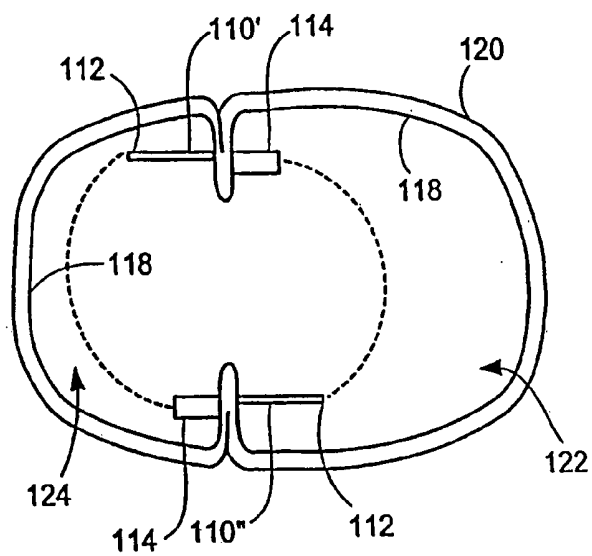


FIG. 6B

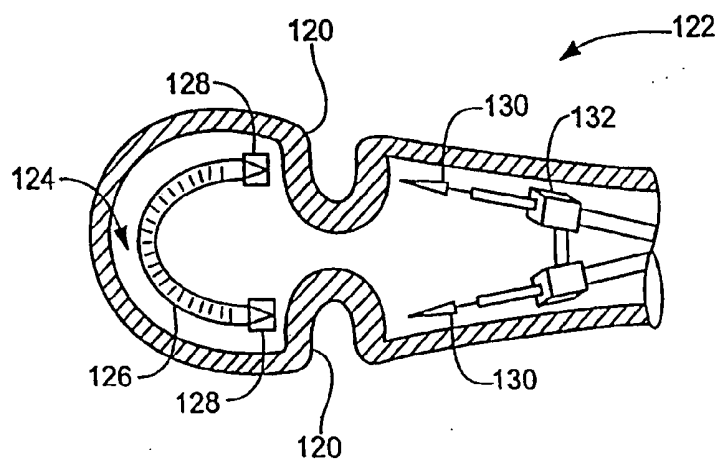


FIG. 6C

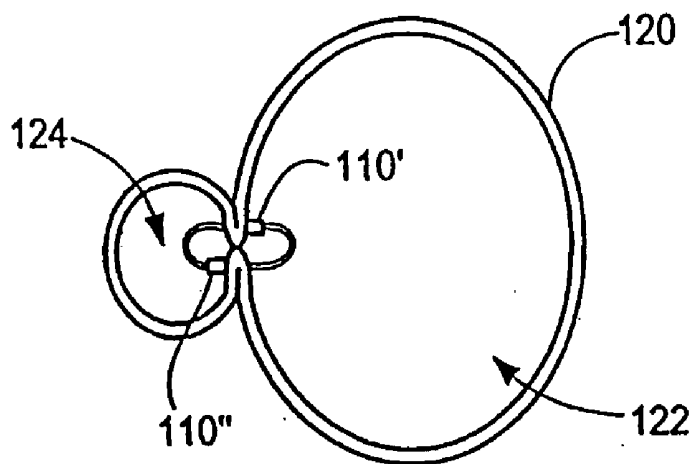


FIG. 6D

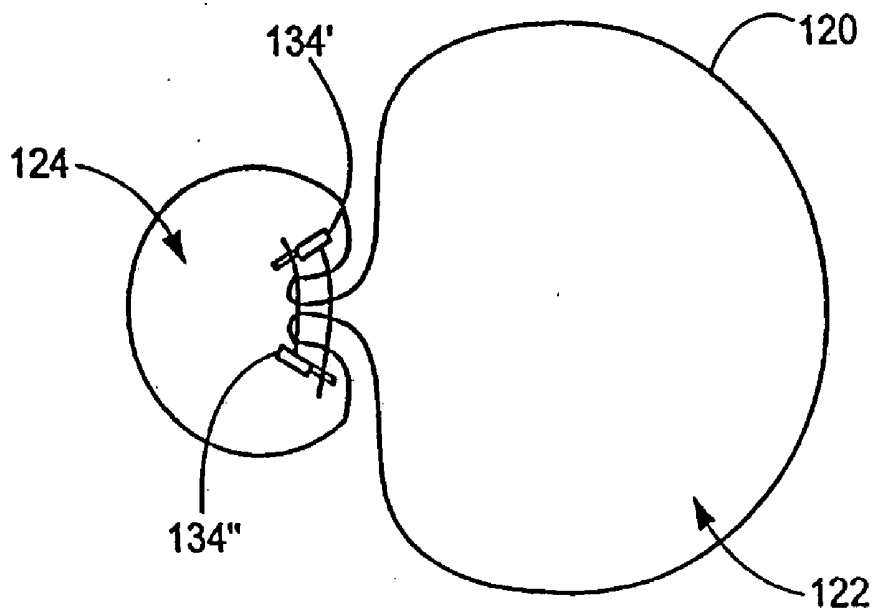


FIG. 6E

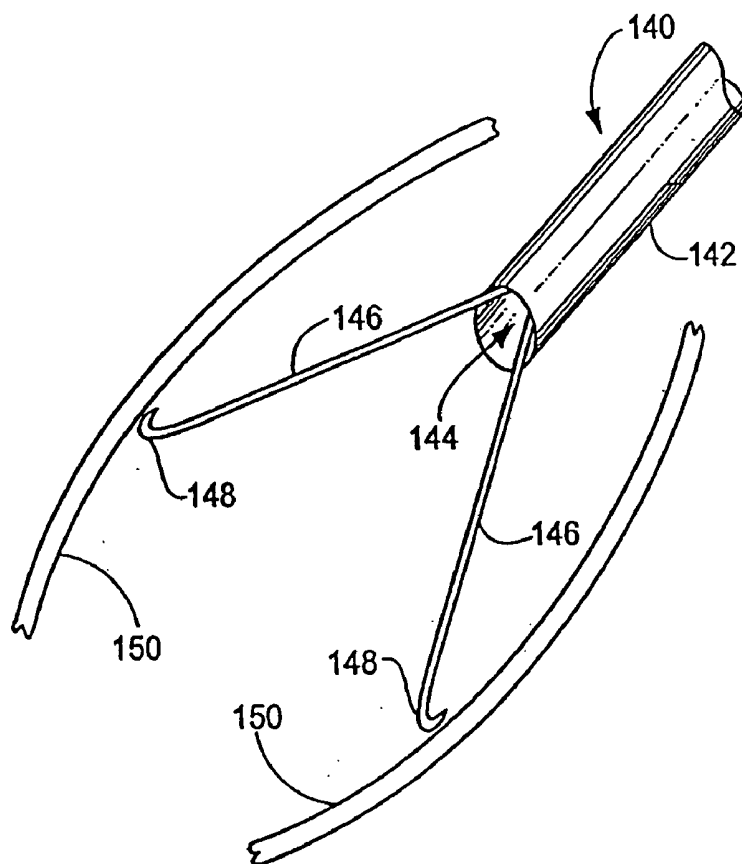


FIG. 7A

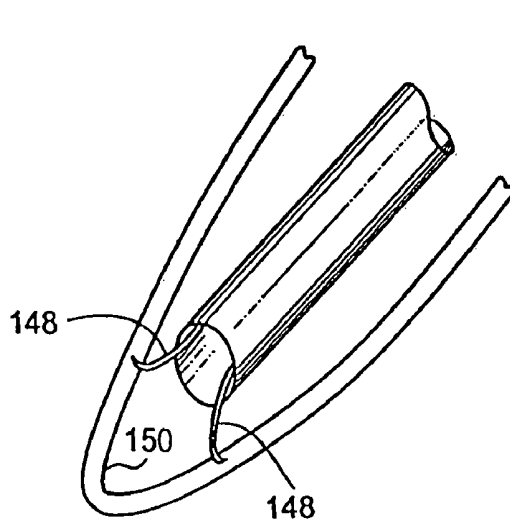


FIG. 7B

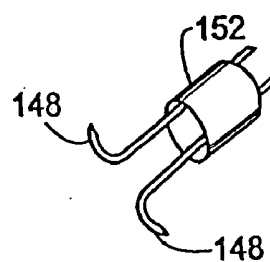


FIG. 7C

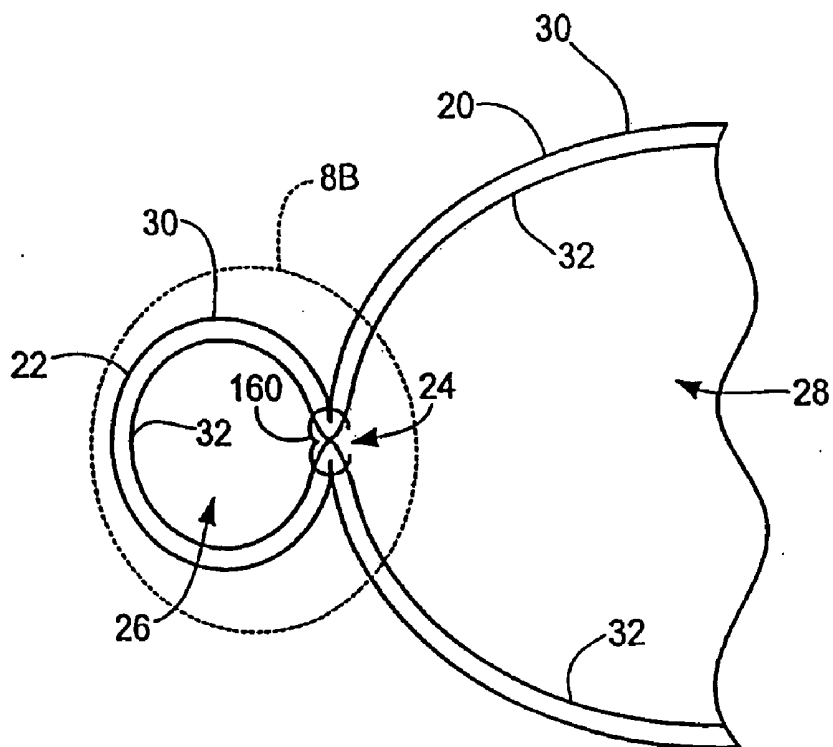


FIG. 8A

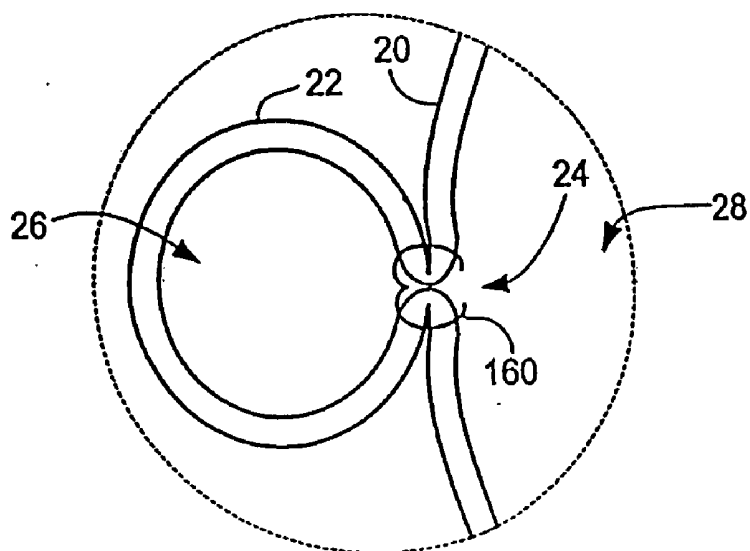


FIG. 8B

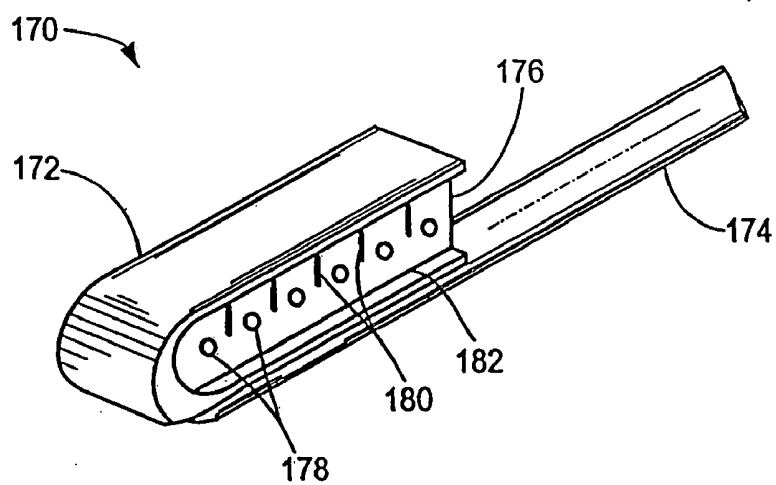


FIG. 9A

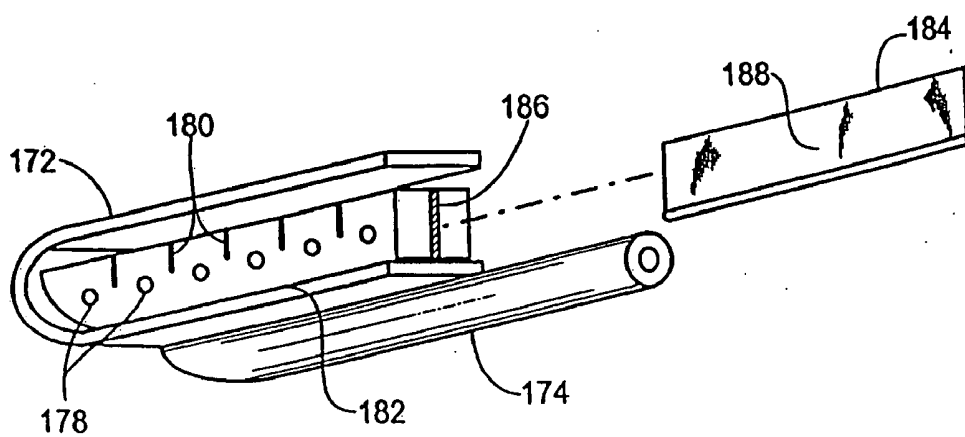


FIG. 9B

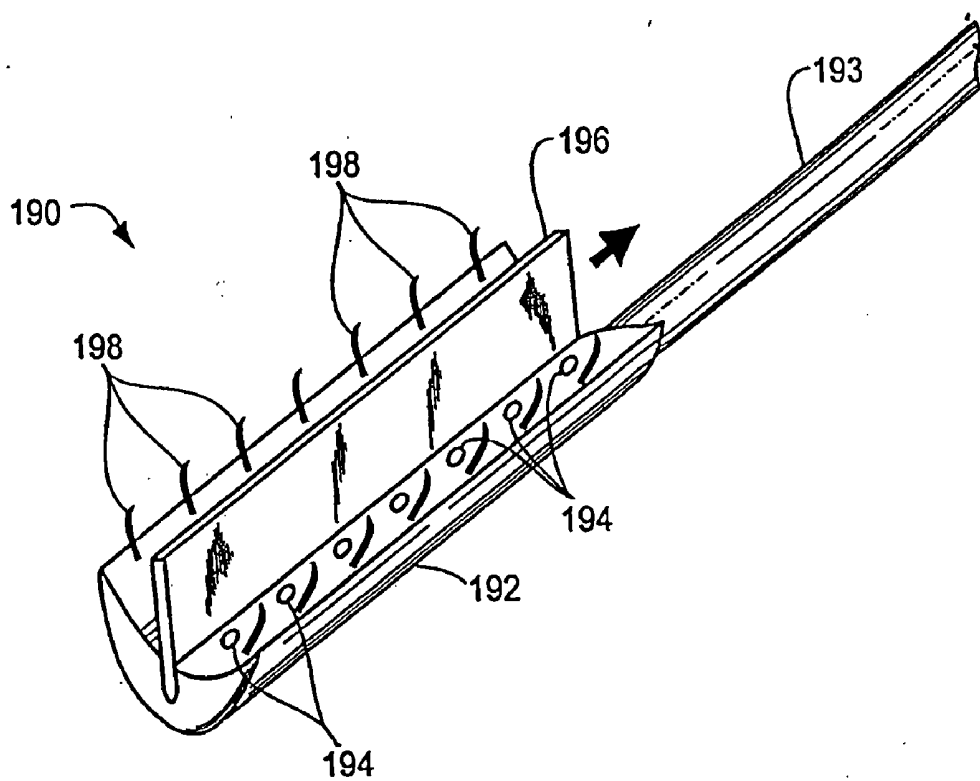


FIG. 10

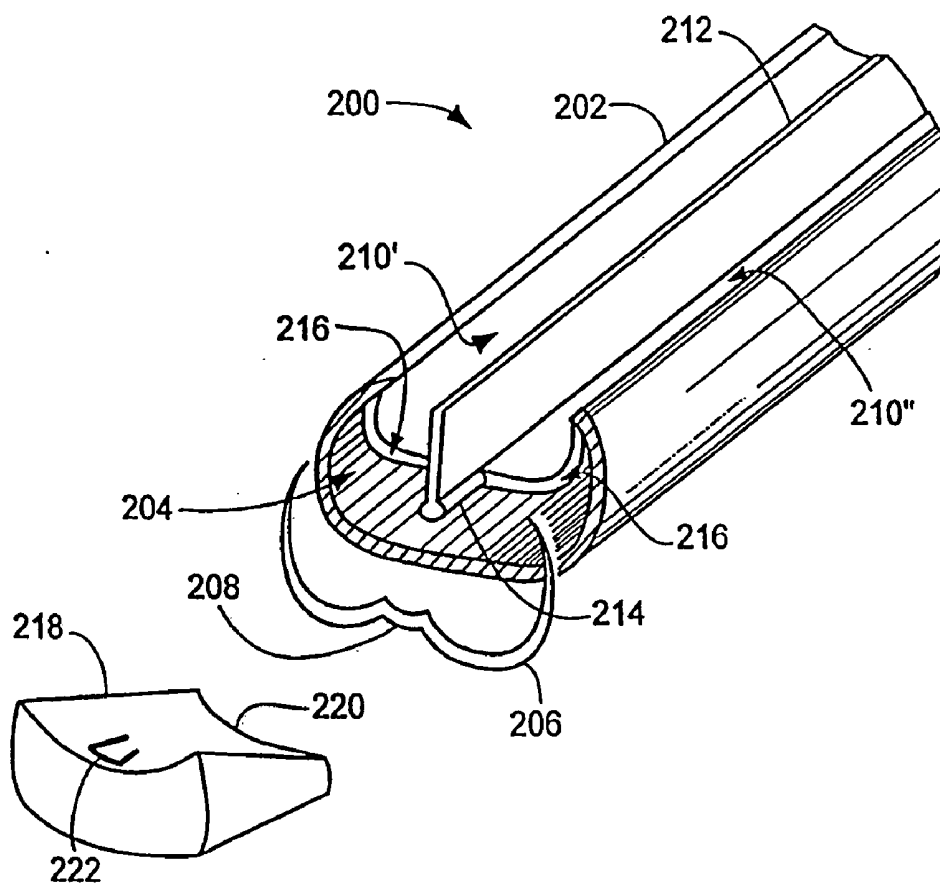


