FIG. 8
Published:

— with international search report (Art. 21(3))
System and Methods for Internalization of Components of an Adjustable Intragastric Balloon

CROSS-REFERENCE TO RELATED APPLICATIONS

[0001] This application claims priority to U.S. Provisional Application Nos. 61/729,654 filed November 26, 2012 and 61/770,364 filed February 28, 2013, the disclosures of which are incorporated herein by reference in their entireties.


TECHNICAL FIELD OF THE INVENTION

[0003] The present invention relates generally to systems and methods directed to adjustable intragastric balloons, and more particularly, to systems and methods for internalizing components of an adjustable intragastric balloon.

BACKGROUND OF THE INVENTION

[0004] Morbid obesity remains an ever-growing problem in the world. Varying forms of gastric bypass surgery have developed and have improved over the last few decades. Recently, laparoscopic gastric banding has emerged as a less invasive surgical option. However, bariatric surgery is fraught with morbidity of up to 20%, with a re-operation rate approaching 25% at 3-5 years post-op. Bariatric surgery
carries an operative mortality of 0.5%. Diet and pharmaceutical alternatives have not been very effective, with a high recidivism rate. Intragastric balloons in use can achieve weight loss and a drop in BMI. However, balloon deflation can result in unwarranted migration leading to obstruction.

[U0005] U.S. Patent Nos. 8,430,894 and 8,430,895 describe balloon systems having a balloon coupled to a gastrointestinal anchor that is configurable to prevent passing of the anchor into a duodenum of a subject. These balloon systems also each describe a stretchable inflation tube that can stretch from a stomach to a mouth of the subject, about 2.5-10 times its length, to facilitate inflation or adjustment of volume of the balloon. The inflation tube is partly inside and partly outside of the balloon. U.S. Patent Application Publication No. 2006/0142731 and U.S. Patent No. 8,403,952 describe a floating anchor that can be used, for example, with a gastrointestinal balloon. However, further improvements can be made to limit side effects from implantable devices including, but not limited to, tissue trauma, difficulty finding the tube, and difficulty grasping the tube, some of which may result from components of the devices that are located on the outside of the gastrointestinal balloon.

[U0006] Intragastric balloons still need to be improved upon with respect to adaptation to the body, monitoring, adjustment, and removal.

**BRIEF SUMMARY OF THE INVENTION**

[U0007] Embodiments of the invention may provide an apparatus for use in a stomach of a subject. The apparatus comprises a balloon adapted for placement in the stomach and having an opening to an interior of the balloon, an internal tube in the interior of the balloon with a proximal end coupled to the opening of the balloon, and an inflation tube having a distal end in the interior of the balloon and a proximal end configured to receive a fluid on an exterior of the balloon, the inflation tube being configured to stretch from the stomach to a mouth of the subject to facilitate inflation of the balloon and to have a resting position that is substantially within the interior of the balloon, the internal tube being disposed around at least a portion of the inflation tube that is in the interior of the balloon.
[0008] Embodiments of the invention may also include a method of adjusting an inflation level of a balloon inserted into a stomach of a subject where the balloon is in fluid communication with an inflation tube. The method comprises stretching the inflation tube while keeping the balloon in the stomach, the inflation tube extending from within the balloon when stretching, transferring a fluid to or from the balloon via the inflation tube which may or may not require connection to an extension tube, to adjust the inflation level of the balloon, and returning the inflation tube to a resting position after adjusting the inflation level, a substantial portion of the inflation tube being contained within the balloon in the resting position.

[0009] Additional features, advantages, and embodiments of the invention are set forth or apparent from consideration of the following detailed description, drawings and claims. Moreover, it is to be understood that both the foregoing summary of the invention and the following detailed description are exemplary and intended to provide further explanation without limiting the scope of the invention as claimed.

**BRIEF DESCRIPTION OF DRAWINGS**

[0010] The accompanying drawings, which are included to provide a further understanding of the invention and are incorporated in and constitute a part of this specification, illustrate preferred embodiments of the invention and together with the detailed description serve to explain the principles of the invention. In the drawings:

[0011] FIG. 1A shows an intragastric balloon with an internal tube in an extended position and an inflation tube disposed substantially inside the balloon in accordance with an embodiment of the invention.

[0012] FIG. 1B shows the intragastric balloon of FIG. 1A with the internal tube in a shortened position and the inflation tube disposed with a substantial portion outside the balloon in accordance with an embodiment of the invention.

[0013] FIG. 2A shows an intragastric balloon with an internal tube in an extended position within a rigid support and a tether at a resting length, and an inflation tube
disposed substantially inside the balloon in accordance with an embodiment of the invention.

[0014] FIG. 2B shows the intragastric balloon of FIG. 2A with the internal tube in a shortened position within the rigid support and tether at a stretched length, and the inflation tube disposed with a substantial portion outside the balloon in accordance with an embodiment of the invention.

[0015] FIG. 3A shows an intragastric balloon with a telescoping internal tube in an extended position with the spring at the resting length and an inflation tube disposed substantially inside the balloon in accordance with an embodiment of the invention.

[0016] FIG. 3B shows the intragastric balloon of FIG. 3A with the telescoping internal tube in a shortened position, with the spring at the compressed length and the inflation tube disposed with a substantial portion outside the balloon in accordance with an embodiment of the invention.

[0017] FIG. 4A shows an intragastric balloon with a telescoping internal tube in an extended position within a rigid support and a tether at a resting length, and an inflation tube disposed substantially inside the balloon in accordance with an embodiment of the invention.

[0018] FIG. 4B shows the intragastric balloon of FIG. 4A with the telescoping internal tube in a shortened position within the rigid support and tether at a stretched length, and the inflation tube disposed with a substantial portion outside the balloon in accordance with an embodiment of the invention.

[0019] FIG. 5 shows an intragastric balloon with an internal tube and a recess to accommodate components of the intragastric balloon assembly in accordance with an embodiment of the invention.

[0020] FIG. 6 shows an intragastric balloon with an elongated internal tube and a recess to accommodate components of the intragastric balloon assembly in accordance with an embodiment of the invention.
FIG. 7 shows an intragastric balloon with a curved internal tube and a recess to accommodate components of the intragastric balloon assembly in accordance with an embodiment of the invention.

FIG. 8 shows a close-up view of a portion of an intragastric balloon with a recess in accordance with an embodiment of the invention.

FIG. 9 shows an intragastric balloon with a support and internal curved tube in accordance with an embodiment of the invention.

FIG. 10A shows an intragastric balloon and endoscope assembly before expansion of the balloon and with a containing member having perforations in accordance with an embodiment of the invention.

FIG. 10B shows the intragastric balloon and endoscope assembly of FIG. 10A during expansion of the balloon and shows widening of the perforations from a partially expanded state of the balloon in accordance with an embodiment of the invention.

FIG. 10C shows the intragastric balloon and endoscope assembly of FIG. 10A from a side view during expansion of the balloon and shows expansion of the balloon through the perforations in accordance with an embodiment of the invention.

FIG. 10D shows the intragastric balloon and endoscope assembly of FIG. 10A during further expansion of the balloon after tearing of the perforations in accordance with an embodiment of the invention.

FIG. 11A shows an intragastric balloon and endoscope assembly with a containing member in a state for insertion of the assembly in a patient's body in accordance with an embodiment of the invention.

FIG. 11B shows the intragastric balloon and endoscope assembly of FIG. 11A during removal of the containing member in accordance with an embodiment of the invention.
FIG. 11C shows the intragastric balloon and endoscope assembly of FIG. 11A after the containing member has been substantially removed from the balloon in accordance with an embodiment of the invention.

DETAILED DESCRIPTION

An adjustable balloon may be a fluid filled balloon that has a stretchable inflation tube connected to the balloon. The inflation tube exits the balloon within a surrounding tube and ends in a valve. During an insertion procedure, a pusher and a firm guidewire within the pusher may connect to the balloon and an external catheter in one straight continuum. Alternatively, the balloon may be carried by the endoscope into the stomach using a sleeve connected to the endoscope, and without need of a guidewire. The inflation tube may stretch about 6 to 10 times its length, and may allow the valve to be pulled out of the stomach while the balloon remains in the stomach. This can allow adjustment of the volume of the balloon. Applicant’s other patent applications referenced and incorporated above discuss some features of a similar balloon.

An adjustable balloon system may have a stretchable inflation tube that is partly within and partly outside of the balloon. An external tube can produce challenges in several ways, including but not limited to causing tissue trauma, difficulty in finding the tube, and difficulty in grasping the tube.

Embodiments described herein provide systems and methods to internalize the external components into the balloon. In an embodiment, an internal rigid catheter may house a stretchable inflation tube that can be bonded in its most distal 1 cm to an internal surface of the rigid catheter, thereby allowing a remainder of the length of the stretchable inflation tube to slide freely within a lumen of the rigid catheter and thereby attaining its maximum stretchability outside of the balloon. For example, a 10 cm long inflation tube bonded in this way in its most distal 1 cm has a remaining 9 cm of length that is unbound and may stretch to lengths of 60-90 cm. The rigid catheter inside the balloon may be one stagnant length and may only hold a set length of stretchable inflation tube. In some embodiments, the internal rigid catheter can be made to have changeable lengths such that at a resting state it is one length, but when
pulled by the bonded end of the inflation tube it can shorten partially and thereby send
more stretchable inflation tube to the exterior of the balloon. This in turn can shorten
the necessary length of the external inflation tube.

[0034] In an embodiment, the internal rigid catheter may have have segments of
accordion-like walls that may contract when pulled to shorten the length of the rigid
catheter. The thickness of the accordion walls can be varied so that they can have
more or less propensity to return to a baseline longer length. Additionally, a metal
spring can be provided (internal to or immediately surrounding the catheter) that
promotes lengthening of the rigid catheter. Alternatively, a distal end of the variable
sized catheter can be bound to a distal end of a surrounding fixed length catheter by a
stretchable tether or spring that promotes lengthening of the catheter to its longer
length. Alternatively, the distal end of the variable sized catheter can be bound to a
distal balloon wall by a stretchable tether or spring that promotes lengthening of the
catheter to its longer length.