FIG. 11A

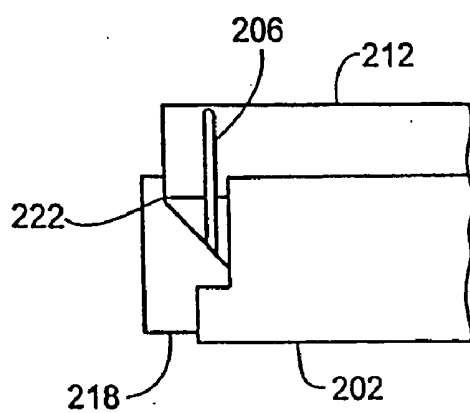


FIG. 11B

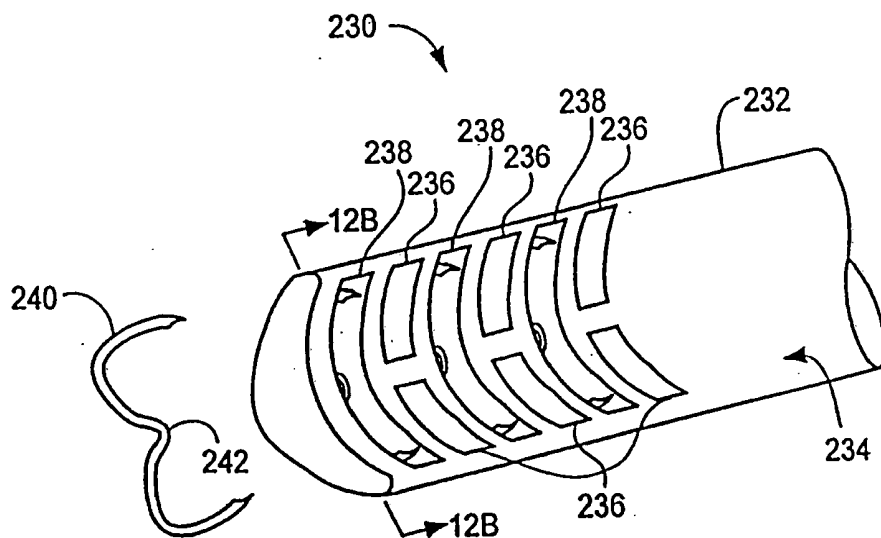


FIG. 12A

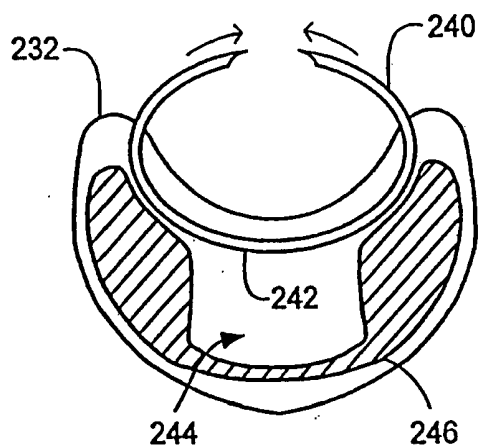


FIG. 12B

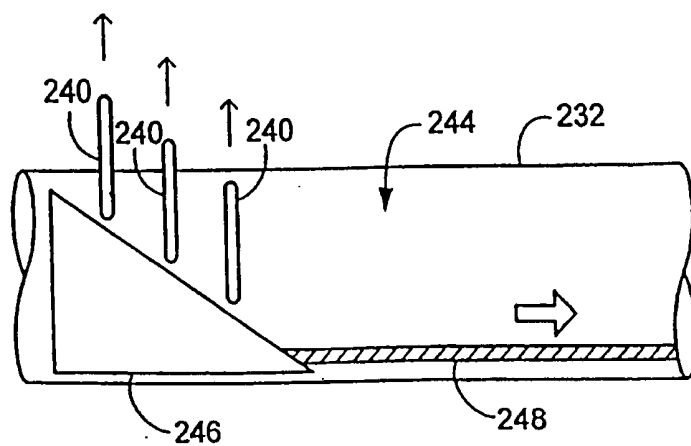


FIG. 12C

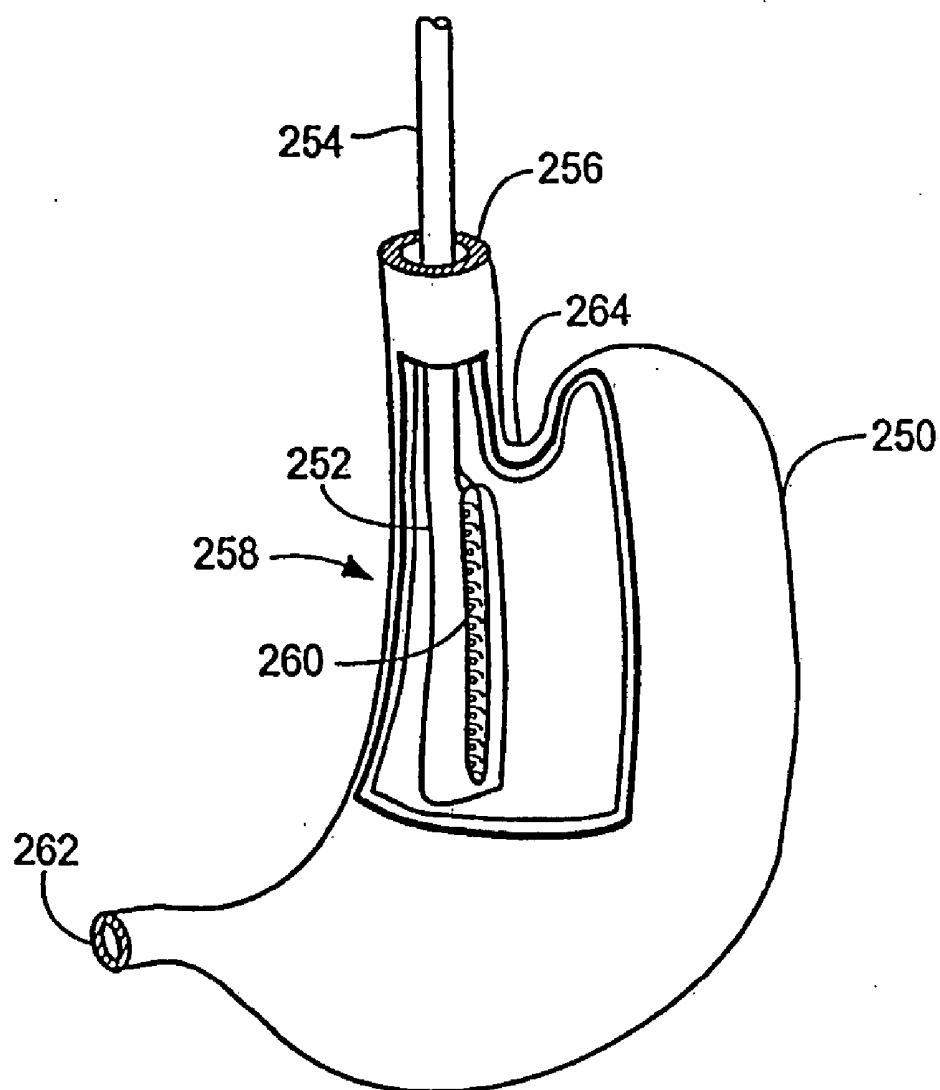


FIG. 13

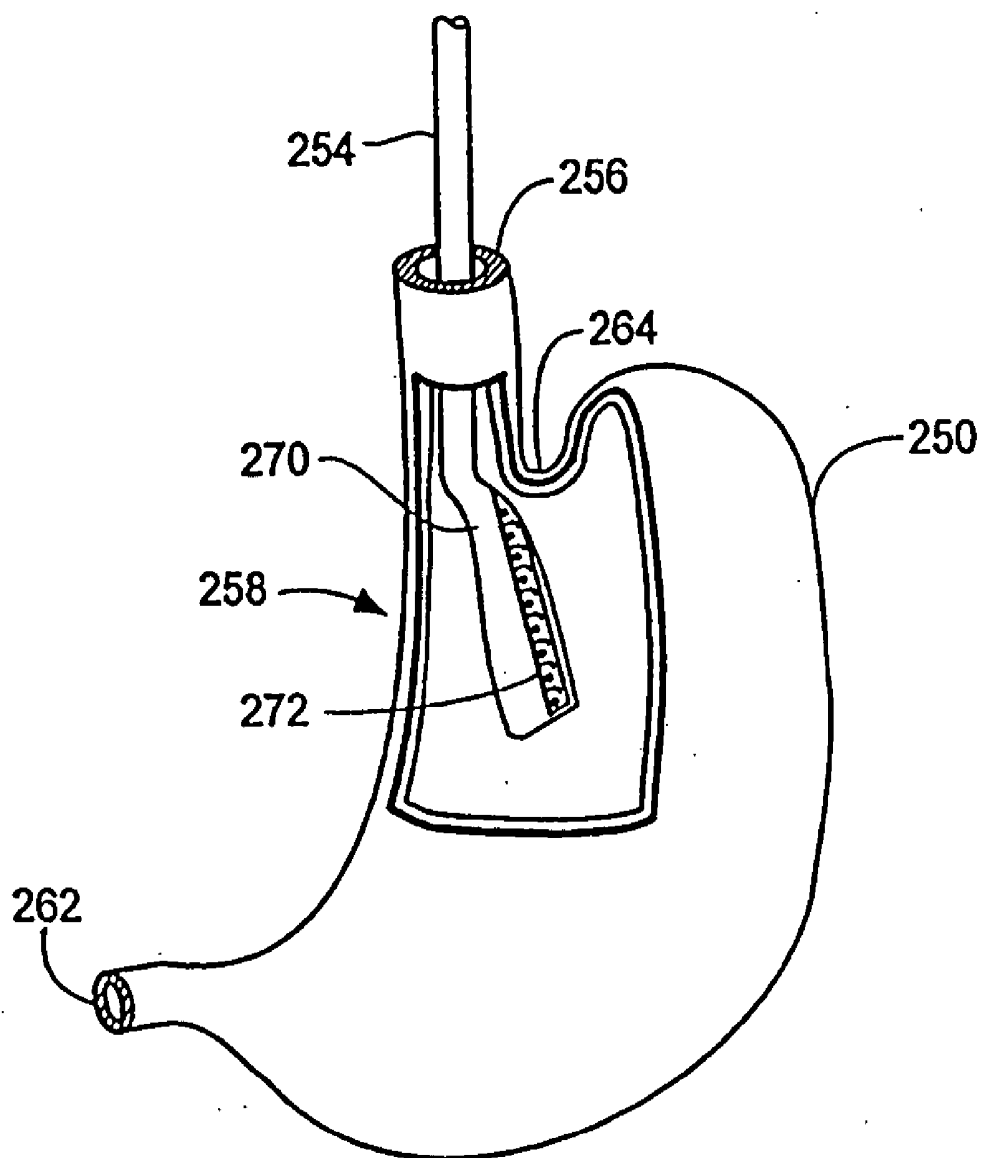
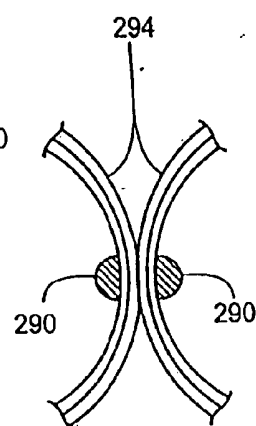
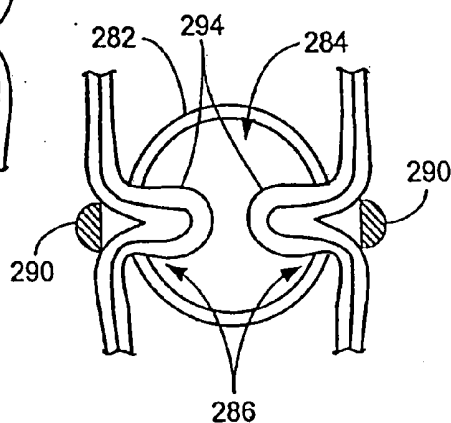
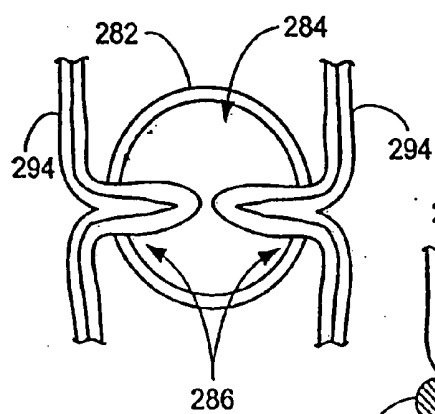
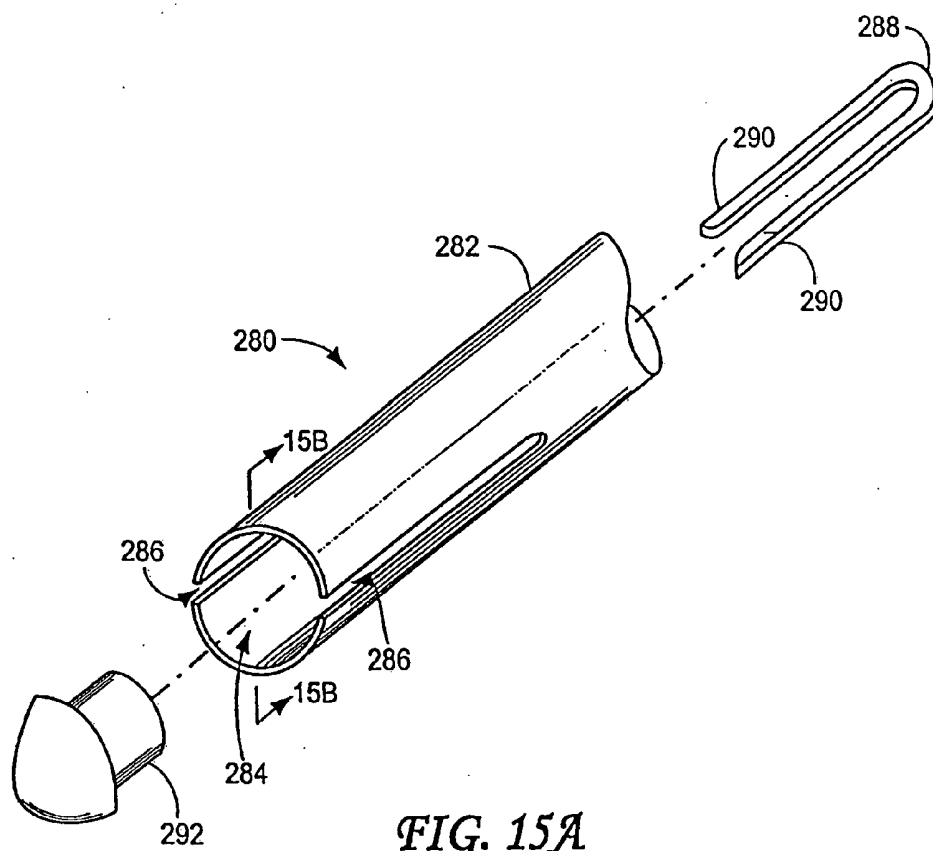
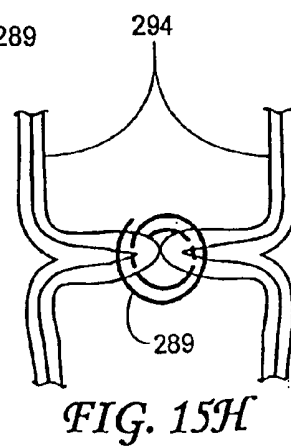
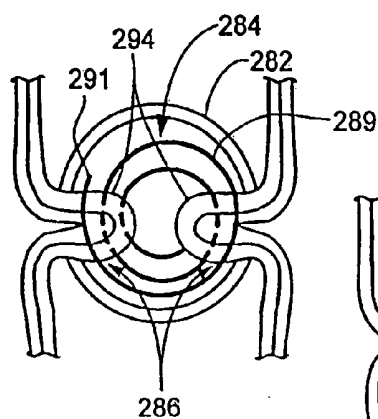
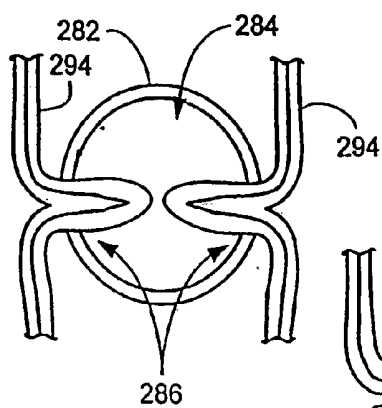
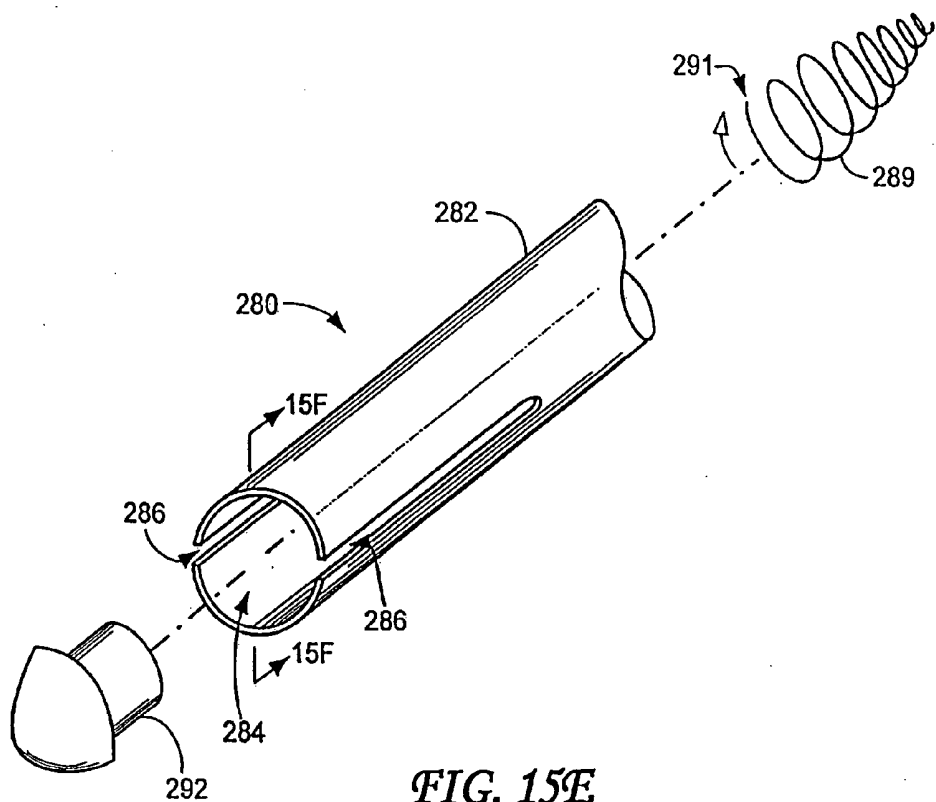


FIG. 14





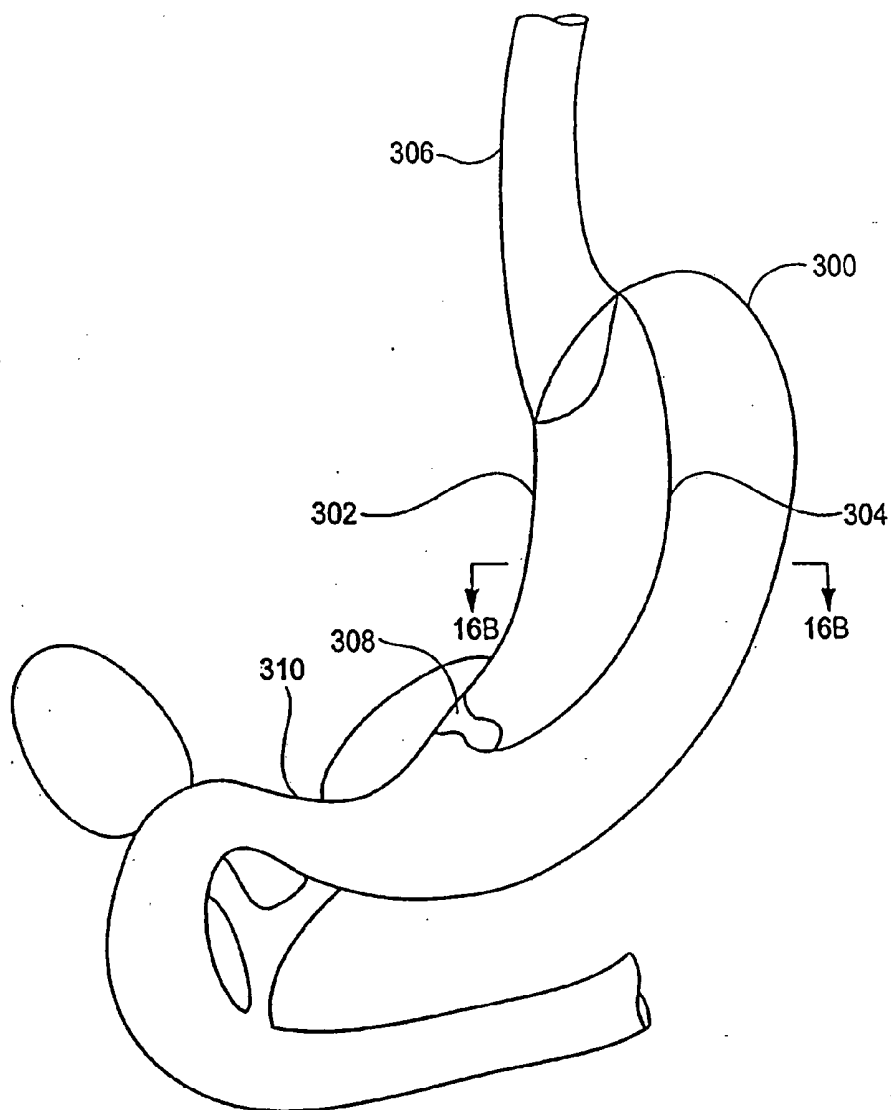


FIG. 16A

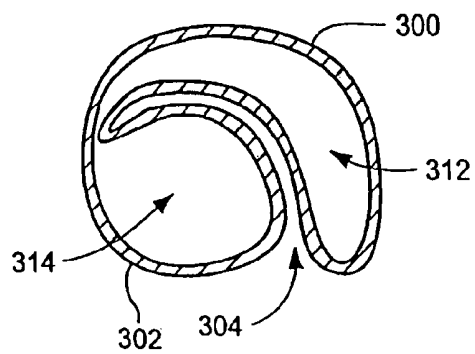


FIG. 16B

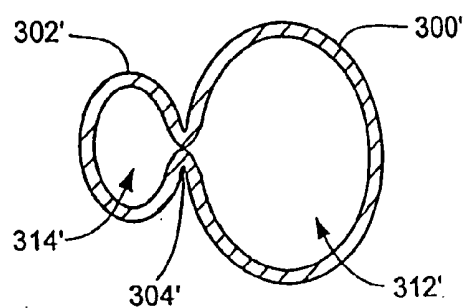


FIG. 16C

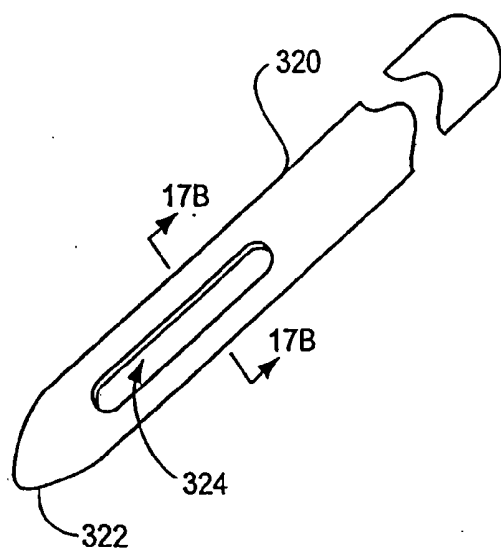


FIG. 17A

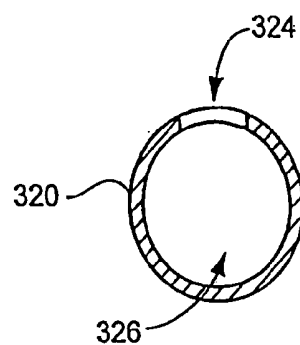


FIG. 17B

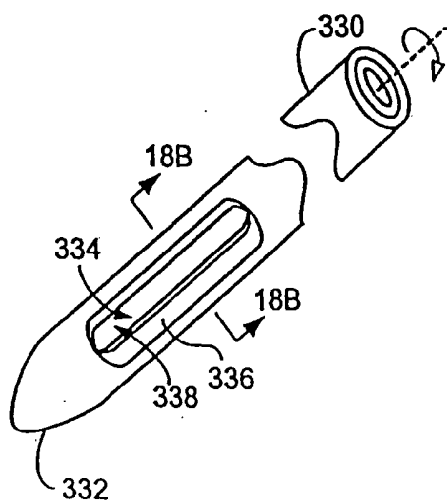


FIG. 18A

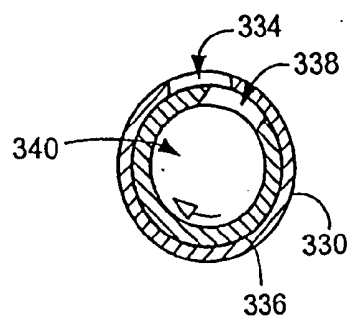


FIG. 18B

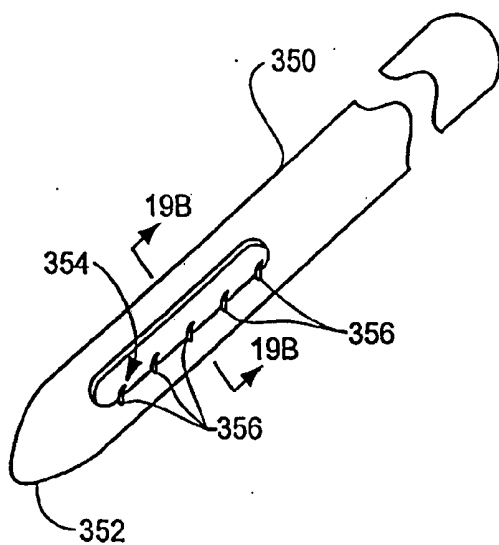


FIG. 19A

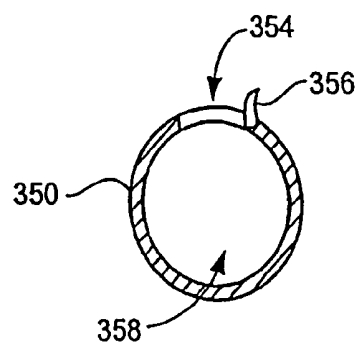


FIG. 19B

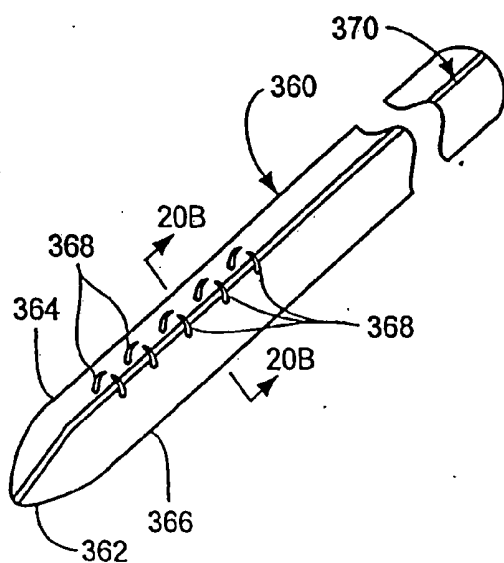


FIG. 20A

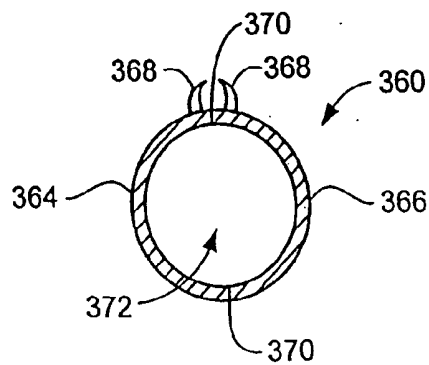


FIG. 20B

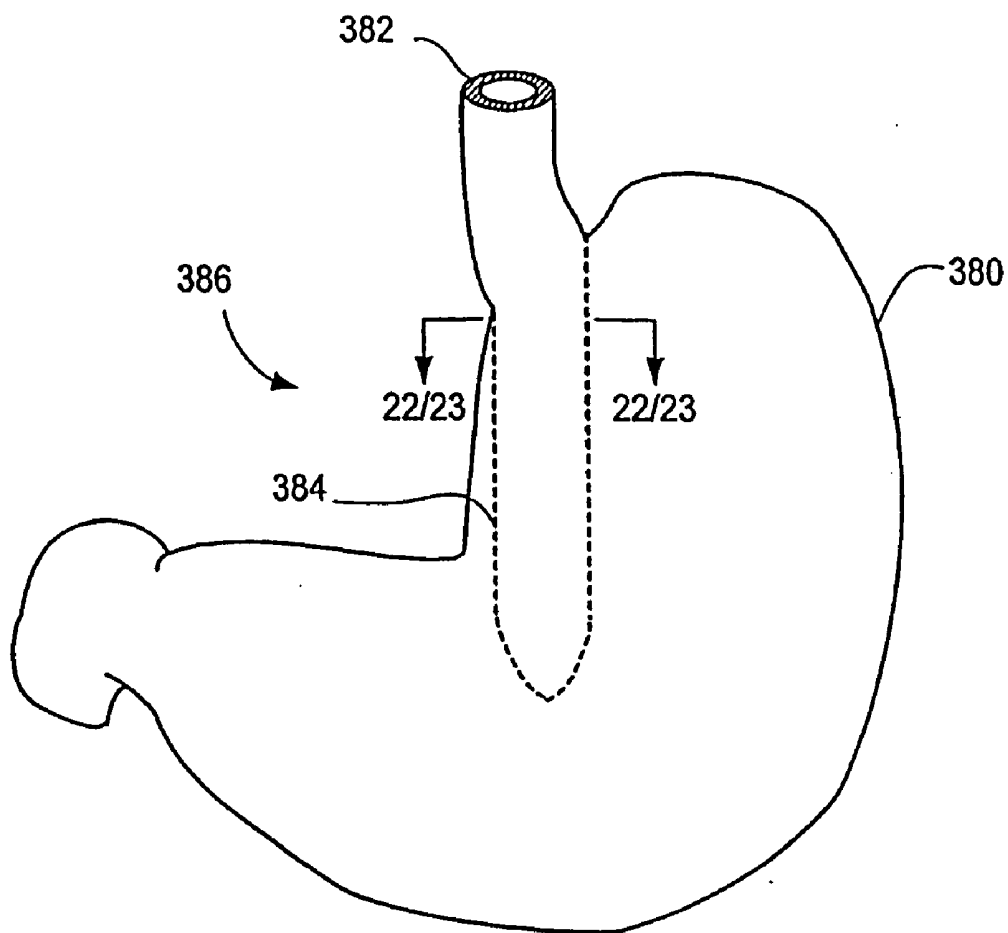


FIG. 21

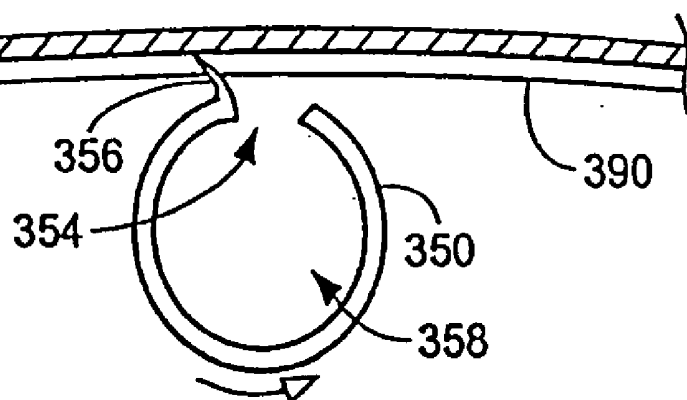


FIG. 22A

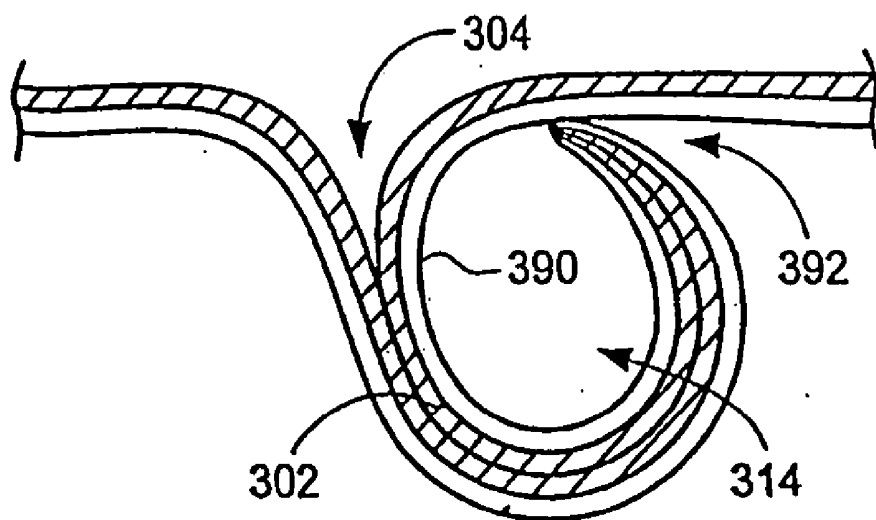


FIG. 22B

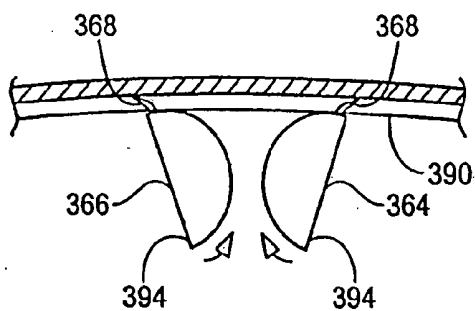


FIG. 23A

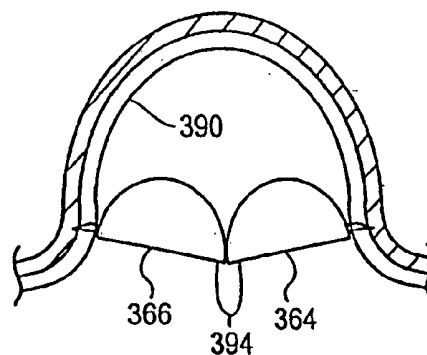


FIG. 23B

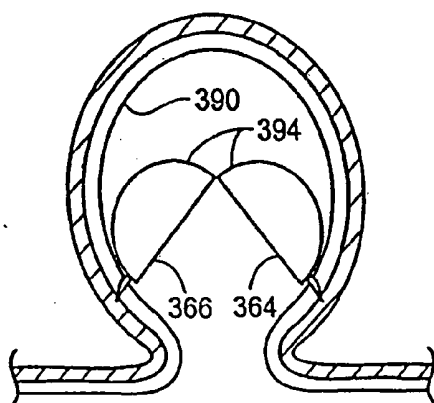


FIG. 23C

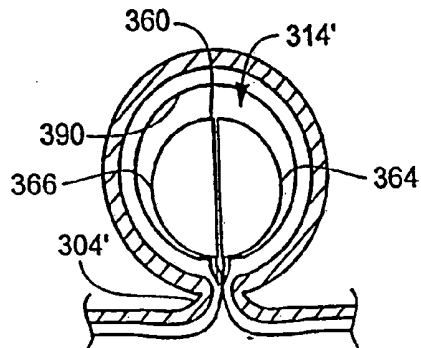


FIG. 23D

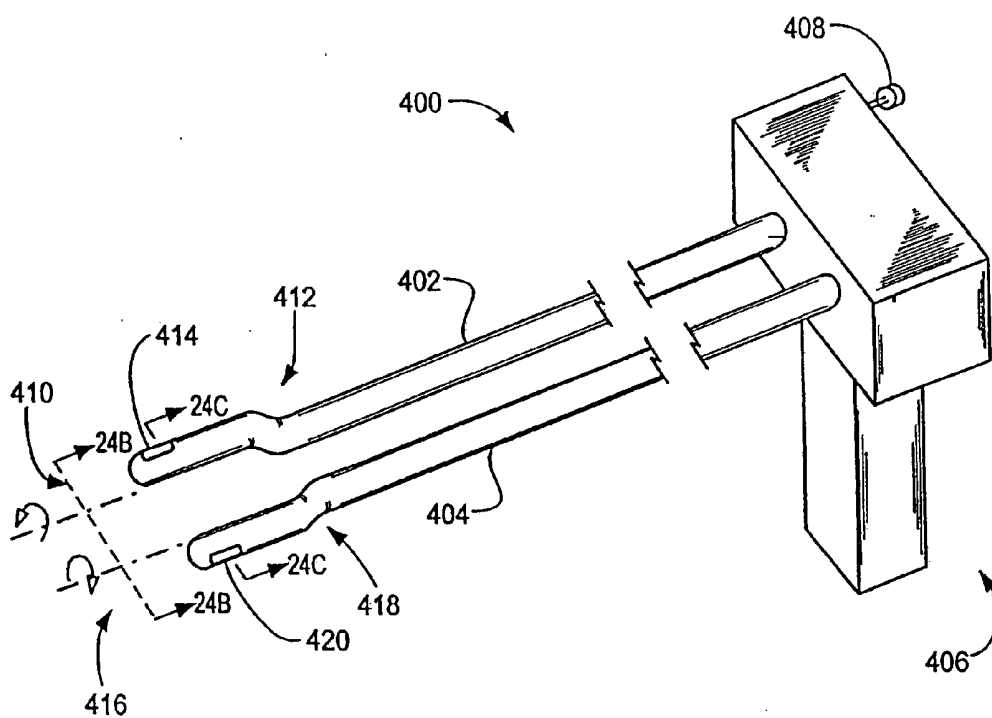


FIG. 24A

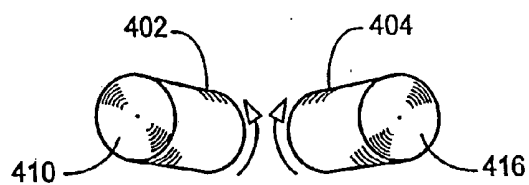


FIG. 24B