[0035] In an embodiment, the variable sized catheter can be formed as an antenna-
type tube, similar to an old-fashioned television antenna, that houses progressively
smaller diameter tubes so that the variable sized catheter can shorten or lengthen as
desired. The inflation tube can sit inside the smallest diameter segment. The smallest
diameter segment would therefore be large enough to house the inflation tube and
may have a clearance, for example, at least 1 mm of space, between the inflation tube
and catheter wall. Additionally, a metal spring can be provided (internal to or
immediately surrounding the catheter) to promote lengthening of the variable sized
catheter. The variable sized catheter can also be can be bound to the distal end of a
surrounding fixed length catheter by a stretchable tether or spring that promotes
lengthening of the collapsible catheter to its longer length. Alternatively, the variable
sized catheter can be bound to the distal balloon wall by a stretchable tether or spring
that promotes lengthening of the catheter to its longer length.

[0036] In some embodiments, systems and methods are provided for internalizing
within the balloon an external catheter, valve and cap. The result may provide a
smooth balloon without substantial external protrusions. This result can be
accomplished according to various embodiments discussed herein. In an embodiment,
the area of the balloon from which exits the catheter will be shaped to form a
concavity or recess, that is deep enough to hold the valve and cap. For example, the concavity may be approximately 1-5 cm deep. The concavity may be created by differing the density of a material such as silicone or adding other materials to the wall of the balloon in the area of the concavity. Alternatively, the balloon can be lengthened such that the internal catheter can be longer and house more inflation tube so that less protrudes outside of the balloon. The longer balloon can be spherical, pear shaped, cylindrical, or have other shapes that fit in the stomach. The inflation tube may stretch about 6-10 times its length. Alternatively, the inflation tube may be constructed of different materials, such as a different silicone material, and may be able to stretch 10-20 times its length. Alternatively, the internal catheter that houses the inflation tube, and may be a solid straight silicone catheter in an embodiment, can be modified to house a substantially frictionless internal canal that houses the inflation tube. For example, the substantially frictionless internal canal may be lined with Teflon ® or other similar materials, or may be filled with liquid silicone oil or other friction reducing material. Reducing the friction in these or other ways would allow the catheter to have a curved shape, such as a shape in a sinusoidal pattern, effectively lengthening the catheter and enabling more inflation tube to be inside the balloon. The catheter may be one piece that starts internally and is glued to the top of the balloon after which it exits the balloon. Alternatively, the catheter may be completely internal catheter that ends in a receptacle at its glue point with the balloon. The receptacle may be big enough to house all or substantially all of the valve and cap. Any combination of the above can accomplish the internalization of the external catheter, valve and cap. Additionally, a supporting frame that can also act as an anchor may also be provided in combination with any of the above features.

[0037] Figs. 1A and IB show an intragastric balloon assembly 100 according to an embodiment of the invention. The assembly 100 generally includes balloon 102 with a substantial internally disposed inflation assembly that can maintain the balloon 102 profile when the inflation assembly is dormant (Fig. 1A), but can be adapted for operational use outside the balloon 102 profile, as shown in Fig. IB. This can be accomplished through the use of an internal tube 114 alone or in combination with a recess 140. The assembly 100 includes a balloon 102 adapted for placement in the stomach of a subject. The balloon 102 has an opening 115 to an interior 104 of the balloon 102. An inflation tube 108 has a distal end 118 and a proximal end 119. The
distal end 118 is positioned in the interior 104 of the balloon 102. The proximal end 119 is adapted to receive a fluid and may be positionable on an exterior 106 of the balloon 102. An internal tube 114 in the interior 104 has a proximal end 121 coupled to the opening 115 of the balloon 102. The inflation tube 108 is contained within a lumen 122 defined by an inner wall 116 of the internal tube 114. A distal end 120 of the internal tube 114 is coupled to the distal end 118 of the inflation tube 108.

The inflation tube 108 can stretch from the stomach to a mouth of the subject to facilitate inflation of the balloon 102. However, in a resting position as shown in Fig. 1A, the inflation tube 108 is substantially within the balloon 102. It can be in the interior 104 of the balloon 102, substantially within the interior 104 of the balloon, or within the general profile of the balloon, which is shown in Fig. 1A by a representative broken line 103. In some embodiments, the recess 140 may have a depth ranging from 10 to 50 mm. The resting position corresponds to a position before the inflation tube 108 is stretched away from the balloon or after the inflation tube 108 has been released from the stretched position and returned to the resting position. According to this internalization of components in the resting position, it may be possible to limit or avoid trauma to tissue of the patient that may otherwise be caused by portions of an intragastric balloon assembly, such as a inflation tube, cap and valve, or other catheter, that protrude from the balloon or profile of the balloon. The returning to the resting position after being stretched may be facilitated by the shrinking or un-stretching of the inflation tube 108, or by the elongation of the internal tube 114 which pulls the distal end 118 of the inflation tube 108, or by a combination of both mechanisms. The internal tube 114 is positioned around at least a portion of the inflation tube 108 that is in the interior 104 of the balloon 102. In some embodiments, a distance x separates the inflation tube 108 from the interior wall 166 of the internal tube 114 to facilitate the stretching and/or sliding of the inflation tube 108 relative to the interior wall 166.

As shown in Figs. 1A and 1B, the internal tube 114 may have a variable length including a first length L1 that accommodates a substantial portion of the inflation tube 108 when the inflation tube 108 is in the resting position (see Fig. 1A), and a second length L2, which is shorter than the first length L1, when the inflation tube 108
is stretched outwardly from the balloon 102 (see Fig. IB). The inflation tube 108 has an un-stretched length \( L_3 \) in the resting position and a stretched length \( L_4 \) in the stretched position.

[0040] The wall of the internal tube 114 in Fig. 1 has a plurality of accordion structures 124 to facilitate the variable length of the internal tube 114. In the example shown in Figs. 1A and IB, each of the two accordion structures 124 includes three peaks 126 and two valleys 128 between the peaks 126. However, the internal tube 114 is not limited to this configuration and may have one or more accordion structures 124 that each has at least one, but possibly more, peak 126 and/or valley 128. Multiple accordion structures 124 may be separated by flat portions 125 of the internal tube 114. As shown in Fig. IB, the accordion structures 124 contract or collapse to allow shortening of the internal tube 114 away from a distal end 138 of the balloon when the inflation tube 108 is stretch outwardly from the balloon 102. Further, upon release of the stretched inflation tube 108 shown in Fig. IB, the inflation tube 108 and internal tube 114 may return to the resting position show in Fig. 1A. By returning to this resting position, components of the balloon assembly 100 may be contained within or substantially within the balloon or the balloon profile. Thus, trauma to tissues of the patient may be limited or prevented that may have otherwise been caused from protruding portions or an irregular profile of an intragastric balloon according to other constructions.

[0041] The opening 115 of the balloon 102 may be formed as a cylindrical structure that is recessed with respect to the walls of the balloon 102 on either side of the opening 115. In addition, the opening 115 may be sized to accommodate one or more structures, such as a cap and valve 110, on the proximal end 119 of the inflation tube 108.

[0042] Figs. 2A and 2B show an intragastric balloon assembly 100 according to an embodiment of the invention that includes a rigid support structure 132. The rigid support structure 132 remains fixed as the internal tube 114 varies in length. A distal end 134 of the rigid support structure 132 may be coupled to the distal end 120 of the internal tube 114 via one or more stretchable tethers or springs 136. The tether or spring 136 biases the internal tube 114 to its lengthened position so that the inflation tube 108 and any other external structures will be pulled in or toward the interior 104
of the balloon 102. In some embodiments, a tether or spring may be coupled to the
distal wall 138 of the balloon 102, rather than the rigid support structure 132.

[0043] The rigid support structure 132 may define an opening 135 at its distal end 134
allow fluid to flow to and from the inflation tube 108 to the remainder of the balloon
102. The rigid support structure 132 may be substantially coaxial with the internal
tube 114.

[0044] Figs. 3A and 3B show an intragastric balloon assembly 200 according to
another embodiment of the invention. In this embodiment, structures similar to the
preceeding embodiments are similarly numbered and may not be discussed in full
detail below. The internal tube 214 of Figs. 3A and 3B is a telescoping tube having
more than one telescoping segments 224 slideably engaged with each other. Fig. 3A
shows the internal tube 214 in an extended position, whereas Fig. 3B shows the
internal tube 214 in a contracted position.

[0045] The corresponding diameters D1, D2, D3 of the various telescoping segments
224 allow for one segment to slide within or over another. The diameter D1 of the
smallest segment 224 must be sufficiently wide to accommodate the inflation tube
208. For example, there may be a space x' between the inflation tube 208 and the
internal wall 216 of the telescoping segment 224 with diameter D1. The space x' may
facilitate stretching and movement and/or sliding of the inflation tube 208 relative to
the internal wall 216 of telescoping segment 224.

[0046] Figs. 4A and 4B show an embodiment of the intragastric balloon assembly
200 that includes a rigid support structure 232 with one or more stretchable tethers or
springs 236. Although only three telescoping segments 224 are shown in Figs. 3A,
3B, 4A, and 4B, more or fewer may be used.

[0047] Figs. 5-9 show additional embodiments of an intragastric balloon assembly
300 according to the invention. In Figs. 5-9, a proximal end 319 of the inflation tube
308 includes one or more structures, including a valve 310 and a removable cap 312.
The valve 310 can be selectively opened to perform an adjustment of the inflation
level of the balloon 302, and closed when the adjustment is completed. The cap 310
can cover the valve 310 to seal it or to keep foreign particles or bacteria away from
the valve.
To minimize the extent to which these external structures project outward from the balloon 302, a recess 340 is formed on the outer surface of the balloon 302. The recess 340 may have a depth d, as shown in Fig. 5, that is sufficient to house substantially all or some portion of the external structures, including valve 310 and cap 312, of the inflation tube 308 when the inflation tube 308 is in a resting (or non-stretched) position. In other words, a depth of the recess 340 may be greater than or equal to a height of the inflation tube 308 that is within the recess 340 in the resting position.

By allowing more of the inflation tube 308 and other related elements to be stored within the balloon 302 or recess 340, a smoother overall appearance of the balloon may be achieved by reducing or eliminating extension of those elements past a profile of the balloon, which is shown in Fig. 5 by a representative broken line 303. However, it may not be necessary for the inflation tube 308 and related elements to be completely within the balloon 302 or recess 340 in some embodiments. For example, a portion of the inflation tube 308, valve 310, or cap 312 may extend beyond the profile of the balloon.