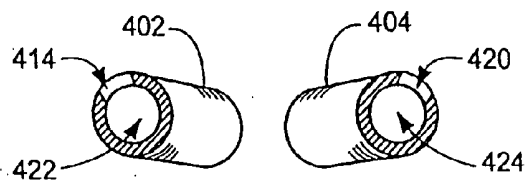


FIG. 24C

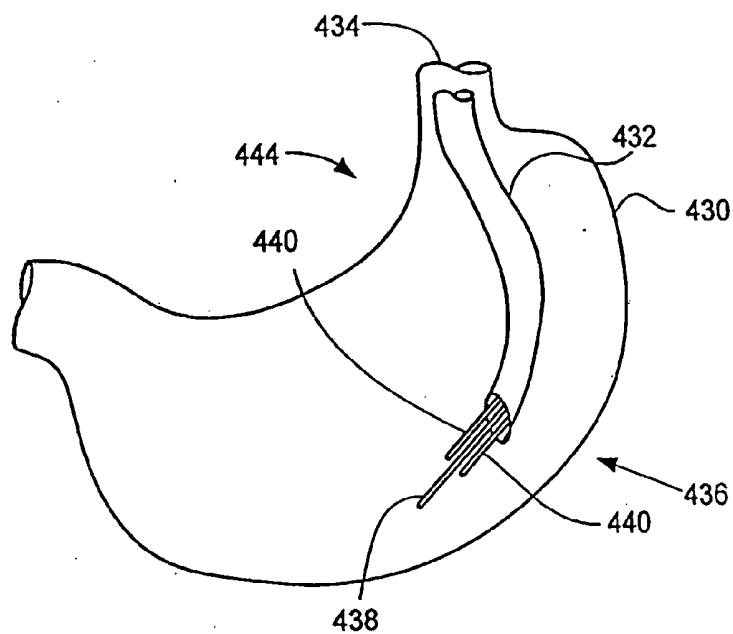


FIG. 25A

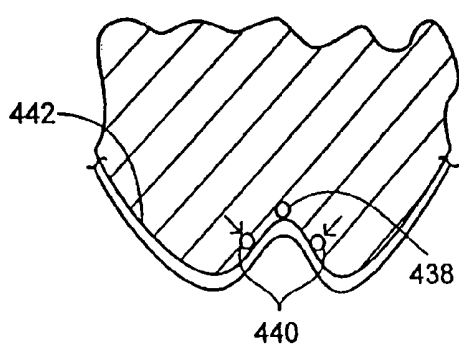


FIG. 25B

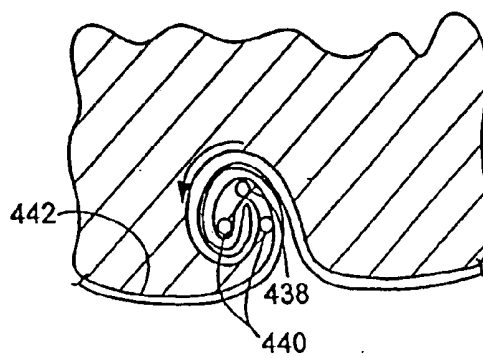


FIG. 25C

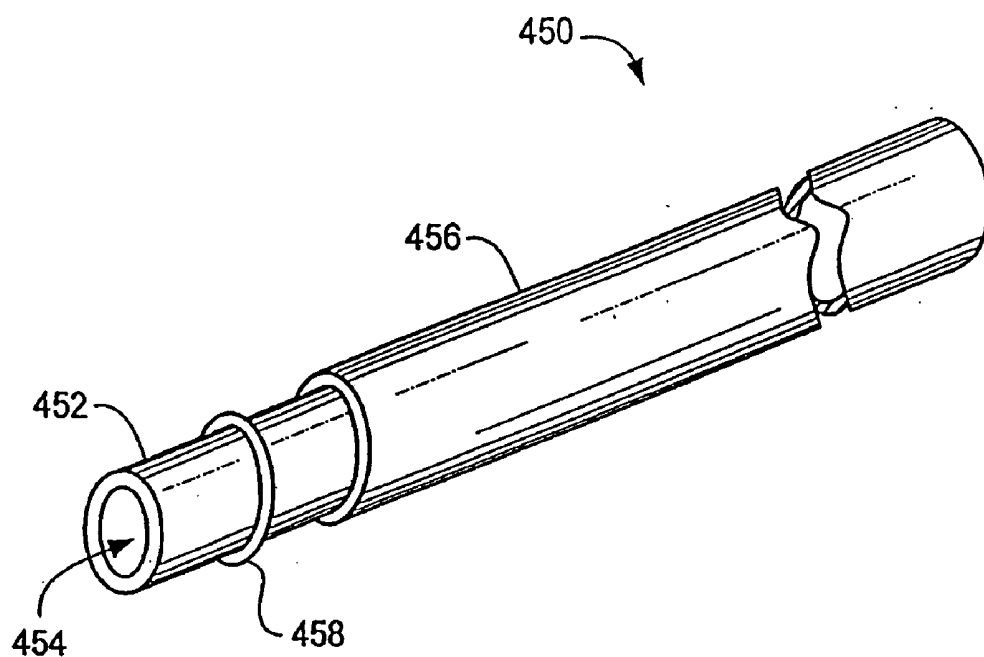


FIG. 26

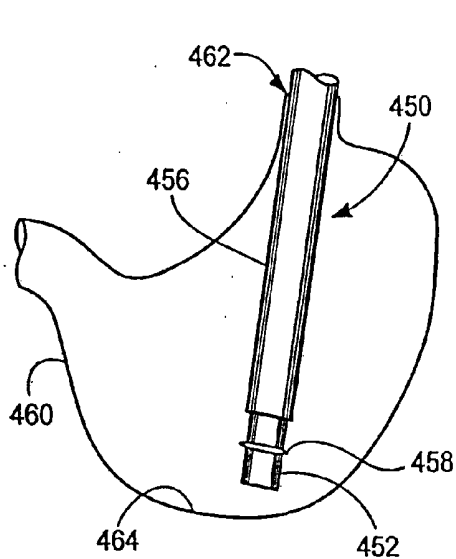


FIG. 27A

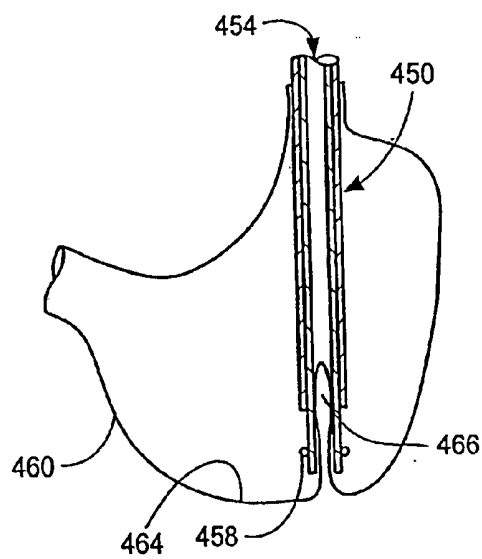


FIG. 27B

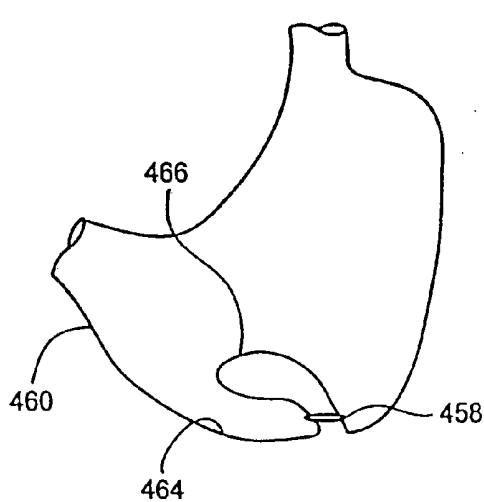


FIG. 27C

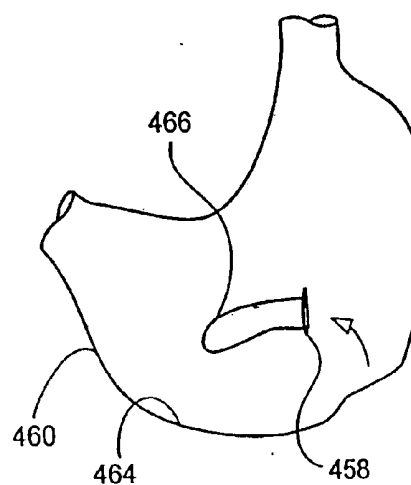


FIG. 27D

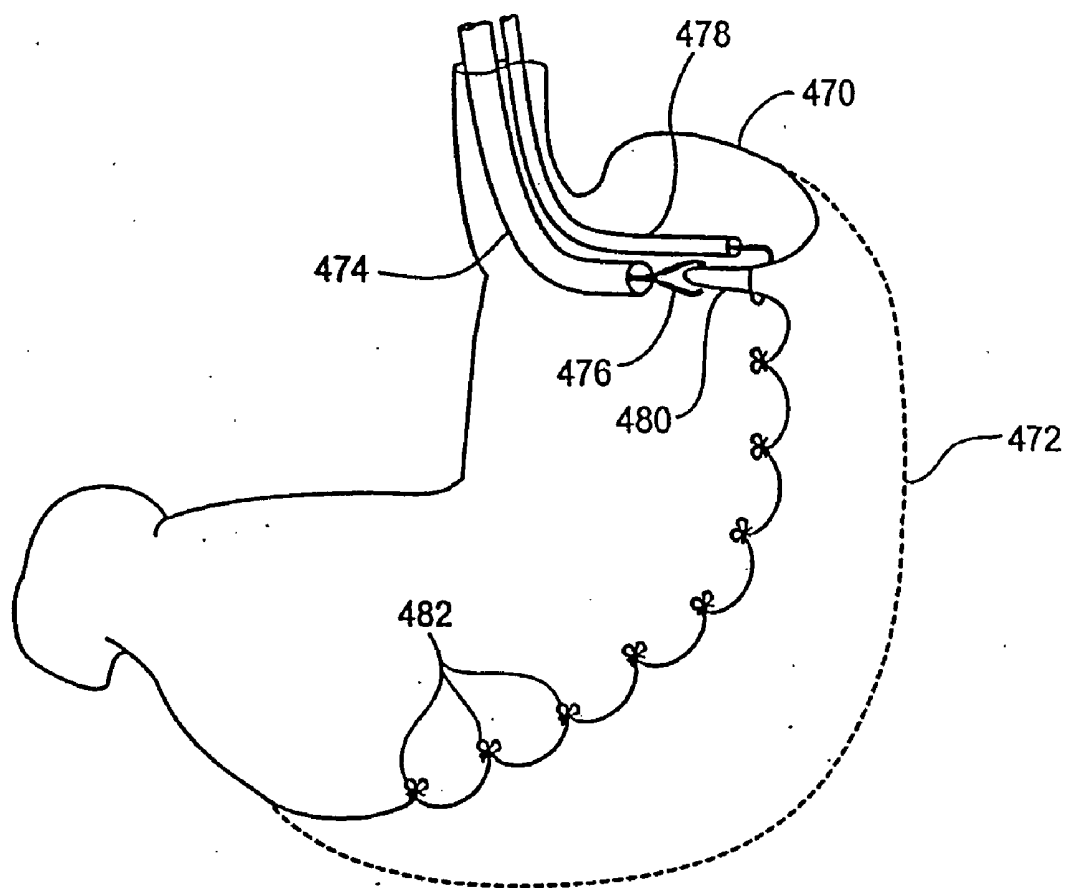


FIG. 28

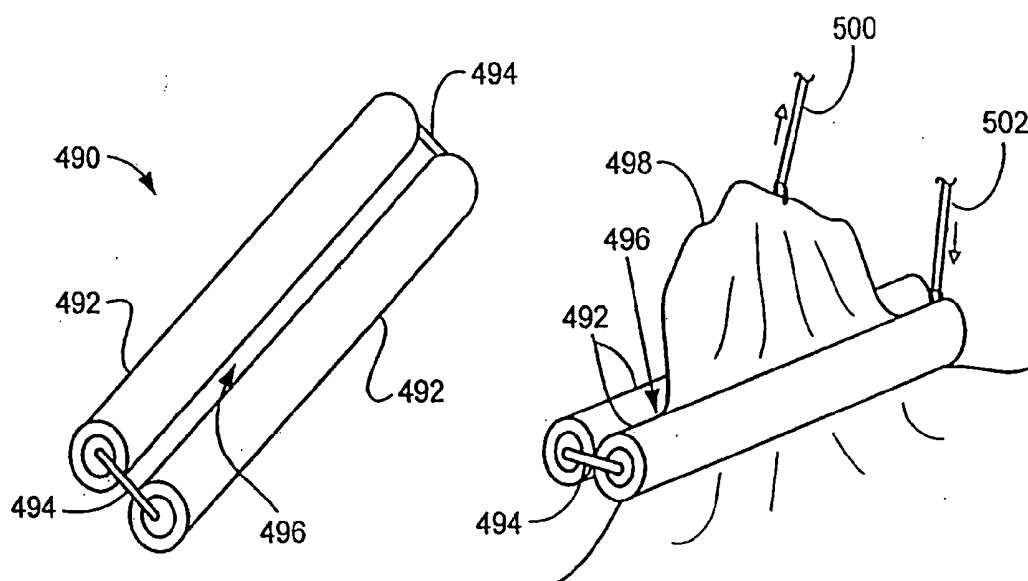


FIG. 29A

FIG. 29B

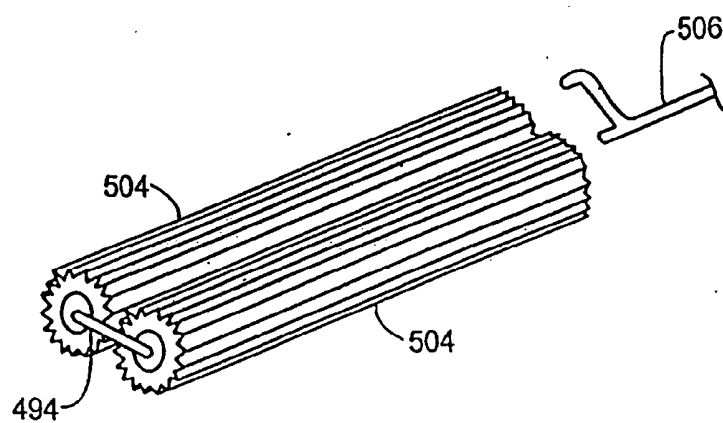


FIG. 29C

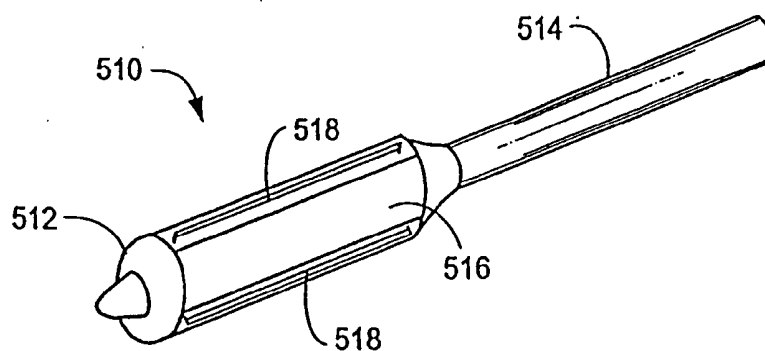


FIG. 30

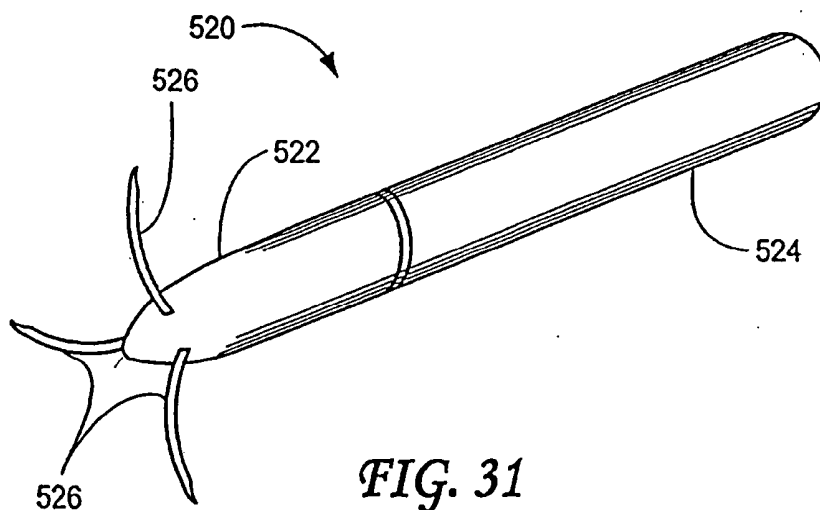


FIG. 31

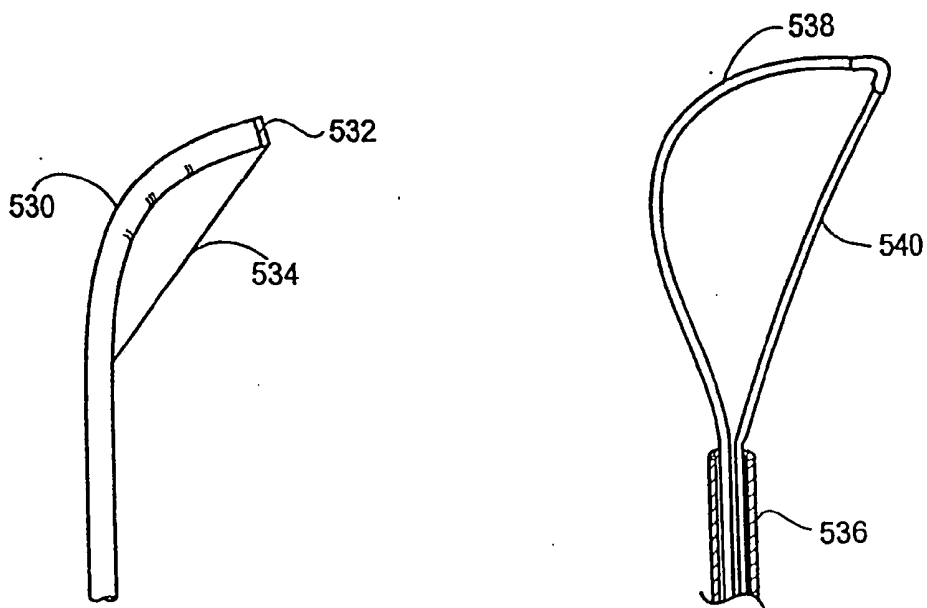


FIG. 32A

FIG. 32B

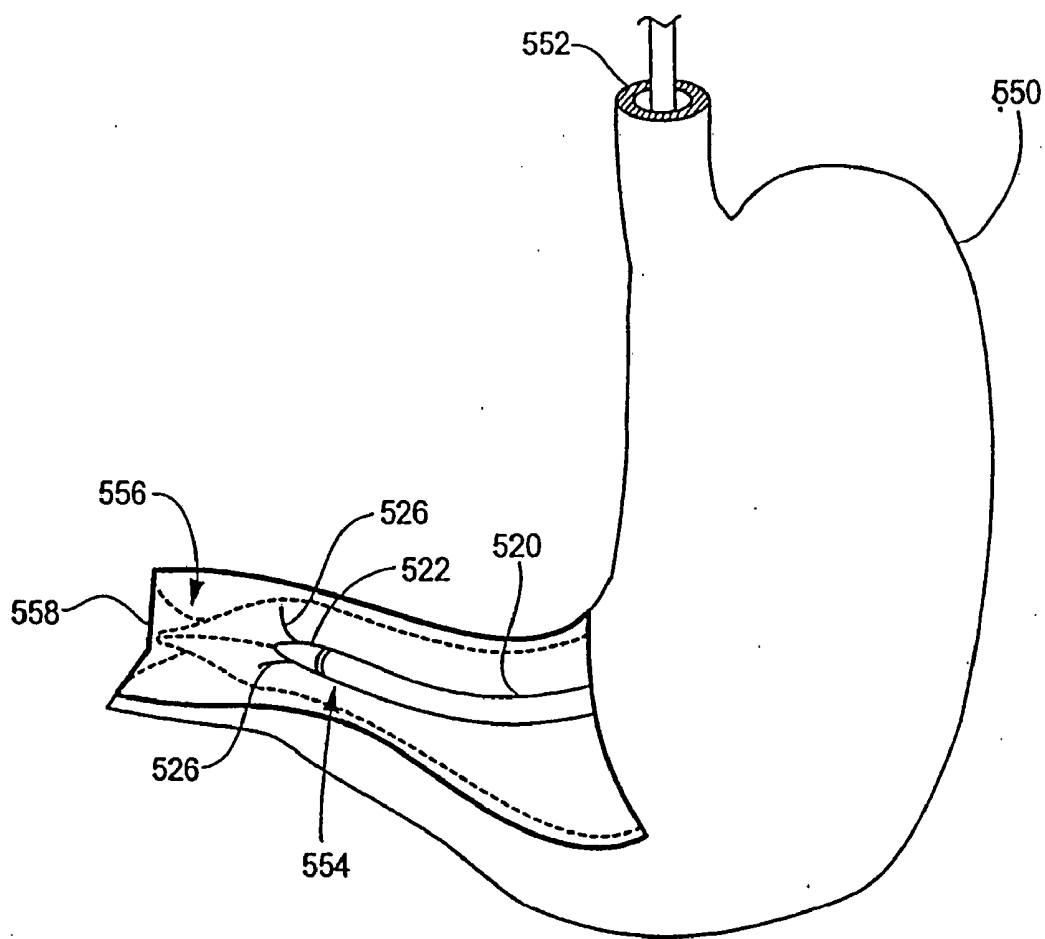


FIG. 33

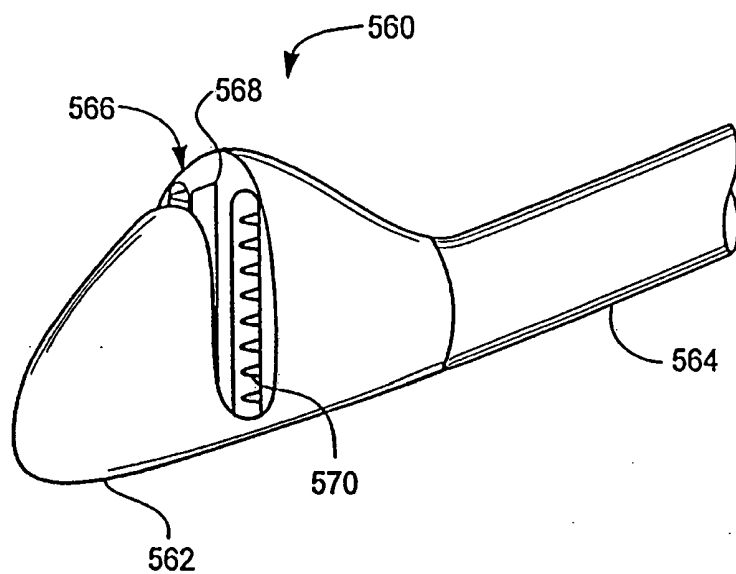


FIG. 34A

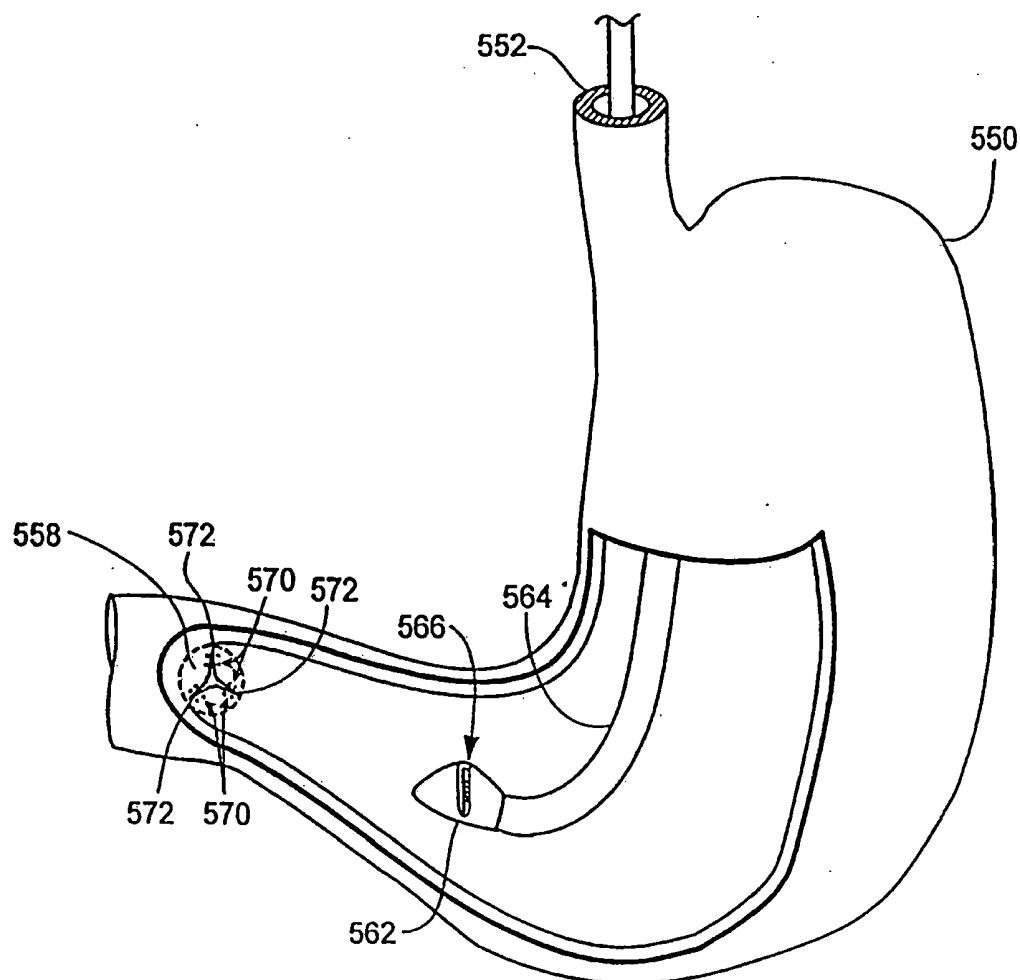


FIG. 34B

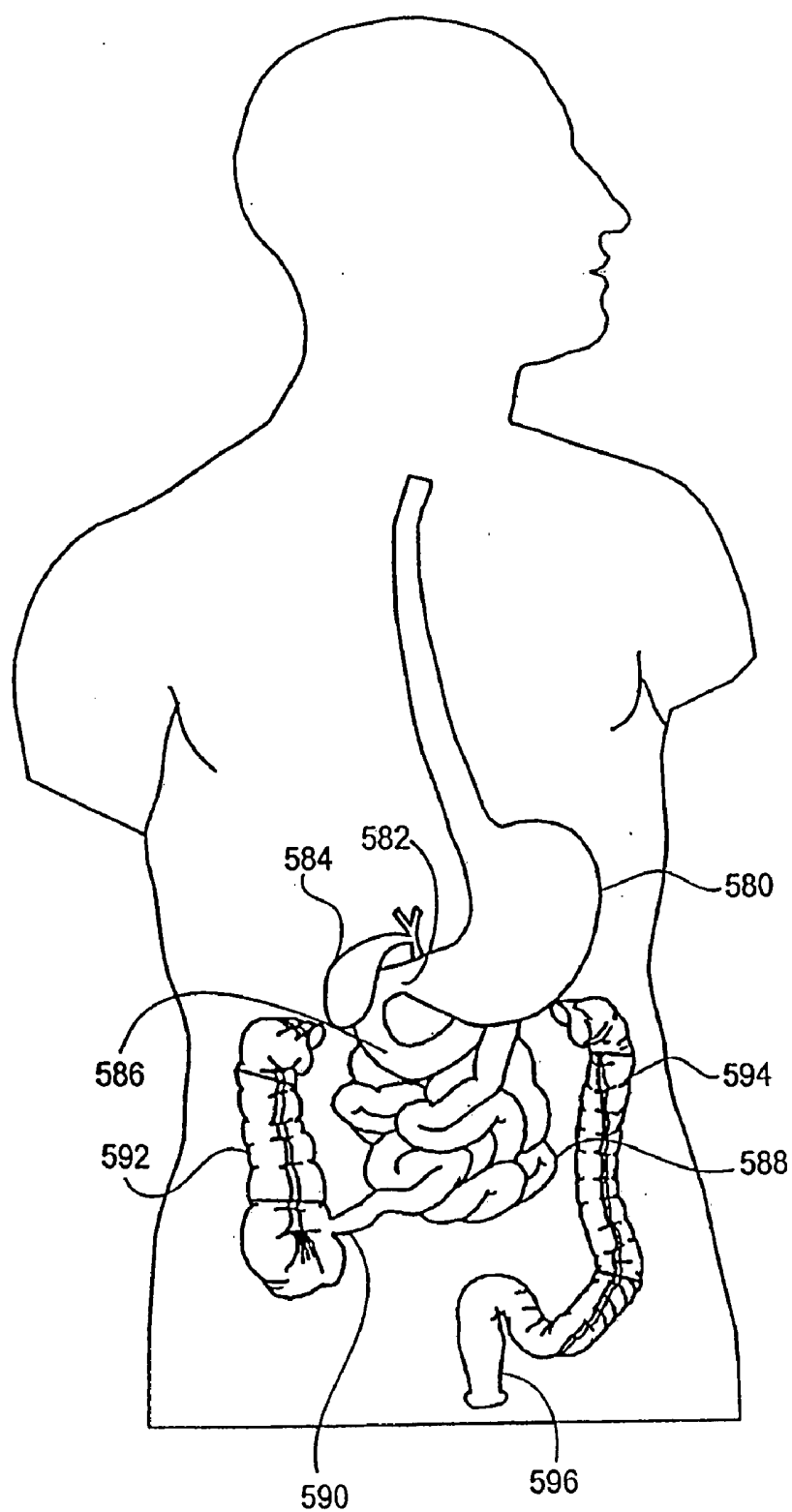


FIG. 35

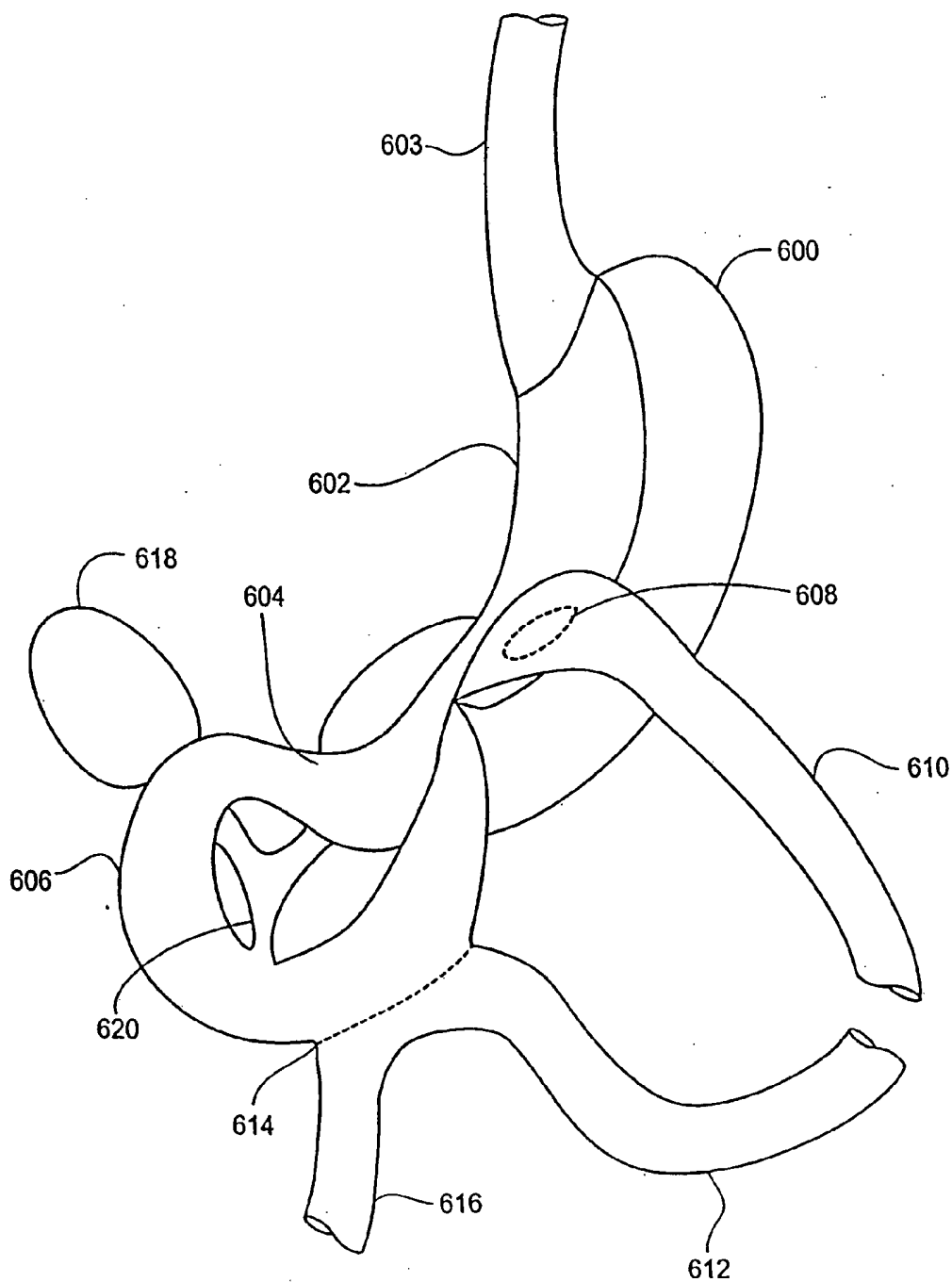
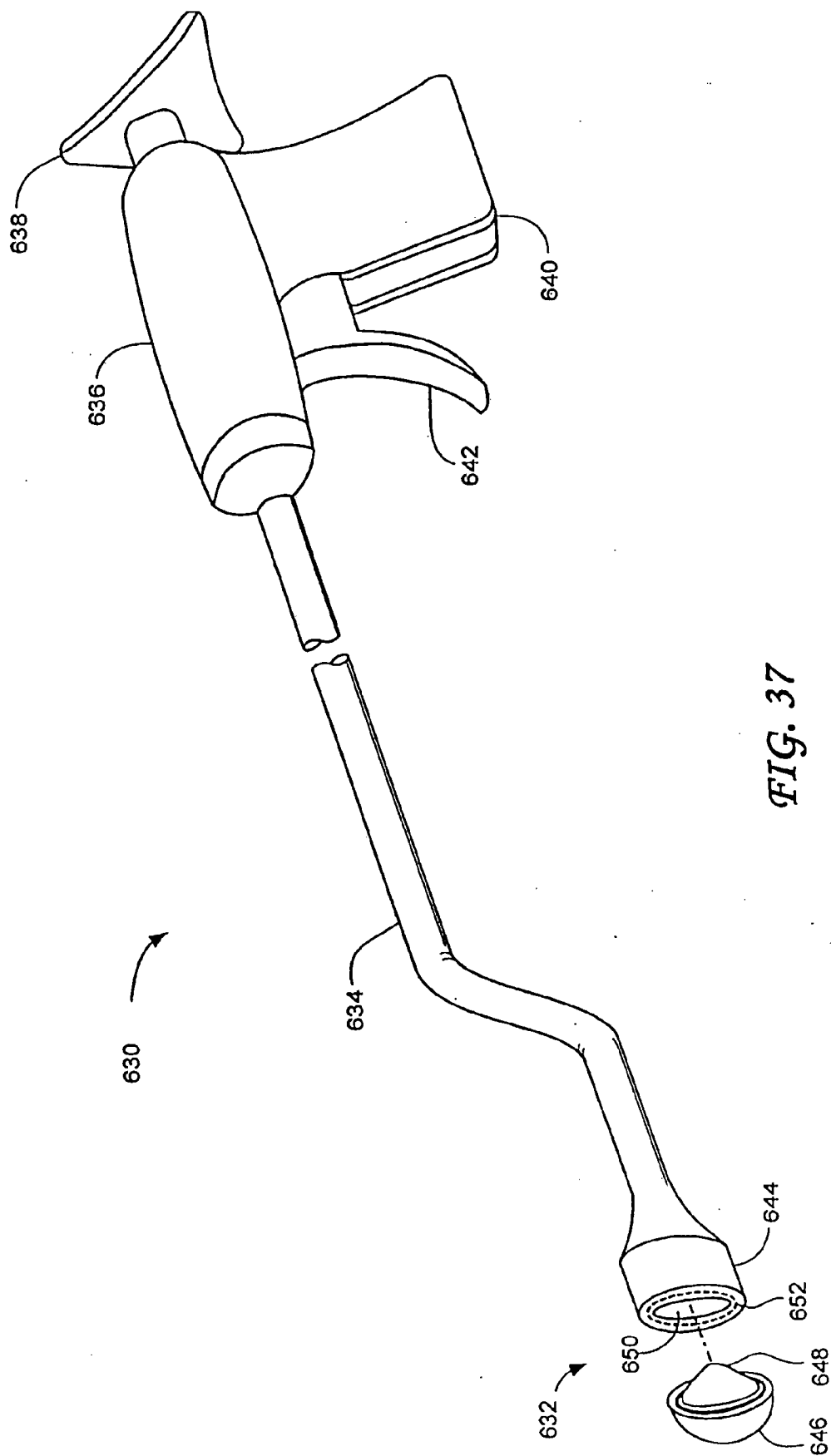


FIG. 36



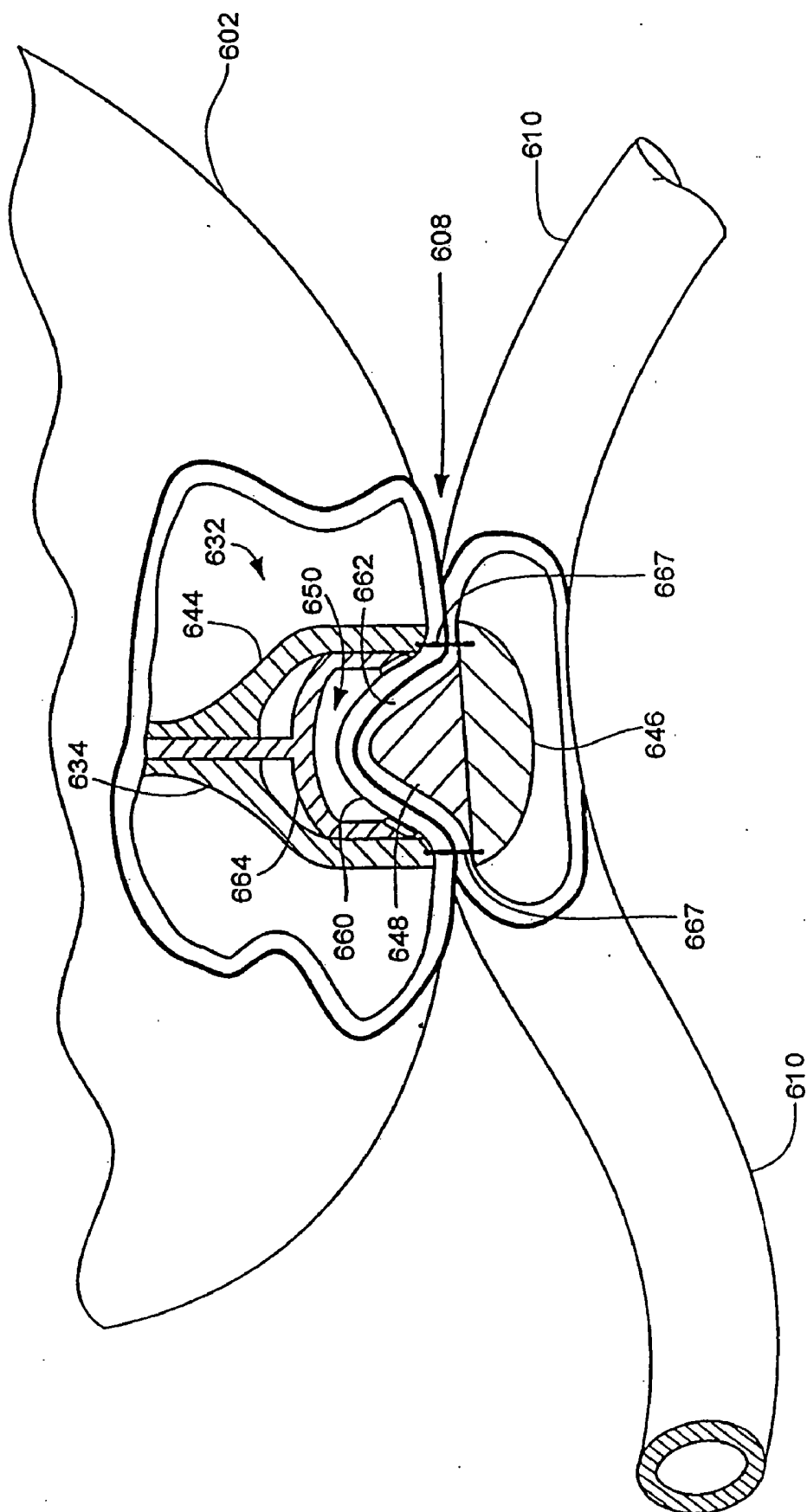


FIG. 38

OBESITY TREATMENT TOOLS AND METHODS

CROSS-REFERENCE TO RELATED APPLICATION

[0001] This application is a continuation of co-pending application Ser. No. 10/402,061, filed Mar. 27, 2003, which is a continuation of U.S. patent application Ser. No. 09/871,297, filed May 30, 2001, which is now U.S. Pat. No. 6,558,400, both of which are incorporated herein by reference.

BACKGROUND OF THE INVENTION

[0002] 1. Field of the Invention

[0003] The present invention relates generally to tools and methods for the treatment of obesity. More particularly, the present invention relates to tools and methods for performing less traumatic gastroplasty procedures.

[0004] 2. General Background and State of the Art

[0005] Obesity is considered a major health problem with annual associated costs reaching \$100 billion in the U.S. alone. Morbid obesity is a condition of obesity with the presence of a secondary debilitating progressive disease and is generally associated with a body mass index (BMI) ≥ 40 kg/m². While the basic mechanism of obesity is simply an imbalance between caloric intake and burn rate, the underlying factors are varied and complex and conservative attempts at sustained weight loss with this population are almost always unsuccessful. Often, there are genetic and other biological influences that may override environmental causes. Consequently, obesity is a disease that eludes a simple treatment, with a recurrence rate above 90% for those who attempt to lose weight. Moreover, long-term results using conservative treatments for morbid obesity are generally unsuccessful and are typically associated with further loss of self-esteem with the regaining of weight. Hypertension, cardiovascular disease, diabetes, along with a host of other comorbidities all make morbid obesity second only to smoking as a preventable cause of death.

[0006] Surgical procedures for obesity date back to 1889 (Billroth) with the earliest peer reviewed procedure being the jejuno-ileal bypass in 1954 (Kreman). A successful procedure is commonly defined as one that results in at least 50% excess weight loss at 2 years. Today, the most commonly done operation is the Roux-en-Y gastric bypass (RYGB), with around 35,000 performed annually in the U.S. Other forms of bariatric surgery include Fobi pouch, bilio-pancreatic diversion, and gastroplasty or "stomach stapling". The single existing procedure that involves an implanted device is the Lap-Band, which is a laparoscopically installed inflatable cuff that is placed around the top of the stomach just below the lower esophageal sphincter (LES). This device affects satiety only (no reduced caloric absorption). Because there is more to obesity than simple overeating, it is unlikely that Lap-Band by itself will ever be as effective as a surgery that includes other physiologic feedback mechanisms.

[0007] The RYGB procedure is a procedure which has become very common in bariatric surgery. This procedure facilitates the movement of the jejunum to a high position by using a retrocolic Roux-en-Y loop. The procedure is generally performed through a 6-8 inch incision extending from

the end of the breastbone to just above the navel. The stomach is completely divided into 2 unequal portions (a smaller upper and a larger lower gastric pouch) using an automatic stapling device with the raw surface reinforced with additional sutures. The upper pouch typically measures less than about 1 ounce or 20 cc, while the lower larger pouch remains generally intact and continues to secrete stomach juices flowing through the intestinal tract.

[0008] A segment of the small intestine (just distal of the duodenum or proximal of the jejunum) is then brought from the lower abdomen and joined with the upper pouch to form an end-to-end anastomosis created through a half-inch opening, also called the stoma. This segment of the small intestine is called the "Roux loop" and carries food from the upper pouch to the remainder of the intestines, where the food is digested. The remaining lower pouch and the attached segment of duodenum are then reconnected to form another anastomotic connection to the Roux loop at a location approximately 50-150 cm (1.6-4.9 ft) from the stoma, typically using a stapling instrument. It is at this connection that the digestive juices from the bypassed stomach, pancreas, and liver enter the jejunum or ileum to aid in the digesting of food. Due to the small size of the upper pouch, patients are forced to eat at a slower rate and are satiated much more quickly, thereby reducing the caloric intake (typically between about 1000-1200 Calories).

[0009] Because the food enters the intestines directly, conditions known as the "dumping syndrome" are created when certain types of "junk foods" are consumed (usually sweets and other simple carbohydrates). This creates unpleasant feelings of nausea, diarrhea, nervousness, and sweating, which in turn discourages patients from developing unhealthy eating patterns. With the RYGB procedure, a loss of at least 50% of excess body weight (EBW) is maintained in approximately 60% of patients at 5 years with a reduced complication rate than other procedures.

[0010] In creating the anastomoses in the RYGB procedure, several methods have previously been developed to maintain channel integrity. However, the conventional RYGB procedure requires a great deal of operative time and because of the degree of invasiveness, post-operative recovery time can be quite lengthy and painful.

[0011] Aside from the RYGB procedure, another gastrointestinal disease which relates to the stomach is gastroesophageal reflux disease (GERD). The lower esophageal sphincter is located in a distal portion of the esophagus adjacent to the junction between the esophagus and the stomach. When food is digested, a properly functioning lower esophageal sphincter would allow food to pass from the esophagus to the stomach while preventing reverse flow. However, GERD is a disorder where the esophageal sphincter allows the stomach contents, which includes gastric acid and bile, to flow back into the distal portion of the esophagus. Some complications associated with GERD include heartburn, pulmonary disorders, chest pain, esophageal ulcers, esophagitis, Barrett's esophagus, and esophageal carcinoma.

[0012] Common treatments for GERD include the administration of prescription acid blockers. But these drugs afford only short term relief; additionally, these drugs can be expensive and may have long-term side effects. Surgical procedures have included a procedure called the Nissen

fundoplication, where a portion of the gastric fundus is wrapped around the esophagus. The wrapped fundus applies pressure to the esophagus to limit the reverse flow of the stomach contents. Effectively elongating the esophagus by fundoplication or by extending it via a staple line may be done to treat GERD. Conventional fundoplication procedures may be effective at treating GERD, but they also have disadvantages. For instance, many of these procedures require large incisions to be made in a patient. Laparoscopic procedures typically require several smaller incisions formed in the abdominal wall for the insertion of instruments into the patient's body. However, such procedures can be expensive and they can increase the risks of post-operative hernias, accidental organ perforations, and other related drawbacks.

[0013] Examples related to the field of gastropasty are described below.

[0014] U.S. Pat. No. 5,549,621 to Bessler et al., which is incorporated herein by reference in its entirety, pertains to an apparatus and method for performing vertical banded gastropasty without the use of staples. The described device uses at least two clamping bars to create a tubular-shaped pouch. However, the device is deployed laparoscopically onto the external surface of the stomach.

[0015] U.S. Pat. No. 5,382,231 to Shlain, which is incorporated herein by reference in its entirety, describes a device for transesophageal stomach retraction by a device having vacuum ports utilized to draw the stomach over the device. However, this device is used for manipulating and retracting a patient's stomach from the inside during a variety of surgical procedures and is not a permanent procedure for creating an internal pouch within the stomach itself.

[0016] U.S. Pat. No. 5,345,949 to Shlain, which is incorporated herein by reference in its entirety, relates to laparoscopic methods and tools for inserting a banding device to bring the walls of the stomach adjacent to one another between the proximal pouch and the distal region of the stomach. But there is no procedure for the creation of an internal pouch internally created from the stomach.

[0017] Examples related to the field of GERD treatment are described below.

[0018] U.S. Pat. No. 6,159,146 to El Gazayerli, which is incorporated herein by reference in its entirety, relates to a device which is inserted transesophageally and engages the inside anterior wall of the fundus and secures it to the side of the esophagus.

[0019] U.S. Pat. No. 6,113,609 to Adams, which is incorporated herein by reference in its entirety, pertains to a system which includes placement of a distal anchor through a hole formed in the wall of the esophagus and through a hole formed in the gastric wall, which are then fastened together.

[0020] U.S. Pat. No. 5,571,116 to Bolanos et al., which is incorporated herein by reference in its entirety, pertains to an invagination device which approximates the lower esophagus and the fundus of the stomach.

[0021] However, all of these examples are limited to treatments for GERD which involves the attachment of the fundus, or upper portion of the stomach, to the esophagus.

SUMMARY OF THE INVENTION

[0022] Various tools and methods of treatment for obesity are described herein which are less traumatic and less invasive than procedures currently available. A variety of methods for the treatment of obesity, as well as other gastric-related diseases, e.g., gastroesophageal reflux disease (GERD), are disclosed. One method involves reducing the size of the stomach pouch to limit the caloric intake as well as to provide an earlier feeling of satiety. This may be done by creating a smaller gastric pouch within the stomach. This procedure optionally may be enhanced by performing a pyloroplasty prior to and/or in conjunction with the pouch size reduction, i.e., rendering the pyloric sphincter incompetent. This increases the rate of stomach emptying, allowing sugars and fats to pass directly into the bowel, thereby inducing dumping. Moreover, the food in the stomach may be made to also bypass a proximal portion of the bowel, i.e., a portion of the duodenum and jejunum, by creating a gastric anastomosis thereby creating a malabsorption of sugars and fats which are mostly absorbed in the bypassed portion of the duodenum and jejunum. Sugars and fats entering the bowel directly from the stomach rather than passing through the pylorus and proximal duodenum and jejunum may cause "dumping" syndrome and diarrhea. This in turn may create enforced behavioral modifications, thereby discouraging the patient from eating these types of high-caloric foods.

[0023] In forming a modified pouch, a marking device, such as a bougie, may be used at the beginning of the procedure, to create a dye marker "road map" on the interior surface of the stomach from the pylorus to the esophagus. This may enable visualization by, e.g., an endoscope, to give the physician a clear reference point for staple or fixation element placement. A distal balloon, which is preferably attached to an inflation tip at a distal end, may be inserted into the pylorus to stabilize the bougie during the procedure and may be inflated from the proximal end of the tubing by the physician.

[0024] In reducing the stomach size, one variation involves grasping the interior walls of the stomach, preferably via an endoscope advanced transesophageally, and placing one to several individual fixation elements on opposing interior walls and then bringing those fixation elements together. The stomach pouch may be modified and/or created by a variety of other device variations utilizing other methods, e.g., stapling opposing sides of a stomach together to form two separate lumens from within the interior surface of the stomach. An endoscopic stapling device may be used to accomplish such a task. Such an endoscopic stapler preferably brings two regions of tissue into apposition and may then apply a fastening element, e.g., staples, clips, tags, screws, etc., into the two regions of tissue to affix them together.

[0025] In addition to endoscopically applied stapling and clip devices, rotating and rotatable probes may also be used to form a modified smaller lumen within a main lumen. Such probes generally may be inserted into a stomach endoscopically and may engage a portion of the interior lining of the stomach and may then be rotated to roll the engaged portion of the stomach wall around the probe itself to bring the wall in apposition with another portion of the stomach wall. Such rotating probes may be used to create a blind-ended pouch of stomach within the main stomach lumen, or as with the

other devices, may be used to create a smaller pouch exiting into the pylorus. Once the roll of stomach wall is brought into apposition, a row or a plurality of fasteners, e.g., staples, blind staples, clips, tags, adhesives, screws, etc., may be used to maintain the stomach. Moreover, other variations may include gastric volume reduction devices as part of the present invention. Such volume reduction devices generally may be inserted into a stomach trans-esophageally through the use of, e.g., an endoscope. The reduction device may be used to draw or engage a portion of the interior lining of the stomach; the drawn or engaged portion may then be eventually removed, either actively or through natural processes, e.g., pressure necrosis.

[0026] To aid in the overall effect, a pyloroplasty procedure may also be performed to enhance treatment. The pyloroplasty may be performed prior to (preferable), in conjunction with, or following the gastric reduction procedure. A pyloroplasty procedure typically results in the pyloric sphincter being rendered incompetent. Generally, a pyloroplasty device may be passed endoscopically through the esophagus, into the stomach, and preferably into position in or across the pylorus. Energy or a stimulus is then preferably applied to the pylorus to render it incompetent.

[0027] Moreover, an additional anastomosis gastric bypass procedure may also be performed to further enhance treatment. The anastomosis procedure may be performed preferably prior to, in conjunction with, or following the gastric reduction and pyloroplasty procedures (if performed at all). The procedure generally involves endoscopically or laparoscopically creating a side-to-side anastomosis preferably from within the stomach and bowel and within the digestive tract. This procedure may be similar to the Roux-en-Y gastric bypass procedure but with minimal trauma.

BRIEF DESCRIPTION OF THE DRAWINGS

[0028] FIG. 1A shows an example of a modified stomach having a smaller pouch created from the interior surface lining.

[0029] FIG. 1B shows a partial superior view of the cross section from FIG. 1A.

[0030] FIG. 2 shows a variation on a marking device or bougie for marking the interior surface of a stomach.

[0031] FIG. 3A shows a variation on positioning a marking device inserted into a stomach.

[0032] FIG. 3B shows a cross section view from FIG. 3A of a deflated stomach around the marking device.

[0033] FIG. 3C shows the cross section view from FIG. 3B of an insufflated stomach with the resulting marks.

[0034] FIG. 4A shows a view of the interior of the lesser curvature of a stomach with anchors attached.

[0035] FIG. 4B shows a cross section view from FIG. 4A with the anchors attached.

[0036] FIG. 5A shows a side view of a crimping variation on a fastening device.

[0037] FIGS. 5B and 5C show a superior and side view, respectively, of several interlocked crimping devices from FIG. 5A.

[0038] FIG. 6A shows an isometric view of a zip-tie or ratcheted variation on a fastening device.

[0039] FIG. 6B shows a superior view of the device of FIG. 6A attached to the stomach wall.

[0040] FIG. 6C shows a superior view of another double zip-tie variation on a fastening device.

[0041] FIG. 6D shows the stomach of FIG. 6B with the fasteners cinched.

[0042] FIG. 6E shows a superior view of another perpendicular zip-tie variation on a fastening device.

[0043] FIGS. 7A and 7B show a superior view of an extendable double hook device attaching to a stomach wall.

[0044] FIG. 7C shows the device of FIG. 7A locked by a crimping variation.

[0045] FIGS. 8A and 8B show a superior view of a modified stomach maintained by a fastening staple.

[0046] FIGS. 9A and 9B show isometric views of a variation on an endoscopic stapling device.

[0047] FIG. 10 shows an isometric view of a variation on a box stapling device.

[0048] FIG. 11A shows an assembly view of another stapling device variation.

[0049] FIG. 11B shows a side view of the device of FIG. 11A.

[0050] FIG. 12A shows an isometric view of a crescent shaped variation of a stapling device.

[0051] FIG. 12B shows an end view of the device of FIG. 12A showing a staple deploying.

[0052] FIG. 12C shows an interior side view of the device of FIG. 12A with a translating wedge sequentially deploying staples.

[0053] FIG. 13 shows an interior view of a stomach with an example of stapling device placement.

[0054] FIG. 14 shows an interior view of a stomach with an example of a modified stapling device which may be used for the treatment of GERD.

[0055] FIG. 15A shows an assembly view of a variation on an approximating device.

[0056] FIGS. 15B to 15D show the process of invaginating stomach interior lining and fastening using the device of FIG. 15A.

[0057] FIG. 15E shows the assembly view of another variation of the device of FIG. 15A wherein the clip may be replaced by a screw.

[0058] FIGS. 15F to 15H show the process of invaginating stomach interior lining and fastening using the device of FIG. 15E.

[0059] FIG. 16A shows an example of a modified stomach created by a rotating device variation.

[0060] FIG. 16B shows a superior cross section view of the stomach of FIG. 16A where the modified lumen may be created by rotating the interior stomach lining upon itself.

[0061] FIG. 16C shows an alternate superior cross section view of the stomach of FIG. 16A where the modified lumen may be created by rotating apposed portions of the interior stomach lining upon itself.

[0062] FIGS. 17A and 17B show an isometric and cross section view, respectively, of a vacuum tube variation.

[0063] FIGS. 18A and 18B show an isometric and cross section view, respectively, of a counter-rotating vacuum tube variation.

[0064] FIGS. 19A and 19B show an isometric and cross section view, respectively, of a vacuum tube variation with attachment points.

[0065] FIGS. 20A and 20B show an isometric and cross section view, respectively, of a split tube variation.

[0066] FIG. 21 shows an example of placement within a stomach of a rotatable device variation.

[0067] FIGS. 22A and 22B show the possible creation of a rotated lumen using the device of FIGS. 19A and 19B.