Figs. 5 and 6 shows internal tubes 314 of different lengths. By providing an internal tube 314 of a certain length, a certain amount of inflation tube 308 may be stored within the internal tube 314. Thus, in some embodiments the internal tube 314 may be longer than other embodiments (e.g., compare Fig. 6 to Fig. 5) so that a longer inflation tube 308 may be used. A longer inflation tube 308 may also allow the inflation tube 308 to be stretch farther from the balloon 302.

Fig. 7 shows another embodiment where the internal tube 314 has a curved shape. The internal tube 314 includes a number of curved portions 348, corresponding to approximately two periods of a sinusoidal wave. Thus, curves 348 can accommodate an inflation tube 308 having a long effective length in a shorter balloon 302. The shape and curvature of the internal tube 314 can vary. For example, the curvature might be adjusted according to the friction between the internal tube 314 and the inflation tube 308 so that the inflation tube 308 can easily slide past the interior wall 316 of the internal tube 308 when being stretched or returning to the resting position.
To facilitate sliding of the inflation tube 308 within the internal tube 314, the interior wall 316 of the internal tube 314 may be formed of or coated with a low-friction or frictionless material or lubricant, or the internal tube 314 can be filled with a substance to reduce friction.

The inflation tube 308 of Figs. 5-7 also includes a curved end portion 309 which can help prevent fluid flow from being blocked during inflation or deflation of the balloon. For example, it is possible that during deflation of some intragastric balloons an internal structure or tissue of the patient may be suctioned against the opening on the distal end of the inflation tube, which could block further deflation. The curved end portion 309 may prevent such a blockage. The curved end portion 309 may also be provided as an atraumatic end on the distal end 318 of the inflation tube 308. Thus, the curved end portion 309 may minimize the potential of the relatively sharp edges of the opening on the distal end 318 of the inflation tube 308 from contacting and possibly damaging an internal structure such as the balloon or the tissue of the patient.

In Fig. 8, a receptacle 315 is provided within the recess 340. The receptacle 315 is sized to accommodate the external structures, such as the cap 312 and valve 310. A filler substance 350 may be used to fill a space 352 between the receptacle 315 and the walls of the recess 340. Such a filler substance 350 can be used to bond the receptacle 315 to the balloon 302 and to keep external matter or bacteria from becoming trapped in the space 352. In an embodiment, the filler substance 350 may be some type of glue or epoxy. In addition, the receptacle 315 may serve to form or hold the shape of the recess 340.

In some embodiments, walls of the balloon 302 surrounding the opening or receptacle 315 may involute and be attached or glued to the receptacle 315 and/or internal tube 314, as shown in Fig. 8, for example. The receptacle 315 may either be formed from the proximal end of the internal tube 314 or be a structure that is attached to the proximal end of the internal tube 314 via glue, adhesive, or some structure attachment mechanism. The receptacle 315 that is outside of the balloon 302, though it may still be within the balloon profile shown in Fig. 8 by a representative broken line 303, may be made formed as a soft flexible catheter. However, in an embodiment, the receptacle 315 nonetheless has sufficient firmness to
help maintain the shape of the recess 340, even as the volume of the balloon is adjusted or the shape of the balloon changes. Furthermore, the material and stiffness of the receptacle 315 can prevent the components disposed within the receptacle 315 from making harmful or injurious contact with internal tissues of the patient. Additionally, the receptacle 315 is itself of suitable construction to not injure the tissue of the patient while maintaining the shape of the balloon under normal operating conditions and pressures. The receptacle 315 may therefore provide protection for the inflation tube 308, valve 310, and cap 312 that are disposed within it, while also not being injurious to internal organs or tissue of a patient.

Fig. 9 shows an embodiment with a rigid support structure 332. The internal tube 314 in Fig. 9 has a curved shape with the curved sections 348 being attached to the rigid support structure 332 at anchor points 349. The internal tube 314 of this embodiment may have a frictionless or low-friction internal canal 344. For example, the internal canal 344 may have a lining 346 to facilitate movement of the inflation tube 308 relative to the internal tube 314.

In the above embodiments, the rigid support structure 132, 232, 332 may be formed, for example, as a tubular or cylindrical structure, or may be formed as one or more beams in a support frame.

Embodiments of the invention may provide a system and method for internalizing external components of an intragastric balloon within the balloon. However, some embodiments provide for a portion of these external components to remain on an outside of the balloon. In some embodiments, an internal tube houses a stretchable inflation tube. The stretchable tube may be bonded at its distal end to an inner surface of the internal tube, thereby allowing the remainder of the inflation tube to slide freely within the lumen of the internal tube and thereby attain its maximum stretched length outside of the balloon. For example, the stretchable inflation tube may be bonded in its most distal 1 cm to the internal surface of the rigid catheter. In such an example, a 10 cm long inflation tube can stretch the 9 cm that are unbound to lengths of 60-90 cm.

The internal tube may be a fixed length and therefore hold only a set length of the stretchable inflation tube. However, in embodiments of the invention, the internal
tube can be made to have a changeable length such that at a resting state it is one length, but when pulled it can shorten. In a shortened state of the internal tube, the stretchable inflation tube may then extend farther from the balloon. In addition, the length of the inflation tube can be decreased due to the increase distance provided by the shortened internal tube.