[0068] FIGS. 23A to 23D show the possible creation of a rotated lumen using the device of FIGS. 20A and 20B.

[0069] FIG. 24A shows an isometric view of a variation on a dual rotatable tube device.

[0070] FIGS. 24B and 24C show an end view and cross section view, respectively, of the device of FIG. 24A.

[0071] FIG. 25A shows a variation on an endoscopic vacuum device in a stomach.

[0072] FIGS. 25B and 25C show an end view of a variation on lumen creation from the interior surface of the stomach using the device of FIG. 25A.

[0073] FIG. 26 shows an isometric view of a variation on a gastric volume reduction device.

[0074] FIGS. 27A to 27D show the device of FIG. 26 inserted into a stomach to draw or cinch up lining tissue to reduce a volume of the stomach.

[0075] FIG. 28 shows another variation on a gastric volume reduction device utilizing a grasping device and a ligating device.

[0076] FIGS. 29A and 29B show an isometric view on a variation of a gastric volume reduction device utilizing tractive rollers to draw tissue up between them.

[0077] FIG. 29C shows another variation of the device of FIGS. 29A and 29B with ratcheted rollers.

[0078] FIG. 30 shows an isometric view of a variation on a pyloroplasty device with an angioplasty balloon.

[0079] FIG. 31 shows an isometric view of another variation on a pyloroplasty device with extendable probes.

[0080] FIGS. 32A and 32B show variations on sphincterotomy arms for use in a pyloroplasty procedure.

[0081] FIG. 33 shows a stomach with a distal portion of the wall of the lesser curvature removed to show a possible use for the device of FIG. 31.

[0082] FIG. 34A shows an isometric view of another variation on a pyloroplasty device with a combination cutting and stapling notch.

[0083] FIG. 34B shows the device of FIG. 34A in a possible use in a stomach.

[0084] FIG. 35 shows a representative and normal gastro-intestinal system of a person.

[0085] FIG. 36 shows an example of a gastro-intestinal system modified by a preferable anastomosis procedure.

[0086] FIG. 37 shows an isometric view of a variation on an anastomosis deployment device.

[0087] FIG. 38 shows a cross section view of an anastomosis assembly mating a portion of the stomach with a portion of the intestinal tract.

[0088] FIG. 39 shows a cross section view of another anastomosis assembly mating two different portions of the intestinal tract.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

[0089] With obesity becoming an increasing problem, various tools and methods of treatment are described herein which are less traumatic and less invasive than procedures currently available. As described in further detail below, a variety of methods for the treatment of obesity, as well as other gastric-related diseases, are disclosed. Generally, the size of the stomach pouch may be reduced to limit the caloric intake as well as to provide an earlier feeling of satiety. This may be accomplished by creating a smaller gastric pouch within the stomach by a variety of methods. This procedure optionally may be enhanced by performing a pyloroplasty prior to and/or in conjunction with the pouch size reduction, i.e., rendering the pyloric sphincter incompetent. Additionally, the food in the stomach may be made to also bypass a proximal portion of the bowel, i.e., a portion of the duodenum and jejunum, by creating a gastric anastomosis thereby creating a malabsorption of sugars and fats which are mostly absorbed in the bypassed portion of the duodenum and jejunum. Sugars and fats entering the bowel directly from the stomach rather than passing through the pylorus and proximal duodenum and jejunum may cause "dumping" syndrome and diarrhea. Moreover, rendering the pylorus incompetent may also lead to dumping syndrome partly because of the rapid gastric emptying which may occur. This in turn may create enforced behavioral modifications, thereby discouraging the patient from eating these types of high-caloric foods.

[0090] FIG. 1A shows an example of a modified stomach 10 which may be created, by any one of the methods described below, as part of the present invention. Greater curvature 12 and lesser curvature 14 is seen in modified stomach 10, as well as the distal end of esophagus 16 and pylorus 18. As part of the present invention, stomach 10 may be divided along junction 24 into modified pouch 22, which is preferably less than about 1 ounce in volume, and main pouch 20. FIG. 1B shows a partial superior view of the cross section of main pouch 20 and modified pouch 22 as viewed from cutting plane P from FIG. 1A. As seen, modified lumen 26 is preferably formed by junction 24 from main lumen 28 by joining a portion of stomach wall 30. During ingestion of food, modified pouch 22 accepts food from esophagus 16 and preferably passes it directly through modified lumen 26 into pylorus 18. Main pouch 20 may remain intact and function normally, but preferably sees little or no food. Acids

and other fluids that may be generated in main lumen 28 may drain through the reduced outlet near pylorus 18 and may pass through the digestive system normally.

[0091] Marking Tools and Methods

[0092] As part of forming a modified pouch, a marking device may be used, preferably at the beginning of the procedure, to create a dye marker “road map” on the interior surface of the stomach from the pylorus to the esophagus. Once such dye marks are placed, they may be visualized, e.g., endoscopically, thereby giving the physician a clear reference point for staple or fixation element placement. An example of such a marking device is shown in FIG. 2 as marking device or bougie 40. Bougie 40 is preferably an elongated device made from tubing member 44 which may have several channels defined within. Tubing 44 may be made from any variety of biocompatible materials, e.g., stainless steel, plastics, etc., and preferably has a diameter and cross section which is similar to that of the finished modified lesser pouch. Along the length may be defined a series of dye ports 46 through which the marking dye may be channeled through from the proximal end of bougie 40. Any variety of biocompatible dyes which preferably enhance visualization may be used, e.g., methylene blue, thionine, acridine orange, acridine yellow, acriflavine, quinacrine and its derivatives, brilliant green, gentian violet, crystal violet, triphenyl methane, bis naphthalene, trypan blue, and trypan red. Also along the length and on either side of dye ports 46 may be a series of vacuum ports 48, which are optional. A distal balloon 52, which may be inserted into the pylorus to stabilize bougie 40 during the procedure, is preferably attached to inflation tip 50 at distal end 42 and may be inflated from the proximal end of tubing 44 by the physician.

[0093] FIGS. 3A to 3C show bougie 40 during one method of use. FIG. 3A shows stomach 60 as bougie 40 is inserted down through esophagus 62. As bougie 40 is advanced down to pylorus 76, distal balloon 52 may be inflated through inflation tip 50, thus securing the device. Bougie 40 preferably follows lesser curvature 64 and may alternatively be shaped to approximate lesser curvature 64. Bougie 40 is also preferably rotated such that dye ports 46 face away from lesser curvature 64 and face towards greater curvature 66. Then the air and fluids contained within stomach 60 are preferably removed, either through vacuum ports 48, if they are included in bougie 40, or through another vacuum port which may be introduced endoscopically through esophagus 62. FIG. 3B shows cross section 3B-3B from FIG. 3A as deflated stomach 60. Once deflated, modified lumen 70 may take shape around bougie 40, separate from deflated main lumen 68. In this deflated state, the dye may be channeled through dye ports 46, thereby leaving dye marks 72 on interior lining 74. Once the staining has been performed, lumen 68 may be insufflated, as shown in FIG. 3C, and bougie 40 may then be removed. As seen in FIG. 3C, dye marks 72 mark or delineate the junction region where anchors or fasteners may be placed to draw interior lining 74 together to form the modified lumen.

[0094] Gastric Reduction Tools and Methods Using Fasteners

[0095] One variation of reducing the stomach size involves grasping the interior walls of the stomach, preferably via an endoscope advanced transesophageally, and

placing one to several fixation elements on opposing interior walls and then bringing those fixation elements together.

[0096] Several examples of different possible variations on fasteners are shown and described below. These variations are not intended to be limiting but are merely given as illustrative examples.

[0097] FIG. 4A shows a view of the interior of the lesser curvature of stomach 60 with part of the greater curvature wall removed. As seen, individual anchors 80 may be secured to the interior surface along the junction 24 where modified pouch 22 from FIG. 1A would form. Anchors 80 may be of any biocompatible material, e.g., stainless steel, polymers, etc., which may be formed into a variety of fasteners, e.g., staples, ratcheted wires, zip ties, clips, tags, eyelets, crimps, and screws. Anchors 80 may be placed by estimating the junction boundary, but they are preferably located along dye mark 72, which may be formed by methods and tools described above, prior to anchor 80 placement, as shown in FIG. 4B, which is cross section 4B-4B from FIG. 4A. After anchors 80 have been fastened, suture 82 may be drawn through each of the anchors 80, preferably in a zig-zag manner, and then suture 82 may be drawn tight to bring the opposing surfaces of interior lining 74 together in apposition along dye marks 72 to form the modified lumen. Alternatively, individual anchors 80 may be preloaded or prefasted by suture 82, and anchors 80 may be fastened to interior lining 74 in this manner.

[0098] FIG. 5A shows a side view of a variation on a fastening device in crimping member 90. Crimping member 90 is preferably made from a biocompatible material, e.g., stainless steel, nitinol, etc., and may be formed to have elbow 92 extend into two opposing anchoring ends 94. FIG. 5B shows a superior view of a created modified lumen 100 formed from main lumen 98 by any of the methods described herein. In this variation, several crimping members 90 may be attached or fastened to interior lining 96 by anchoring ends 94. As they become attached, each of the members 90 are preferably configured to interlock with an adjacent crimping member 90, much like a zipper. FIG. 5B shows the interlocked members 90 from the top to form lumen 100 and FIG. 5C shows the view from 5C-5C from FIG. 5B where each of the crimping members 90 are shown interlocking at their elbows 92 like a zipper.

[0099] FIG. 6A shows an isometric view of another variation on a fastening device in ratcheted wire or zip tie 110. This particular variation shows a distal tip or male end 112 and a corresponding proximal end or female end 114, with ratcheted length 116 between those two ends. FIG. 6B shows a superior view of stomach wall 120 just prior to the formation of modified lumen 124 from main lumen 122. As seen, male end 112 of first zip tie 110' maybe pierced through one side of interior lining 118 and second zip tie 110" may be pierced through the opposing side of interior lining 118 such that the male ends 112 of each zip tie preferably correspond to the female ends 114 of the other zip tie. To then form the lumen 124, each zip tie 110', 110" may be drawn together and tightened accordingly, as shown in FIG. 6D. A plurality of zip ties 110 are preferably used to form modified lumen 124 by aligning them by any of the methods described above.

[0100] An alternative zip tie device which may be used is a perpendicular type version of zip tie 110. As shown in FIG.

6E, first perpendicular zip tie **134'** and second perpendicular zip tie **134''** may be used in place of zip tie **110** and lumen **124** may be formed in much the same manner as described above to result in the modified stomach as shown in FIG. 6E. A further alternative is shown in FIG. 6C where male zip tie **126** preferably has dual piercing male ends with catcher tubes **128**. In this variation, a vacuum-type device, as described below in detail, or forceps may be used to draw portions of stomach wall **120** in apposition. As the apposed stomach walls **120** are positioned, needles **130**, which are preferably passed through a double female zip tie **132**, may be used to pierce through tissue **120** and lock into catcher tubes **128**. Needles **130** may then be drawn back through tissue **120**, while simultaneously pulling male ends/catcher tubes **128** back through tissue **120** and into the corresponding double female zip tie **132**. The locked zip tie **126** may then be drawn tight against female zip tie **132**, trimmed, and then released. This procedure may be repeated for any number of zip ties which may be used to draw the stomach lining together to form the smaller pouch and may also be used with the dye marking device **40** and procedure as described above.

[0101] A further variation on the individual anchoring fasteners is shown in FIG. 7A. This variation shows gasping device **140** with retaining tube **142** and extendable members **146** which may extend from distal opening **144**. Extendable members **146** are preferably made from a biocompatible material, e.g., superelastic or shape memory alloy such as nitinol, which may be biased to urge away from a longitudinal axis defined by tube **142** once extended beyond distal opening **144**. As members **146** extend, they may reach out to grasp apposed portions of interior lining **150** by hooks **148**. As above, the locations where hooks **148** grasp may be defined by the marking device as described above and viewed by the physician through, e.g., an endoscope. Once hooks **148** have grasped the appropriate portion of lining **150**, members **146** may then be drawn back through distal opening **144**, as shown in FIG. 7B, and a retaining device, such as crimp **152**, may be slid over a distal section of members **146**, as shown in FIG. 7C, to maintain the position of hooks **148** and apposed lining **150** to create the desired lumen.

[0102] Gastric Reduction Tools and Methods Using Stapling Devices

[0103] Aside from individual anchoring and fastening devices, the stomach pouch may be modified and/or created by a variety of other device variations utilizing other methods. FIG. 8A shows the cross sectioned superior view of FIG. 1B with the addition of staple **160** maintaining junction **24**. The figure shows an example of how, e.g., an endoscopically applied stapler, may be used to retain and hold junction **24** to form modified lumen **26**. FIG. 8B shows a close-up view of the junction **24** and staple **160** which was applied from within lumen **26**.

[0104] To staple opposing sides of a stomach together to form two separate lumens from within the interior surface of the stomach, an endoscopic stapling device may be used to accomplish such a task. Such an endoscopic stapler preferably brings two regions of tissue into apposition and may then apply a fastening element, e.g., staples, clips, tags, etc., into the two regions of tissue to affix them together. These stapling devices may optionally incorporate the use of the

marking device or bougie **40**, as described above, as a preliminary step as a guide to vacuum placement and/or stapling to form the desired modified lumen. The fastening elements, e.g., staples, are preferably made of a biocompatible material such as stainless steel, titanium, polymers, sutures, nitinol, or any other similar metals and alloys, etc. and may be in any conventional shape such as C-shaped and U-shaped staples or any of the other shapes as described herein. The two regions of tissue may be adhered to the stapling device by a variety of attachment methods, e.g., tines, barbs, hooks, vacuum, or any combinations thereof. In an adhering device utilizing a vacuum to hold the apposing regions of tissue together, such a device may be a tubular or wand-shaped member and preferably has at least two windows which may be spaced about the circumference of the tube or wand. These windows may be separated by an arc in a range of about 20° to 180° about the longitudinal axis defined by the length of the tube or wand, and are preferably separated by an arc in a range of about 90° to 180°.

[0105] Several examples of different possible variations on the stapling device are shown and described below. These variations are not intended to be limiting but are merely given as illustrative examples.

[0106] FIG. 9A shows a variation of an endoscopic stapling device in the isometric view of anvil stapling device **170**. Stapling unit **172** is shown attached to the distal end of tube **174**. Within stapling unit **172** is staple enclosure **176** where staples may be loaded and vacuum ports **178** which are seen in an alternating fashion with staple slots **180**, through which the staples may be deployed. FIG. 9B shows a reverse isometric view of the device of FIG. 9A. As seen, stapling unit **172** may have septum **184** insertable into septum slot **186**, which is preferably midway between the sides of staple enclosure **176** and which may separate the interior of staple enclosure **176** into two separate chambers. Septum **184** may serve several functions, one of which may be to allow selective activation of opposing sides of vacuum ports **178** of unit **172** as tissue is selectively adhered to the device. Other functions of septum **184** are discussed below.

[0107] In operation, stapling unit **172** may be inserted trans-esophageally into a stomach and a first portion of the interior lining may be adhered to a single side of staple enclosure **176** through a vacuum created within vacuum ports **178**. The vacuum may be created in stapling unit **172** through tube **174** and activated from the proximal end of tube **174** from outside the patient's body. Once the first portion of the interior lining is adhered to one side of staple enclosure **176**, the opposite set of vacuum ports **178** may be activated and unit **172** may be used to draw the first portion to an opposing second portion of the interior lining, which may then be adhered to the device such that the first portion and the second portion are preferably in apposition to each other. This action preferably forms the modified lumen **26** of FIGS. 8A and 8B. As the tissue is held to unit **172**, septum **184** may be withdrawn from septum slot **186** by introduced forceps through, e.g., an endoscopic or through an integral actuator, to form a single chamber within staple enclosure **176**. Removal of septum **184** may then bring the first and second portions of tissue into contact apposition. The side surfaces **188** of septum **184** may incorporate a cutting, abrading, scoring, heating, freezing, chemically damaging, or some other damaging surface to tissue. Such a surface **188** may damage the interior lining contacting each other upon

removal of septum **184** as surface **188** slides past. This damage may encourage a more vigorous healing response and a more permanent fixation between the damaged tissue once stapled or affixed together.

[0108] After removal of septum **184**, the staples loaded within staple enclosure **176** may be fired through staple slots **180** to affix the tissue. As the staples are fired, anvil **182** may be used as an anvil to secure the staples to the tissue, thereby resulting in the modified lumen **26** as shown in FIG. 8B. The length of stapling device **170** may be made according to the desired junction length and the size of the patient's stomach. This particular variation may be withdrawn from the area after the stapling procedure by first pushing the stapling device **170** past the resulting staple line.

[0109] FIG. 10 shows an isometric view of another variation in box stapling device **190**. Stapling unit **192** is shown as being attached in fluid communication to vacuum tube **193**. Stapling device **190** may be inserted and operated in the same manner as device **170** described above. Stapling unit **192** may have vacuum ports **194** activated selectively on either side of septum **196** as described above. The tips of staples **198** are shown partially deployed for illustration purposes, but are preferably not deployed until septum **196** is first retracted preferably in the direction as indicated. Septum **196** may also be configured to damage the contacting tissue upon septum **196** withdrawal in the same manner as described above. Stapling device **190** may be easily applied and removed after staples **198** have been deployed.

[0110] FIG. 11A shows an assembly isometric view of another variation in stapling device **200**. This variation **200** shows curved tube **202** which may have lumen **204** house staples **206** as well as act as a combination vacuum and staple slot **216**. Tube **202** may be shaped in a variety of ways but is shown here as a C-shaped or U-shaped tube with first channel **210'** and second channel **210''**, for adhering the two apposed portions of tissue, preferably separated by removable septum **212**. With this variation **200**, tissue may be adhered within the channels **210'**, **210''** through vacuum/staple slot **216** and once positioned, staples **206** may be deployed while septum **212** is removed simultaneously by the use of curved wedge **218**. In operation, curved wedge **218** may be drawn within lumen **204** from the tube **202** distal end to the proximal end by, e.g., a pull-wire attached to wedge **218**. As wedge **218** is advanced proximally, wedge **218** would preferably force pivot **208** of staple **206** against contact edge **214** of septum **212**. As wedge **218** is advanced further proximally, urging end **220** may then urge the curved ends of staple **206** to rotate about pivot **208** and deploy through slot **216**. While staple **206** is deploying, notch **222**, preferably located at a distal end of wedge **218**, may engage contact edge **214** and begin to slide septum **212** simultaneously towards the proximal end of tube **202**. FIG. 11B shows a side view of stapling device **200** of FIG. 11A. As seen, curved wedge **218** preferably contacts septum **212** via notch **222** and pushes while simultaneously urging staple **206** to deploy. The figures show a single staple **206** for illustrative purposes only and any plurality of staples **206** may be used in practice depending upon the desired results.

[0111] FIG. 12A shows an isometric view of yet another variation in stapling device **230**. This variation may omit a removable septum. Curved tube **232** is preferably curved in this variation in a crescent shape forming contact channel

234. Within contact channel **234**, a number of vacuum ports **236** and staple slots **238** may be defined in an alternating pattern, as shown. A possible W-shaped staple **240** preferably having pivot **242** at the staple **240** midpoint is shown outside of tube **232** for illustrative purposes in a possible orientation for insertion within staple slots **238**. FIG. 12B shows cross section **12B-12B** from FIG. 12A. As seen, tube **232** defines lumen **244**, which preferably runs the length of tube **232**, and translating wedge **246** which is preferably slidably disposed within lumen **244**. As seen in FIGS. 12B and 12C, which is a side view of the interior of tube **232**, wedge **246** may be translated by pull-wire **248**. Pull-wire **248**, which may be made of any high-strength material, e.g., stainless steel, nitinol, nylon, polymers, etc., may be manipulated by a physician from the proximal end of tube **232** from outside of the patient's body. Like the device **200** of FIGS. 11A and 11B, once vacuum ports **236** have acquired the interior tissue lining to be approximated, translating wedge **246** may be advanced proximally. Advancing wedge **246** may urge staples **240** to deploy through staple slots **238** sequentially as shown to hold the tissue and form the desired lumen.

[0112] An example of deployment for any of the stapling devices described above is shown in FIG. 13. As shown, stomach **250** with the wall partially cut out is seen with stapling device **252** inserted within. Stapling device **252** is shown merely as an example of insertion and could comprise any of the devices described herein. Device **252**, which is preferably advanced trans-orally into stomach **250** and through esophagus **256**, is preferably located at the distal end of delivery/vacuum tube **254**. Once inserted, device **252** may be located by the assistance of the lesser curvature **258** of stomach **250**. Also shown are vacuum/staple ports **260**, which may be any of the configurations as described herein. In a preferable variation, stapling device **252** may be configured to produce a staple line or junction following the lesser curvature beginning from cardiac notch **264** down towards pylorus **262**. Accordingly, device **252** may have the length and vacuum/staple ports **260** configured such that the distal end of device **252** points towards pylorus **262**.

[0113] FIG. 14 shows stapling device **270** in a slightly different configuration for the treatment of other gastrointestinal diseases such as gastroesophageal reflux disease (GERD), as discussed above. The stomach **250** of FIG. 13 is shown, but for the treatment of GERD, stapling device **270** may be slightly modified such that the device **270** and vacuum/staple ports **272** may be straight or flared away from, rather than towards, lesser curvature **258** and pylorus **262** as described above. As such, vacuum/staple ports **272** would preferably produce a staple line or junction beginning from cardiac notch **264** and then flares away from lesser curvature **258** and pylorus **262**. Device **270** may be any of the devices described and operated herein, but for the flared modification. Likewise, any of the devices described herein may be used for the treatment of GERD by simply angling the device to produce a flared staple line. Alternatively, a simple non-flared staple line may also suffice for treating GERD. The staple line may act as a Heimlich valve which preferably closes down in response to pressure exerted from the greater or main lumen. Moreover, the smaller volume of the modified lumen in-line with esophagus **256** may provide a smaller volume of acid available for esophageal reflux.

[0114] An isometric view of a single channel vacuum device variation is shown in FIG. 15A in approximating device 280. Tube 282 is preferably a tubular device which may be inserted into a stomach through the esophagus of a patient. A lumen 284 may run through tube 282 from a proximal end to the distal end of tube 282. At the distal end, two or more windows or slots 286 are preferably defined opposite of one another, as shown. The lengths and widths of slots 286 may vary and is preferably long enough to approximate the desired length of the boundary or junction line of the modified lumen; likewise, the width is preferably wide enough to accommodate at least two layers of the stomach interior lining. Approximating clip 288 is shown having at least two piercing ends 290 and may be loaded into tube lumen 284 from either the proximal end or distal end of tube 282 preferably prior to inserting the device 280 into the patient. Clip 288 is preferably made of a biocompatible material as described above. Biodegradable plug 292 may be placed into the distal end of tube 282 prior to insertion into the patient and is preferably made of a biocompatible biodegradable material, e.g., biodegradable polymers such as polylactide, polyglycolide, and their copolymers. Plug 292 may be alternatively made from a non-biodegradable material and may simply pass after the procedure. Plug 292 may aid in maintaining a vacuum seal through slots 286 during the approximation procedure, as described below.