In some embodiments, for example, when the inflation tube is pulled or stretched away from the balloon to facilitate inflation or adjustment of the volume of the balloon, the internal tube is shortened or compressed as it is pulled by the distal end of the inflation tube. When the inflation tube is released from its pulled or stretched position, the internal tube will return from the shortened or compressed state to a resting or un-compressed state. In returning to the resting or un-compressed state, the internal tube pulls the distal end of the inflation tube to assist in returning the inflation tube its resting state such that the inflation tube is in the interior of the balloon, substantially within the interior of the balloon, or within the general profile of the balloon.

According to some embodiments, the internal tube may have segments of accordion walls that contract when pulled to shorten the length of the internal tube. The structure of the accordion walls can be varied such that they will have more or less propensity to return to the baseline longer length. For example, a thickness of the accordion walls can be varied. In addition, or alternatively, a spring can be provided that promotes lengthening of the internal tube. Such a spring can be made of metal and can be disposed an internal or an external side of the internal tube.

In some embodiments, the distal end of the variable sized internal tube can be coupled to a distal end of a surrounding structure, such as a rigid support structure. These may be coupled by, for example, a stretchable tether or spring that promotes lengthening of the internal tube to its longer length. Alternatively, the internal tube can be coupled to a distal balloon wall by a stretchable tether or spring that promotes lengthening of the internal tube to its longer length.

According to some embodiments, the internal tube can be formed of telescoping segments (e.g., an old-fashioned television antenna-like structure) that houses progressively smaller diameter tubes that can shorten or lengthen as desired.
similar to the accordion structures of an above-discussed embodiment. The inflation tube may fit within the smallest diameter segment of the telescoping internal tube. The smallest segment can therefore have a diameter large enough to accommodate the inflation tube. In some embodiments, a space between the inflation tube and the internal tube is provided. For example, the space may be at least 1 mm.

[0064] In some embodiments, any external catheter, including the inflation tube, as well as the valve and cap on the inflation tube may be internalized within the balloon or within an outer shape of the balloon. As a result, a smooth balloon without any external protrusions may be provided. This can be accomplished in several ways. In some embodiments, the area of the balloon from which the catheter or external components exit will be shaped to form a recess or concavity. The recess may be deep enough to hold the external portion of the catheter, including the valve and cap. For example, the recess may be approximately 1-5 cm deep. The recess may be created by differing the density of the balloon wall or silicone, or by adding other materials to the wall of the balloon in the area of the recess. For example, the balloon may comprise a silicone wall having varying thickness, or multiple silicone materials of different stiffnesses. Additionally, materials, such as internal layers of materials, may be added to the balloon. Such layers may be formed in mesh or solid structures. In an embodiment, multiple layers of silicon may be used with intervening mesh layers. Other materials may also be used, including, for example, polypropylene, polyethylene, nylon, and latex.

[0065] In some embodiments, the valve may comprise a silicone housing with a luer lock neck that can be closed with the cap. The housing may have a funnel or tapered shape. The funnel may contain a one-way or two-way valve that is made from hard material such as polypropylene, or it may contain a valve made of a soft material such as silicone. The funnel could then be closed with a soft material such as silicone in the shape of a thin rubber sheath that rolls on and off. The thin rubber sheath may have a cylindrical shape with a closed first end and an open second end. The soft material version would be utilized to prevent tissue damage.

[0066] In some embodiments, the balloon can be elongated such that an internal tube or catheter can be longer and house more of the inflation tube, leaving less to protrude
outside of the balloon. According to various embodiments, the balloon can be spherical, pear-shaped, cylindrical, or other shapes that fit in the stomach.

[0067] In some embodiments, the inflation tube may stretch 6-10 times its length. In other embodiments, the inflation tube may stretch 10-20 times its length. The inflation tube may be made from different silicone material having different stretchability or elasticity.

[0068] In some embodiments, the internal tube or catheter may be formed as a solid, straight catheter and may be made from, for example, silicone. In some embodiments, the internal tube or catheter that houses the inflation tube can be modified to house a frictionless internal canal that houses the inflation tube. For example, the internal tube can be lined with TEFLON ® or other similar materials, or filled with liquid silicone oil or other friction reducing material. The reduced friction can help enable internal tubes of different shapes and paths according to some embodiments. For example, the internal tube may curve in a sinusoidal pattern. In this way, the effective length of the internal tube is increased over a limited straight-line distance. Therefore, the internal tube may be allowed to contain more of the inflation tube inside the balloon.

[0069] The internal tube may be provided as one piece that starts internally and is glued to the top of the balloon, after which it exits the balloon. According to some embodiments, the internal tube can be changed to a completely internal tube or catheter that ends in a receptacle at its glue point with the balloon. The receptacle is big enough to house the valve and cap of the inflation tube.

[0070] Any combination of all of the above can accomplish the internalization of the external catheter, valve and cap, with a supporting frame that can also act as an anchor.