[0115] FIG. 15B shows an end view from section 15B-15B from FIG. 15A of tube 282 in operation. As shown, opposing portions of stomach interior lining 294 may be drawn into lumen 284 through opposing slots 286 by creating a vacuum within lumen 284. Approximating clip 288 may be urged distally through tube 282 such that each of ends 290 may be drawn through a corresponding slot 286 over and/or pierced through lining 294 within lumen 284. As lining 294 is approximated within lumen 284, biodegradable plug 292 may become invaginated within lining 294. Accordingly, as clip 288 and ends 290 are positioned over lining 294, tube 282 may be withdrawn from the area while clip 288 preferably slides through the distal end of tube 282 leaving the approximated interior lining 294 held in position by ends 290, as seen in FIG. 15D. Removal of tube 282 may urge plug 292 to slide off the distal end of tube 282 and remain within the newly formed lumen to become degraded over time or to pass through the patient's system.

[0116] FIG. 15E shows the device of FIG. 15A, but in this variation, clip 288 may be replaced by screw 289, which is preferably in the shape of a helix or coil having a tapering width or diameter. The first few turns or coils of screw 289 may have the same or similar diameter than the remaining tapering coils; this may enable piercing end 291 to engage interior 294 and may also allow screw 289 to be advanced at the desired orientation through the tissue. Screw 289 preferably maintains a parallel orientation with tube 282 during delivery into the tissue, i.e., a longitudinal axis defined by screw 289 is preferably parallel, or close to parallel, with the longitudinal axis defined by tube 282. Moreover, the outer diameter of the first few turns or coils are preferably the same diameter, or slightly less than, the inner diameter of tube 282. This may further enable screw 289 to be advanced through lumen 284 at the proper orientation prior to engaging interior 294.

[0117] As described above for the device of FIGS. 15A to 15D, opposing portions of stomach interior lining 294 may

be drawn into lumen 284 through opposing slots 286 by creating a vacuum within lumen 284, as shown in FIG. 15F. Screw 289 may then be urged through lumen 284 and rotated in the direction of the arrow shown until piercing end 291 engages the invaginated lining 294. Piercing end 291 preferably is sharp and needle-like to enable piercing through multiple layers of lining 294. As screw 289 is further rotated, it may be further advanced distally through the remaining portion of invaginated lining 294. The tapering diameter and decreasing width may also begin to further approximate the opposing edges of lining 294 towards one another, as shown in FIG. 15G. Finally, as seen in FIG. 15H, further advancement of screw 289 preferably draws the opposing surfaces into contact with one another. Tube 282 may then be removed, as described above. Although the fixation of one screw 289 is described, multiple screws 289 may be fastened one after another to form a continuous fixation line.

[0118] Screw 289 may be made of a bioabsorbable or biocompatible material, as described herein such as a polymer or superelastic alloy, and may be integrally formed with barbs or whisker-like filaments protruding along its length to help prevent screw 289 from backing out once it has been engaged within the lining 294. An example of a spiraling suturing needle or screw which may be used in this variation is shown and described in U.S. Pat. No. 5,330,503 to Yoon, which is incorporated herein by reference in its entirety. Another example of a helical fastener or screw and applicator which may be used in this or another variation is shown and described in U.S. Pat. No. 5,582,616 to Bolduc et al., which is also incorporated herein by reference in its entirety. Other examples of helical fasteners or screws and applicators are also shown in U.S. Pat. No. 5,810,882; U.S. Pat. No. 5,824,008; and U.S. Pat. No. 5,964,772; all to Bolduc et al., each of which is incorporated herein by reference in their entirety.

[0119] Gastric Reduction Tools and Methods Using Rotatable Devices

[0120] Aside from endoscopically applied stapling and clip devices, rotating and rotatable probes may also be used to form a modified smaller lumen within a main lumen. Such probes generally may be inserted into a stomach endoscopically and may engage a portion of the interior lining of the stomach and may then be rotated to roll the engaged portion of the stomach wall around the probe itself to bring the wall in apposition with another portion of the stomach wall. Such rotating probes may be used to create a blind-ended pouch of stomach within the main stomach lumen, or as with the other devices, may be used to create a smaller pouch exiting into the pylorus. Once the roll of stomach wall is brought into apposition, a row or a plurality of fasteners, e.g., staples, blind staples, clips, tags, adhesives, etc., may be used to maintain the stomach. The tubes themselves may be made of any variety of biocompatible materials which preferably have sufficient strength to undergo a torsional load, e.g., stainless steel, nickel, platinum, etc.

[0121] An example of a stomach modified by such a rotating probe or device is shown in FIG. 16A. Main pouch 300 is seen with modified pouch 302 formed along the lesser curvature of the stomach and delineated by junction 304. This example shows modified pouch 302 extending from esophagus 306 and terminating in pouch opening 308 proximally of pylorus 310. Pouch opening 308 may also be made to terminate at pylorus 310.

[0122] FIG. 16B shows a superior view from cross section 16B-16B from FIG. 16A of one variation on producing modified pouch 302 having modified lumen 314 from main pouch 300 having main lumen 312 where junction 304 may be formed by rotating the stomach upon itself. FIG. 16C shows an alternative superior view from cross section 16B-16B from FIG. 16A where modified pouch 302' having modified lumen 314' may be formed from main pouch 300' having main lumen 312'. In this particular variation, junction 304' may be formed by taking apposed sides of the interior stomach lining near the lesser curvature and approximating them to form modified lumen 314'.

[0123] Several examples of different possible variations on the rotating probe or device are shown and described below. These variations are not intended to be limiting but are merely given as illustrative examples.

[0124] FIG. 17A shows vacuum tube 320 which may have an elongate tubular body. Tube 320 may be inserted into a patient's stomach transesophageally via, e.g., an endoscope. Accordingly, distal end 322 is preferably rounded or gently tapered to be atraumatic to the patient. An opening or window 324 may be defined in the wall of tube 320 near distal end 322 and as seen in FIG. 17B, opening 324 is preferably in communication with lumen 326, which may run throughout tube 320. The geometry of opening 324 is preferably large enough to accommodate the invagination of tissue from the interior stomach lining by a vacuum created within lumen 326 and opening 324. The vacuum may be activated by the physician from a proximal end of tube 320 from outside of the patient. Once tissue is invaginated within window 324, a fastening member may be inserted and deployed to secure the interior stomach lining thereby reducing its overall volume, as described in further detail below. As shown in FIG. 17B, which is cross section 17B-17B from FIG. 17A, tube 320 preferably has a diameter and cross section which may approximate a final geometry of the newly created lumen within the stomach.

[0125] FIG. 18A shows an isometric view of another variation in counter-rotating tube 330. Counter-rotating tube 330 may have a gently tapered distal end 332 with an opening 334 defined in the tube wall near distal end 332. Preferably contained within tube 330 is an additional inner tube 336, which may be geometrically similar to tube 330 but with a diameter small enough to allow free rotation about the longitudinal axis preferably shared by both tubes 330 and 336. Inner tube 336 likewise may have inner opening 338, which may allow communication between lumen 340 and openings 334 and 338. As above, a vacuum may be activated from a proximal end of tube 330 to draw tissue from the interior stomach lining through lumen 340 and into openings 334 and 338 when they are aligned. As shown in FIG. 18B, which is cross section 18B-18B from FIG. 18A, once the tissue has become invaginated within openings 334, 338, inner tube 336 may be rotated to effectively pinch and firmly hold the tissue in place, as shown in FIG. 18B. The addition of the pinching action in addition to the vacuum may aid in holding the tissue, thereby aiding in the rotation of both tube 330 and inner tube 336 when forming the modified lumen. Both tubes 330 and 336 may be manipulated and rotated from a proximal end of the tubes from outside of the patient.

[0126] FIG. 19A shows an isometric view of another variation in barbed tube 350. Tube 350 may be similar to

vacuum tube 320 described above. Distal end 352 is preferably tapered and opening 354 may be defined in the wall of tube 350 near distal end 352. Additionally, at least one and preferably several attachment points 356, e.g., tines, barbs, or hooks, may be defined along at least a single edge around opening 354. Attachment points 356 are preferably defined along the leading edge of opening 354 for rotation of tube 350. FIG. 19B, which is cross section 19B-19B from FIG. 19A, shows opening 354 preferably in communication with lumen 358 and a preferred orientation of attachment point 356.

[0127] FIG. 20A shows an isometric view of yet another variation in split tube 360. Split tube 360 may be formed of at least two splittable halves, e.g., first half 364 and second half 366, which may be joined together longitudinally along split 370. When first half 364 and second half 366 are joined together, split tube 360 preferably forms a tapered distal end 362. Split tube 360 may also define a lumen 372 which may run throughout the length of split tube 360. This variation may also comprise at least one and preferably several attachment points 368 on each of first half 364 and second half 366. As shown in the figure, first half 364 may have a row of attachment points 368 preferably aligned along a portion of split 370 and second half 366 may likewise have a row of attachment points 368 juxtaposed and preferably mirroring those located on first half 364. Attachment points 368 may be of any type described above and the number and positioning of attachment points 368 may depend upon the desired length of the resulting junction formed upon rolling the stomach. FIG. 20B, which is cross section 20B-20B from FIG. 20A, shows split 370 and an example of the juxtaposed relationship of attachment points 368.

[0128] FIG. 21 shows an example of a rotatable probe device during insertion into stomach 380. As seen, tube 384 may be inserted into stomach 380 via esophagus 382, preferably endoscopically. Tube 384 may be any of the devices described above and is shown generally as an example of how such devices may be inserted into an organ, e.g., stomach 380. As tube 384 is inserted, it may engage a portion of the interior of stomach 380, preferably along lesser curvature 386. The engagement may be accomplished by any of the methods described herein, e.g., attachment points partially piercing the stomach lining, a vacuum adhering a portion of the lining, etc. Once engaged, tube 384 may then be rotated to roll the engaged portion of the stomach wall around the probe itself to bring the wall in apposition with another portion of the stomach wall.

[0129] FIG. 22A shows a variation on partial cross section 22/23-22/23 from FIG. 21 with tube 350 from FIGS. 19A and 19B in a preferred operation. As shown, interior lining 390 may be adhered to tube 350 via a vacuum created in opening 354 through lumen 358 and/or via attachment points 356 which may partially pierce lining 390, as described above. The location for adhering tube 350 may also be determined or aided by the use of marking device 40, as described above. Once the desired location of interior lining 390 has been established, tube 350 may be rotated about its longitudinal axis, following the arrow as shown, by at least about 180° and preferably at least about 360°. Lining 390 is preferably rotated until the adhered portion contacts a second portion of lining 390 to result in the modified lumen 314 of FIG. 22B, also shown in FIG. 16B. Once modified lumen 314 has been formed, fasteners may be fired

or deployed through opening 354 or via a separate endoscopic stapling device at location 392 to secure and maintain modified lumen 314. Fasteners may comprise any of the fasteners as described herein, e.g., staples. Once modified lumen 314 has been secured, tube 350 may then be removed. FIG. 16B shows newly created modified pouch 302 with modified lumen 314 and, as seen, interior lining 390 also forms the interior surface defining modified lumen 314.

[0130] FIGS. 23A to 23D show another variation on partial cross section 22/23-22/23 from FIG. 21 with split tube 360 from FIGS. 20A and 20B. Split tube 360 may be inserted into the stomach either as separate halves 364, 366 individually or as a whole tube which may then be split while in the stomach. Once separated, first half 364 and second half 366 may be engaged to interior lining 390 by attachment points 368 at a slight distance from one another. The separation distance may be determined by the desired resulting size of the lumen. Alternatively, the separation distance may be determined or aided by the use of marking device 40, as described above.

[0131] Once first half 364 and second half 366 have engaged interior lining 390, as shown in FIG. 23A, each of free ends 394 of halves 364, 366 may then be rotated in the direction of the arrow, as shown. Free ends 394 may be configured to simply contact each other or to interlock with each other and rotate about a hinge or pivot. As first half 364 and second half 366 continue to be rotated, FIGS. 23B and 23C show the progression of lumen formation as attachment points 368 draw around and towards one another. Finally in FIG. 23D, as split tube 360 is preferably formed again, modified lumen 314' may be formed, as also shown in FIG. 16C, to then be secured or maintained preferably by fasteners, e.g., staples, which may be deployed through junction 304'.

[0132] A further variation on a rotating device is shown in the isometric view of dual tube device 400 shown in FIG. 24A. Dual tube device 400 may have at least two elongate members, first member 402 and second member 404, which may be rotatably attached to controlling device 406 and may be parallel to each other. The members 402, 404 are preferably counter-rotating and may be rotated by a rotation control 408, which is preferably located on controlling device 406. First member 402 may have first distal end 410 offset slightly from the longitudinal axis of first member 402 by first bend 412. First opening 414 is also preferably defined in the wall of first member 402 proximally of first distal end 410. Second member 404 is preferably similar to first member 402 and may have second distal end 416 offset slightly from the longitudinal axis of second member 404 by second bend 418. Near second distal end 416, second opening 420 may be defined in the wall of second member 404.

[0133] FIG. 24B shows end view 24B-24B from FIG. 24A. Distal ends 410, 416 are seen as preferably being parallel and mirror images of one another. Also, the preferable counter-rotating action may be seen by the directional arrows. FIG. 24C shows cross section 24C-24C from FIG. 24A. As shown, the relationship between first and second opening 414, 420, respectively, and first and second lumen 422, 424, respectively, may be seen in the figure. Lumens 422, 424 preferably run through the length of members 402, 404, respectively, and are in communication with openings

414, 420. A vacuum may be created in openings 414, 420 through lumens 422, 424, respectively, from the controlling device 406. In operation, members 402, 404 may be inserted trans-esophageally into a patient's stomach. A vacuum may then be created in first and second openings 414, 420 to engage a portion of the stomach interior lining. Once engaged, a modified pouch may be created from the interior lining in much the same manner as described for FIGS. 23A to 23D, except the individual counter-rotating members 402, 404 do not form a split tube. The operation of the vacuum application and counter-rotation may be controlled through controlling device 406 which is preferably located outside the patient's body.

[0134] FIG. 25A shows yet another variation in vacuum device 432 shown inserted into stomach 430. Vacuum device 432 may be an endoscopic device inserted trans-esophageally into stomach 430 through esophagus 434. Device 432 may have vacuum member 438 and at least two grasping members 440, preferably disposed on either side of vacuum member 438. Once device 432 has been introduced into stomach 430, vacuum member 438 may be steered towards a desired area of interior lining 442, as seen in FIG. 25B which is a cross section view of device 432 attached to stomach interior lining 442. The desirable area of interior lining 442 may be located along greater curvature 436 or alternatively along lesser curvature 444, depending upon the desired results. In position, a vacuum may be activated in member 438 to draw a portion of interior lining 442 preferably between grasping members 440. As lining 442 is adhered to vacuum member 438, grasping members 440 may be used to pinch and grasp the drawn portion of lining 442. Then, device 432 may be rotated in the direction of the arrow indicated in FIG. 25C to result in the formation of a modified lumen. Afterwards, grasping members 440 may be locked in place, disengaged from device 432, and left as an implant. Alternatively, lining 442 may be fastened to maintain the created lumen by any of the methods described herein and grasping members 440, along with the rest of device 432, may be removed from stomach 430.

[0135] Gastric Reduction Tools and Methods Using Volume Reduction Devices

[0136] Aside from the use of rotating and rotatable probes, gastric volume reduction devices may also be used as part of the present invention. Such volume reduction devices generally may be inserted into a stomach trans-esophageally through the use of, e.g., an endoscope. The reduction device may be used to draw or engage a portion of the interior lining of the stomach; the drawn or engaged portion may then be eventually removed, either actively or through natural processes.

[0137] Several examples of different possible variations on the gastric volume reduction devices are shown and described below. These variations are not intended to be limiting but are merely given as illustrative examples.

[0138] FIG. 26 shows an isometric view of a variation on the gastric volume reduction device in concentric tube device 450. Device 450 may have inner tube 452 defining lumen 454, which preferably runs throughout inner tube 452. Pusher sleeve 456 may be disposed concentrically over inner tube 452 such that pusher sleeve 456 may be allowed to slide freely along inner tube 452. Pusher sleeve 456 is also preferably disposed over inner tube 452 such that the distal

end of inner tube 452 is open to allow ring 458 to be rolled or stretched onto the distal end. Ring 458 is preferably made of an elastic type material which would allow ring 458 to elastically cinch onto inner tube 452.

[0139] During use, FIG. 27A shows a view of concentric tube device 450 within stomach 460 preferably inserted through esophagus 462. The distal end of device 450, particularly inner tube 452, may be brought into position near a location of interior surface 464 where tissue may be desirably removed. As shown in FIG. 27B, once device 450 is in place, a vacuum may be actuated within lumen 454. The vacuum may then draw a portion of withdrawn lining tissue 466 up into lumen 454, as seen in the cross section of device 450. While lining tissue 466 is held within lumen 454, pusher sleeve 456 may be pushed or urged distally along inner tube 452. As pusher sleeve 456 advances, it may also push or urge elastic ring 458 distally along inner tube 452 until ring 458 is pushed entirely off the distal end of inner tube 452 and onto a portion of lining tissue 466, as seen in FIG. 27C. Device 450 may then be removed from stomach 460 after ceasing the vacuum, thereby leaving lining tissue 466 with elastic ring 458. After time, as seen in FIG. 27D, pressure necrosis may cause lining tissue 466 and ring 458 to simply fall off from the rest of interior surface 464 to be passed normally through the rest of the patient's body. The action of drawing up and removing a portion of interior surface 464 may effectively reduce the overall volume of stomach 460, thereby reducing the available volume for the ingestion of foods. As such, this procedure may be repeated several times either sequentially or simultaneously until the overall volume of stomach 460 is reduced to a desirable volume depending upon the desired results.

[0140] FIG. 28 shows another variation on the gastric volume reduction device. As shown, an endoscope 474 preferably having grasping device 476, e.g., biopsy forceps, may be inserted into stomach 472. A ligating apparatus, e.g., ring stapler, zip tie, etc., either as part of endoscope 474 or as a separately introduced ligation device 478, is preferably also introduced within stomach 472. Forceps 476 and ligation device 478 may be used in conjunction with one another by, e.g., having forceps 476 grasp withdrawn tissue 480 and then having ligation device 478 tie or ligate tissue 480. Forceps 476 may then be used to excise and remove withdrawn tissue 480 above ties 482 to reduce the overall stomach volume. An example of a jaw structure which may be utilized is shown and described in U.S. Pat. No. 5,749,893 to Vidal et al., which is incorporated herein by reference in its entirety. Alternatively, ligated withdrawn tissue 480 may be left attached to stomach 470 to be removed naturally by pressure necrosis. Several excisions may be performed in reducing stomach volume from, e.g., stomach 472 (as shown by the dashed lines) down to a final reduced stomach 470.

[0141] FIG. 29A shows yet another variation with tractive rollers 490. This device may have at least two rigid rollers 492, which are preferably elongated, connected to one another preferably at both ends by, e.g., elastic members 494. The connection of rollers 492 may create channel 496 therebetween through which tissue may be drawn. FIG. 29B shows rollers 492 with a portion of stomach interior surface 498 being drawn through channel 496 by a grasping device, e.g., forceps 500. Meanwhile, rollers 492 may be maintained within the stomach by, e.g., retaining forceps 502, which may be used to hold rollers 492 relative to interior surface

498. Elastic members 494 may pinch rollers 492 together, thereby creating a zone of pressure necrosis in withdrawn interior surface 498. Also, as interior surface 498 is drawn up through channel 496, rollers 492 may contain a ratcheting device within to prevent surface 498 from rolling out back through channel 496. Once the desired amount of surface 498 has been drawn, it may either be excised or simply left to be removed naturally by necrosis. FIG. 29C shows an alternative variation with ratcheted rollers 504. Ratcheted rollers 504 may be operated in the same manner as described for rollers 492 but they preferably have a tractive surface to enhance traction between the tissue and the rollers 504. Torquing device 506 may be used with ratcheted rollers 504 and may be introduced into the stomach endoscopically to mesh with one of rollers 504 for the purpose of causing it to rotate. Moreover, either rollers 492 or ratcheted rollers 504 may be used simply to gather stomach surface tissue to allow for fastening, e.g., suturing, stapling, etc.

[0142] Pyloroplasty Tools and Methods

[0143] Creating a smaller gastric pouch within the stomach may be accomplished by a variety of methods, as described above. To aid in the overall effect for the treatment of obesity, a pyloroplasty procedure may also be performed to enhance treatment. The pyloroplasty may be performed prior to (preferable), in conjunction with, or following the gastric reduction procedure. A pyloroplasty procedure typically results in the pyloric sphincter being rendered incompetent. However, in the case of treatments for GERD using the devices and methods described above, the pyloroplasty procedure as described herein may be omitted. Conventional pyloroplasty procedures may typically be performed surgically or through the use of standard peripheral angioplasty balloons, e.g., in the 7 mm range. However, in order to render a relatively healthy and normal pylorus permanently incompetent, a more aggressive procedure may be needed.

[0144] To accomplish this generally, a pyloroplasty device may be passed endoscopically through the esophagus, into the stomach, and preferably into position in or across the pylorus. Energy or a stimulus is then preferably applied to the pylorus to render it incompetent. Energy may be in the form of, e.g., heat, electrical, chemical, RF, etc., or a combination. Examples of chemical energy stimulus may comprise alcohol and sotrodecol. The stimulus may be in the form of, e.g., dilatation, cutting, ablation, viral, etc., or a combination. An example of a viral or chemical stimulus may be, e.g., a poison such as the botulinum toxin type A virus (Botox). An example of a method of use for Botox is described in U.S. Pat. No. 5,437,291 to Pasricha et al., which is incorporated herein by reference in its entirety. An incompetent pylorus may allow stomach contents to drain directly into the proximal duodenum with minimal resistance. Moreover, some of the mentioned pyloroplasty treatments may be selected or designed to last only for a specific time period, e.g., a week or several months, etc. For instance, the effects of simple dilatation or the injection of Botox may be designed to render the pylorus incompetent for only a few months, which may be a desirable time period for the patient to obtain the desired results of the procedure.

[0145] Several examples of different possible variations on pyloroplasty devices are shown and described below. These variations are not intended to be limiting but are merely given as illustrative examples.

[0146] FIG. 30 shows an isometric view of one variation of a dilatation device in balloon device 510 which may have angioplasty balloon 512 located near or at the distal end of catheter 514. Angioplasty balloon 512 may be used alone to simply dilate the pylorus. Alternatively, exterior balloon surface 516 may have at least one and preferably several stimulating members 518 disposed about surface 516. Stimulating members 518 are shown in the figure as cutting blades or wires, but alternatively, they may include electrodes, cryogenic dispensing probes or members, chemical dispensing probes, etc. Moreover, balloon 512 may alternatively be a dilation wire basket similarly disposed with stimulating members 518.