[0071] Some embodiments of the present invention relate to the elimination of the guide wire and pusher used during the insertion process of an intragastric balloon. For example, the insertion procedure may include the balloon being attached to an endoscope and pulled into the stomach during a routine endoscopy procedure. The balloon may then be released from the endoscope and inflated. This process can be accomplished in several ways.
In some embodiments, the balloon may be folded in a way to allow it to be wrapped around the endoscope. Additionally, a cylindrical containing member or wrapping piece may be attached to the distal end of the endoscope. Such a wrapping piece can be made to fit snugly on one or all endoscope sizes, and/or may be adjustable to different size endoscopes. The wrapping piece may be made of a pliable material including, for example, silicone, rubber, polymer or other pliable material, or of a non-pliable material such as a fabric that can hold the balloon. The wrapping material may be able to securely hold, for example, about 2-15 cm of the balloon length during the insertion of the endoscope into the stomach.

Once in the stomach, the balloon is separated from the endoscope. This can be accomplished in several ways. In some embodiments, the wrapping piece can have perforations on its surface such that inflation of the balloon will expand the wrapping piece and the perforations will open and release the balloon. Alternatively, the inflation of the balloon may automatically push the balloon out of the wrapping piece's grasp. Alternatively, the proximal edge of the wrapping piece may be connected to a string that enters a biopsy channel at the distal end of the endoscope. The string, which may exit the top of the biopsy channel, may be pulled to pull the proximal edge of the wrapping piece distally and thereby release the balloon.

Some embodiments of the invention relate to a method of adjusting an inflation level of a balloon inserted into a stomach of a subject where the balloon is in fluid communication with an inflation tube. The method includes stretching the inflation tube while keeping the balloon in the stomach. The inflation tube will extend from within the balloon when stretching. In its extended position, a fluid is transferred to or from the balloon via the inflation tube to adjust the inflation level of the balloon. Subsequently, the inflation tube is returned to a resting position after adjusting the inflation level. In this resting position, a substantial portion of the inflation tube is contained within the balloon or within a recess on the outer surface of the balloon. Thus, the balloon has reduced protrusions on the exterior.

Although the foregoing description is directed to the preferred embodiments of the invention, it is noted that other variations and modifications will be apparent to those skilled in the art, and may be made without departing from the spirit or scope of the invention. Moreover, features described in connection with one embodiment of
the invention may be used in conjunction with other embodiments, even if not explicitly stated above.

[0076] In some embodiments, the internal tube may be made of a shape-memory material to facilitate a return of the internal tube to the resting position.

[0077] Figs. 10A-10D and 11A-C show an intragastric balloon and endoscope assembly 400 according to additional embodiments of the invention.

[0078] An adjustable balloon may be a fluid filled balloon that has a stretchable inflation tube connected to the balloon. The inflation tube exits the balloon within a surrounding tube and ends in a valve. During an insertion procedure, a pusher and a firm guidewire within the pusher may connect to the balloon and an external catheter in one straight continuum. The inflation tube may stretch about 6 to 10 times its length, and may allow the valve to be pulled out of the stomach while the balloon remains in the stomach. This can allow adjustment of the volume of the balloon. Applicant’s other patent applications referenced and incorporated above discuss some features of a similar balloon.

[0079] In some embodiments, systems and methods are provided to eliminate the guidewire and pusher sometimes used during the insertion process. In an embodiment, the balloon may be attached to an endoscope and pulled into the stomach during a routine endoscopy procedure. The balloon may then be released from the endoscope and inflated. This release and inflation can be accomplished in several ways. For example, in an embodiment, the balloon is folded in a way to allow it to be wrapped around the endoscope. In addition, a cylindrical wrapping piece may be provided and attached to a distal end of the endoscope. The wrapping piece may fit snugly on different sized endoscopes and/or may be adjustable to different size endoscopes. The wrapping piece can be made of a pliable material such as silicone or rubber, or a non-pliable material such as a fabric. The wrapping piece may be configured to hold the balloon (for example, about 2-15 cm of the length of the balloon) securely during the insertion of the endoscope into the stomach. Once in the stomach, the balloon may be separated from the endoscope. In an embodiment, the wrapping piece can have perforations on its surface such that inflation of the balloon will expand the wrapping piece and the perforations will open and release the balloon. Alternatively, the
inflation of the balloon will automatically push the balloon out of the grasp of the wrapping piece. Alternatively, the proximal edge of the wrapping piece may be connected to a string that enters a biopsy channel at the distal end of the endoscope. The string exits the top of the biopsy channel, and, when pulled, can pull the proximal edge of the wrapping piece distally to release the balloon.

[0080] In Fig. 10A, the balloon 402 is attached to or wrapped around the distal end 458 of an endoscope 456 and covered or held in place by a wrapping 460. The wrapping 460 may include perforations 462. As shown in Figs. 10B and IOC, the perforations can expand and/or allow the wrapping 460 to tear when the balloon 402 is inflated. The perforations may be placed toward a proximal edge 464 of the wrapping 460 to control the tear and to release the balloon on the proximal side, as shown in Fig. 10D.