[0147] FIG. 31 shows an isometric view of another variation in probe device 520. Device 520 may have catheter or delivery member 524 with, e.g., probes 526, which may extend from distal end 522. Although three probes 526 are shown in the figure, at least one and up to several probes of varying thickness and lengths may be used. Probes 526 may be retractable so that during delivery through, e.g., the esophagus or stomach, probes 526 may be withdrawn within distal end 522 and then extended when treating the pylorus. Probes 526 may be electrically connected to a voltage or power source located outside the patient's body to deliver electrical, RF, or heat energy to the pylorus. Alternatively, they may be configured like a needle to deliver chemical or biological stimuli to render the pylorus incompetent. For example, probes 526 may be used to inject chemicals, e.g., alcohol, sotrodecoll, or other ablative chemicals, or biological stimuli, e.g., Botox virus or some other incapacitating virus, into the pylorus. Such stimulants may be carried within distal end 522, delivery catheter 524, or they may also be delivered from the proximal end of catheter 524 and injected through to probes 526.

[0148] Other variations which may be used for the pyloroplasty procedure are shown in FIGS. 32A and 32B. FIG. 32A shows sphincterotome arm 530 having a distal end 532. Arm 530 may be bent as shown to allow cutting member 534 to be drawn between distal end 532 and a location proximal of distal end 532 along arm 530. Another variation is seen in FIG. 32B where delivery member 536 may have an arcuate support member 538' to support cutting member 540. The variations shown in FIGS. 32A and 32B may be delivered via a catheter or endoscope trans-esophageally and through the stomach to the pylorus where either cutting member 534, 540 may be used to cut or saw into the tissue in or around the pylorus to render it incompetent. These particular variations of sphincterotomes shown in FIGS. 32A and 32B may be manufactured by Medi-Globe Corporation, located in Tempe, Ariz.

[0149] FIG. 33 shows stomach 550 with a distal portion of the wall of the lesser curvature removed for clarity. Device 520 may be delivered through esophagus 552 to a location proximal of pylorus 558, e.g., first position 554. If probes 526 were retracted during delivery, they may then be extended, as shown. Distal end 522 of device 520 may be advanced to, e.g., second position 556, such that probes 526 may pierce pylorus 558 to deliver the stimulus.

[0150] FIG. 34A shows an isometric view of another variation with combination device 560. Device 560 may have housing 562 on the distal end of delivery catheter or endoscope 564. Housing 562 defines notch 566 which may

be oriented perpendicularly relative to the longitudinal axis defined by endoscope 564. Notch 566 preferably has a geometry large enough to accommodate part of pylorus 558 and housing 562 may be tapered at its distal end to allow for easy insertion into the pylorus 558 during the procedure. Within notch 566 may be cutting blade 568 and on either side of blade 568 may be fasteners 570, e.g., individual anchors, staples, etc. In operation, FIG. 34B shows housing 562 and endoscope 564 delivered through esophagus 552. The wall of stomach 550 is partially cut away for clarity. Housing 562 may be inserted into pylorus 558, then notch 566 is preferably aligned such that part of the pyloral sphincter lies within notch 566. Alternatively, the pyloral tissue may also be drawn into notch 566 via a vacuum or grasping member. Once the pyloral tissue is within notch 566, cutting blade 568 may be actuated to traverse notch 566 and sever part of the tissue of pylorus 558. Fasteners 570 may then be deployed on either side of incision 572 to affix the incised tissue. The number of incisions 572 may vary depending upon the desired degree of pyloric disablement. Alternatively, an inflatable balloon may be attached on the back of notch 566 and inflated to push housing 562 into apposition with pylorus 558 and cause invagination of the tissue into notch 566.

[0151] Anastomosis Tools and Methods

[0152] In addition to the tools and methods described above for gastric reduction and pyloroplasty procedures, an additional anastomosis gastric bypass procedure may also be performed to further enhance treatment. The anastomosis procedure may be performed preferably prior to, in conjunction with, or following the gastric reduction and pyloroplasty (if performed at all) procedures. In the case of treatments for GERD using the devices and methods described above, the anastomosis procedure as described herein may be omitted. The procedure generally involves endoscopically or laparoscopically creating a side-to-side anastomosis preferably from within the stomach and bowel and within the digestive tract. This procedure may be similar to the Roux-en-Y gastric bypass (RYGB) procedure but with minimal trauma. This procedure may also effectively bypass food from the stomach, past a proximal portion of the bowel, and preferably directly into a lower portion of the bowel. This bypassed portion may be considered a malabsorption zone.

[0153] A representative and normal gastro-intestinal system of a person is shown in FIG. 35 for comparison. Stomach 580 is shown with pyloric sphincter 582 near gallbladder 584 and attached to the proximal section of duodenum 586. The distal section of duodenum 586 is attached to the proximal section of jejunum 588, the distal section of which is further attached to the proximal section of ileum 590. Ileum 590 is then attached to ascending colon 592, which continues through to the transverse colon (which has been removed for clarity), and then to descending colon 594 and finally to rectum 596.

[0154] A gastrointestinal system which may be modified by a preferable anastomosis procedure is shown in FIG. 36. Stomach 600 is shown in this variation as having been modified by creating modified pouch 602, which may be created by any of the methods and tools as described above. Esophagus 603 is preferably connected to a proximal end of pouch 602. As described above, the distal end of pouch 602

may be connected directly to pylorus 604 or alternatively, may be a blind-ended pouch and pylorus 604 is connected to the proximal end of duodenum 606. A first anastomosis 608 may be created preferably between modified pouch 602 and a section of digestive tract either from the distal duodenum 606 or proximal jejunum 610. First anastomosis 608 may be located in a range from about 20 to 50 cm from pylorus 604. A second anastomosis 614 may be created preferably between a section of duodenum 606 and a section of ileum 612. The second anastomosis 614 may be located in a range from about 15 to 55 cm from pylorus 604 or about 150 to 200 cm down along the length of the small intestines from pylorus 604. This procedure may allow for drainage of secretions created by stomach 600 to pass through pylorus 604 and secretions of bile and chyme from the pancreas and gallbladder 618 to pass through biliary duct 620 partly through duodenum 606 and then through second anastomosis 614 and directly into distal ileum 616 and out of the body. The bypassed stomach 600, pylorus 604, and proximal duodenum 606 may act as a malabsorption zone because sugars and fats which might normally be mostly absorbed in this zone may now be directly passed into the distal duodenum 606 or proximal jejunum 610.

[0155] During the anastomosis procedure, both first and second anastomoses 608, 614, respectively, may be created first. Duodenum 606 may then be closed off between the two anastomoses 608, 614. Then, depending upon the length and size of the resulting modified stomach 602, pylorus 604 may be closed off or left open, depending upon the desired result and which of procedures and tools are implemented. Finally, modified pouch 602 may be created after the anastomoses procedures. Alternatively, modified pouch 602 may be created prior to the anastomoses procedures, again depending upon the desired result and which of procedures and tools are implemented. If modified pouch 602 were created first, then the anastomoses procedure may be reversed to essentially end with the same result.

[0156] A conventional RYGB procedure is generally performed through a 6-8 inch incision extending from the end of the breastbone to just above the navel. However, the procedure described above may be performed entirely endoscopically or laparoscopically. FIG. 37 shows an isometric view of an assembly which may be utilized to achieve part of the procedure. Deployment device 630 may have anastomosis assembly 632 preferably connected by steerable length 634 to manipulation handle 636. Assembly 632 may be steerable during insertion, preferably trans-esophageally and through the stomach, by steering grip 638 which may be located on manipulation handle 636. Control by a physician or surgeon of manipulation handle 636 may be facilitated by handle 640.

[0157] Anastomosis assembly 632 may have stapler housing 644 configured to fit intimately with distal element 646 preferably by a magnetic force, the use of which is described below. Distal element 646 is preferably tapered or rounded on one side and may have a coring anvil 648 on its opposing side. Coring anvil 648 may be tapered or rounded and may fit intimately into coring mate 650 which is preferably located near or at the center of stapler housing 644. Stapler housing 644 may also house several staples loaded within staple slots 652, which may be disposed circumferentially around coring mate 650 and may be actuated from the proximal end of length 634 by staple trigger 642.

[0158] FIG. 38 shows a cross sectioned view of anastomosis assembly 632 mated with distal element 646 at first anastomosis 608 between modified pouch 602 and jejunum 610. Part of the walls of modified pouch 602 and jejunum 610 have been removed for clarity. In creating first anastomosis 608, distal element 646 may first be placed within the appropriate section of jejunum 610. This may be done by orally passing distal element 646 through the esophagus, stomach, and then through the duodenum. Distal element 646 is preferably magnetized, either by manufacturing distal element 646 from natural ferrous materials or artificially magnetizing it. Because of the magnetization, distal element 646 may be urged through the body and into place within the duodenum by, e.g., magnetic wands or magnetic pickups, which may be manipulated from outside the patient's body.

[0159] During or after placement of distal element 646, stapler housing 644, which may be attached to steerable length 634, may be introduced transesophageally into the stomach 602 and placed into position along stomach wall 660 at the desired site of first anastomosis 608. Once both stapler housing 644 and distal element 646 are in position, they may then be coupled together preferably by the magnetic force and attraction between the two. Moreover, the two may be brought into alignment either by alignment grooves (not shown) or by the mating of coring anvil 648 into coring mate 650. As the mating occurs, part of stomach wall 660 and intestinal wall 662 are preferably held or maintained between stapler housing 644 and distal element 646. To enhance the mating, fasteners may optionally be deployed from stapler housing 644 through staple slots 652 and preferably through both stomach wall 660 and intestinal wall 662 into distal element 646. FIG. 38 shows staples 667 deployed as fasteners, but they may comprise any type of mechanical fasteners as described above, as well as, e.g., grommet-type swages, snap lock fits, staples, screws, clips, and friction-fittings.

[0160] Once the fitting has been accomplished, the device may be left in apposition to maintain the position of stomach wall 660 and intestinal wall 662 for about one week. This may result in pressure necrosis of the tissue between stapler housing 644 and distal element 646 preferably causing the serosal layers of the gut to fuse, at which point the assembly may drop out and be passed, preferably leaving first anastomosis 608 behind. Alternatively, a coring device 664, which may be slidably contained within stapler housing 644, may first be advanced through the center of stapler housing 644 and both stomach wall 660 and intestinal wall 662 to create first anastomosis 608. The remaining assembly may then be left to cause the pressure necrosis and fusing of tissue, as described. Another alternative may be to use stapler housing 644 and distal element 646 as a mechanism for a conventional end-to-end anastomosis (EEA) stapler. In this case, once they are aligned, a rod may be advanced through the center of the assembly to preferably lock distal element 646 to intestinal wall 662. The rod may be drawn back, preferably pulling a distal stapler segment into stapler housing 644. This action may cause staples to fire and a circumferential blade to cut out the center of the staple ring, thereby creating an anastomosis.

[0161] To create second anastomosis 614, a similar approach may be taken as for creating first anastomosis 608. An example of another magnetic anastomosis device which may also be used in this procedure is shown and described

in U.S. Pat. No. 5,690,656 to Cope et al., which is incorporated herein by reference in its entirety. FIG. 39 shows a portion of duodenum 606 juxtaposed to a portion of ileum 612 and distal ileum 616 with part of the intestinal walls removed for clarity. In this variation, proximal element 670 may be used and is preferably a magnetized mating element for distal element 646. Distal element 646 may first be urged to the desired location preferably in ileum 612 by, e.g., magnetic wands or magnetic pickups, which may be manipulated from outside the patient's body, in the same manner as above. During or after placement of distal element 646, proximal element 670 may also be delivered or urged to the desired location in the same manner. Once both elements 646, 670 are in position, they are preferably mated together by a magnetic force. The mating may optionally be enhanced by fasteners, e.g., staples 667, to hold both elements 646, 670 in position. The intestinal wall inbetween may be cored, as described above, but it may also be simply left to undergo pressure necrosis between elements 646, 670 eventually causing the serosal layers of the gut to fuse, at which point elements 646, 670 may drop out and be passed, preferably leaving second anastomosis 614 behind.

[0162] The applications of the methods and tools discussed above are not limited to the treatment of obesity, but may include any number of further applications, e.g., GERD, which may involve manipulation of an organ interior. Modification of the above-described methods and tools for carrying out the invention, and variations of aspects of the invention that are obvious to those of skill in the art are intended to be within the scope of the claims.

We claim:

1. A system for forming a lumen from an interior of an organ, comprising a rotatable device for insertion into the organ, the device being specially adapted to adhere onto an internal surface while being rotated about a longitudinal axis defined by the device.
2. The system of claim 1, further comprising a fastening device specially adapted to fasten a rotated portion of the organ to a remaining portion of the organ.
3. The system of claim 1, wherein the device is rotated at least about 180°.
4. The system of claim 1, wherein the device is rotated at least about 360°.
5. The system of claim 1, further comprising a side-to-side anastomosis device for coupling adjacent regions of digestive tract, the anastomosis device including a distal segment and a proximal segment, the distal and the proximal segments each being adapted to align in apposition with each other while maintaining the regions of digestive tract therebetween.
6. The system of claim 2, wherein the rotated portion is mechanically fastened to the remaining portion by a fastener selected from the group consisting of staples, blind staples, clips, tags, screws, and adhesives.
7. The system of claim 1, wherein the rotatable device for insertion is inserted into the organ endoscopically via an esophageal passageway.
8. The system of claim 5, wherein the anastomosis device is inserted into the digestive tract endoscopically via an esophageal passageway.
9. The system of claim 6, wherein the anastomosis device is inserted into the digestive tract laparoscopically.

10. The system of claim 1, wherein the rotatable device for insertion includes an outer tube having a proximal end and a distal end with a lumen defined therebetween, the outer tube defining an opening proximal of the distal end.

11. The system of claim 10, wherein the internal surface within the organ is adhered to the device via a vacuum created in the opening.

12. The system of claim 10, wherein the distal end includes an enclosed, tapered end.

13. The system of claim 10, wherein the rotatable device for insertion further comprises an inner tube having a proximal end and a distal end with a lumen defined therebetween, the inner tube defining an opening proximal of the distal end.

14. The system of claim 13, wherein the opening of the inner tube is configured to align with the opening of the outer tube.

15. The system of claim 14, wherein the inner tube is counter-rotatable about the longitudinal axis with the outer tube.

16. The system of claim 10, wherein the outer tube further comprises a plurality of attachment points disposed adjacent to the opening, the internal surface within the organ adhering to the outer tube via the attachment points.

17. The system of claim 16, wherein the attachment points are selected from the group consisting of tines, barbs, and hooks.

18. The system of claim 1, wherein the rotatable device for insertion comprises a split tube having at least two halves, each of the halves being separatable along a partition defined along the tube, the split tube comprising a proximal end and a distal end with a lumen defined therebetween the split tube further comprising a plurality of attachment points disposed on each half-near the distal end.

19. The system of claim 18, wherein the attachment points are selected from the group consisting of tines, barbs, and hooks.

20. The system of claim 1, wherein the rotatable device for insertion comprises at least two adjacent tubes, each of the tubes having a proximal end and a distal end with a lumen defined therebetween, and each of the tubes defining an opening proximal of the distal end.

21. The system of claim 20, wherein each of the two adjacent tubes are adapted to counter-rotate.

22. The system of claim 20, wherein the internal surface within the organ is adhered to the device via a vacuum created in each of the openings.

23. The system of claim 1, wherein the rotatable device for insertion comprises at least one attachment tube and at least two compression members, the attachment tube being specially adapted to adhere to the internal surface within the organ, and the two compression members being adapted to compress a portion of the internal surface within the organ therebetween.

24. The system of claim 23, wherein the attachment tube defines an opening proximal of a distal end.

25. The system of claim 24, wherein the internal surface within the organ is adhered to the attachment tube via a vacuum created in the opening.

26. The system of claim 5, wherein the distal and the proximal segments are magnetically coupled together while maintaining the regions of digestive tract therebetween.

27. The system of claim 26, wherein the distal and the proximal segments are further coupled by mechanical fasteners.

28. The system of claim 27, wherein the mechanical fasteners are selected from the group consisting of grommet-type swages, snap lock fits, staples, screws, clips, and friction-fittings.

29. The system of claim 5, further comprising a deployment device comprising all elongate tubular member having a proximal end and a distal end with a lumen defined therebetween, the distal end being specially adapted to retain the proximal segment for alignment in apposition with the distal segment.

30. The system of claim 29, wherein the deployment device further comprises a coring ring in the distal end, the coring ring being adapted to pass through a center of the distal segment and the proximal segment for removal of the regions of digestive tract therebetween.

31. The system of claim 29, wherein the deployment device distal end is adapted to the manipulatable via the proximal end.

32. The system of claim 5, further comprising a locating device adapted to urge the distal segment into a predetermined location in the digestive tract.

33. The system of claim 32, wherein the locating device comprises a magnetized device selected from the group consisting of magnetic wands and magnetic pickups.

34. A method of creating a separate gastric lumen from a gastric pouch having a main lumen, comprising:

adhering a portion of interior of the gastric pouch to a device specially adapted to rotate;

rotating the device about a longitudinal axis defined by the device while maintaining adherence to the portion of interior, the device being rotated until the separate gastric lumen is defined; and

fastening the portion of interior to the gastric pouch such that the gastric lumen is maintained.

35. The method of claim 34, wherein the device comprises a tube having a proximal end and a distal end with a lumen defined therebetween, the tube defining an opening proximal of the distal end.

36. The method of claim 35, wherein the portion of interior is adhered to the device via a vacuum created in the opening.

37. The method of claim 35, wherein the portion of interior is adhered to the device via a plurality of attachment points disposed adjacent to the opening.

38. The method of claim 38, wherein the attachment points are selected from the group consisting of tines, barbs, and hooks.

39. The method of claim 34, wherein the portion of interior is fastened to the gastric pouch by a mechanical fastener selected from the group consisting of staples, blind staples, clips, tags, screws, and adhesives.

40. A system for organ volume reduction, comprising:

an elongate tubular device having a proximal end, a distal end, with a length therebetween and a lumen defined throughout the length, wherein the device is adapted for insertion into an hollow body organ, the device being further adapted to draw up a portion of an interior wall of the hollow body organ to be rotated about a longitudinal axis of the device; and

a fastening device adapted to fasten the rotated portion of the interior wall.

41. The system of claim 40, wherein the device for insertion is inserted into the hollow body organ endoscopically via an esophageal passageway.

42. The system of claim 40, wherein the lumen defines a passageway for drawing up the portion of the interior wall of the organ via a vacuum.

43. The system of claim 40, wherein the fastening device comprises an elastic band disposed about the distal end of the device, wherein the band is adapted to slide over and secure the portion of the interior wall.

44. The system of claim 40, wherein the device for insertion comprises at least two elongate members, each member having a proximal end and a distal end with a length therebetween, the members being rotatably connected at the proximal end and the distal end such that the members are biased towards each other.

45. The system of claim 44, wherein the elongate members define an uneven surface.

46. The system of claim 40, wherein the device for insertion comprises an instrument having manipulatable grasping members at the distal end for grasping the portion of the interior wall of the hollow body organ.

47. The system of claim 46, wherein the fastening device comprises a catheter having a cinching distal end adapted to cinch the portion of the interior of the organ.

48. The system of claim 40, wherein the fastening device comprises a clip adapted to be inserted into the lumen.

49. The system of claim 48, wherein the clip comprises two adjacent members attached at a proximal end, the distal ends of each member being configured to pierce tissue.

50. The system of claim 40, wherein the fastening device comprises a screw configured to be inserted into a channel defined within the device.

51. The system of claim 50, wherein the screw comprises a helical shape.

52. The system of claim 51, wherein a diameter of a distal end of the screw is larger than a diameter of a proximal end of the screw.

53. The system of claim 52, wherein a length between the distal end of the screw and the proximal end of the screw is tapered.

54. The system of claim 53, wherein the screw is configured to alternately pierce a first area of tissue and a second area of tissue while approximating the first area of tissue and the second area of tissue together.

55. The system of claim 40, further comprising a plurality of additional fastening devices, each of the fastening devices being configured to fasten the portion of interior while juxtaposed linearly to form a continuous fixation line.

56. The system of claim 40, wherein the fastening device comprises at least one staple.

57. The system of claim 40, wherein the device defines a slot along the length for drawing up the portion of the interior wall, wherein the slot is in fluid communication with the lumen.

58. The system of claim 57, further comprising at least one attachment point positioned adjacent to the channel.

59. The system of claim 40, wherein the device comprises a tubular member that is longitudinally splittable, wherein each tubular half has at least one attachment point positioned thereon such that the attachment points are in apposition to one another when the tubular member is joined together.

60. The system of claim 40, wherein the device comprises at least two elongate members, each member having a proximal end, a distal end, with a length therebetween and a lumen defined throughout each length, the members being rotatably connected at the proximal ends to a handle, each member defining a slot along each length in communication with each respective lumen, wherein each member is adapted to counter-rotate relative to one another about a longitudinal axis defined by each member.

61. The system of claim 60, wherein each member is parallel to one another.

62. The system of claim 60, further comprising a vacuum pump in fluid communication with each slot through each respective lumen.

63. A system for organ volume reduction, comprising:

at least two elongate members, each member having a proximal end, a distal end, with a length therebetween and each member being rotatably connected proximally of the distal end, wherein each member is adapted to adhere a portion of an interior wall of a hollow body organ thereto and rotate about a longitudinal axis of the member; and

a fastening device adapted to fasten the rotated portion of the interior wall.

64. The system of claim 63, wherein each member is adapted to rotate relative to one another about the longitudinal axis of each member.

65. The system of claim 63, wherein the at least two elongate members are adapted to counter-rotate relative to one another.

66. The system of claim 63, wherein each member defines a lumen through the length.

67. The system of claim 66, wherein each member defines an opening along the length which is in communication with the lumen.

68. The system of claim 63, wherein each member is adapted for insertion into the hollow body organ via an esophageal passageway.

69. The system of claim 63, wherein each member is adapted to adhere the portion of interior wall via a vacuum.

70. The system of claim 63, further comprising a vacuum pump in fluid communication with each member.

71. The system of claim 63, wherein each member is parallel with one another.

72. The system of claim 63, further comprising at least one attachment point positioned adjacent to the opening.

73. The system of claim 63, wherein the distal end of each member is offset relative to the longitudinal axis.

74. The system of claim 63, wherein the fastening device comprises a clip adapted to be inserted into the member.

75. The system of claim 63, wherein the fastening device comprises at least one staple.

76. The system of claim 63, further comprising a plurality of additional fastening devices, each of the fastening devices being configured to fasten the portion of interior while juxtaposed linearly to form a continuous fixation line.

77. The system of claim 63, further comprising a handle attachable to the proximal end of each member.

78. The system of claim 63, wherein each member is adapted to rotate the portions of interior wall such that an overlying region of tissue is created.

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摘要(译)

本文描述了各种肥胖症治疗工具和方法，以及其他胃相关疾病（例如GERD）的治疗。治疗包括减小胃袋的尺寸以限制热量摄入以及提供更早的饱腹感。这可以通过直接从胃内部在胃内形成较小的胃袋来完成。较小的袋可以通过使用单独的锚固装置，旋转探针或体积减小装置来制造。还可以进行幽门成形术以使幽门括约肌不能胜任。可以另外使用无创伤磁性吻合装置进行胃旁路手术，使得糖和脂肪在绕过胃时直接传递到肠。许多这些程序可以以各种组合进行。治疗可以通过阻止摄入高热量食物来产生强制性的行为改变