[0081] In Fig. 11A, the wrapping 460 is coupled to a string 466. The wrapping 460 is removed by the string 466 being pulled from the biopsy channel 468 to remove the wrapping 460 by pulling the proximal edge 464 of the wrapping 460, as shown in Figs. 11B and 11C.
CLAIMS:

1. An apparatus for use in a stomach of a subject, the apparatus comprising:
   a balloon adapted for placement in the stomach and having an opening to an interior of the balloon;
   an internal tube in the interior of the balloon with a proximal end coupled to the opening of the balloon; and
   an inflation tube having a distal end in the interior of the balloon and a proximal end configured to receive a fluid on an exterior of the balloon, the inflation tube being configured to stretch from the stomach to a mouth of the subject to facilitate inflation of the balloon and to have a resting position that is substantially within the interior of the balloon, the internal tube being disposed around at least a portion of the inflation tube that is in the interior of the balloon.

2. The apparatus according to claim 1, wherein a distal end of the internal tube is coupled to the distal end of the inflation tube.

3. The apparatus according to claim 2, wherein the internal tube has a variable length including a first length that accommodates a substantial portion of the inflation tube when the inflation tube is in the resting position, and a second length, which is shorter than the first length, when the inflation tube is stretched outwardly from the balloon.

4. The apparatus according to claim 3, wherein a wall of the internal tube has one or more accordion structures to facilitate the variable length of the internal tube.

5. The apparatus according to claim 3, further comprising:
a spring coupled to the internal tube, the spring biasing the internal tube to the first position.

6. The apparatus according to claim 3, wherein the internal tube comprises a plurality of telescoping segments slideably engaged with each other.

7. The apparatus according to claim 3, further comprising:

   a rigid support in the interior of the balloon disposed around, and extending coaxially with, the internal tube.

8. The apparatus according to claim 7, wherein the distal end of the internal tube is coupled to a distal end of the rigid support via a stretchable tether such that the stretchable tether is stretched when the internal tube is in the second position thereby biasing the internal tube toward the first position.

9. The apparatus according to claim 1, wherein the inflation tube comprises:

   a valve on the proximal end of the inflation tube configured to selectively open and close the inflation tube, and

   a removable cap on the proximal end of the inflation tube covering the valve.

10. The apparatus according to claim 1, wherein the opening of the balloon is disposed within a recess formed in an exterior surface of the balloon.

11. The apparatus according to claim 10, wherein the proximal end of the inflation tube is disposed within the recess in the resting position of the inflation tube.
12. The apparatus according to claim 11, wherein a depth of the recess is greater than or equal to a height of the inflation tube that is within the recess in the resting position.

13. The apparatus according to claim 10, the apparatus further comprising:

   a receptacle disposed within the recess and configured to surround the proximal end of the inflation tube when in the resting position.

14. The apparatus according to claim 13, wherein a gap between a wall of the recess and an outer wall of the receptacle is filled with a filler substance.

15. The apparatus according to claim 1, wherein the internal tube has a curved shape to increase a length of the inflation tube contained within the internal tube.

16. The apparatus according to claim 1, wherein a lumen wall of the internal tube is configured to allow sliding of the inflation tube relative to the lumen wall.

17. A method of adjusting an inflation level of a balloon inserted into a stomach of a subject where the balloon is in fluid communication with an inflation tube, the method comprising:

   stretching the inflation tube while keeping the balloon in the stomach, the inflation tube extending from within the balloon when stretching;

   transferring a fluid to or from the balloon via the inflation tube which may or may not require connection to an extension tube, to adjust the inflation level of the balloon; and
returning the inflation tube to a resting position after adjusting the inflation level, a substantial portion of the inflation tube being contained within the balloon in the resting position.

18. The method according to claim 17, further comprising:

positioning a proximal end of the inflation tube within a recess formed on an exterior of the balloon when returning the inflation tube to the resting position.
INTERNATIONAL SEARCH REPORT

Box No. II Observations where certain claims were found unsearchable (Continuation of item 2 of first sheet)

This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. ☒ Claims Nos.: 17-18
   because they relate to subject matter not required to be searched by this Authority, namely:
   Claims 17-18 pertain to methods for treatment of the human and thus relate to a subject-matter which this International Searching Authority is not required, under Article 17(2)(a)(i) of the PCT and Rule 39.1(iv) of the Regulations under the PCT, to search.

2. ☐ Claims Nos.:
   because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:

3. ☐ Claims Nos.:
   because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).

Box No. III Observations where unity of invention is lacking (Continuation of item 3 of first sheet)

This International Searching Authority found multiple inventions in this international application, as follows:

1. ☒ As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.

2. ☐ As all searchable claims could be searched without effort justifying an additional fees, this Authority did not invite payment of any additional fees.

3. ☒ As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.: 

4. ☐ No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos. :

Remark on Protest

☐ The additional search fees were accompanied by the applicant's protest and, where applicable, the payment of a protest fee.

☒ The additional search fees were accompanied by the applicant's protest but the applicable protest fee was not paid within the time limit specified in the invitation.

☒ No protest accompanied the payment of additional search fees.
A. CLASSIFICATION OF SUBJECT MATTER

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

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

- A61M 29/02; A61M 29/00; A61B 19/00; A61F 2/04; A61M 25/10

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

- Korean utility models and applications for utility models
- Japanese utility models and applications for utility models

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)
eKOMPASS (KIPO internal) & Keywords: stomach, balloon, internal tube, inflation tube, stretch

C. DOCUMENTS CONSIDERED TO BE RELEVANT

<table>
<thead>
<tr>
<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>X</td>
<td>US 2011-0218563  Al (BROOKS, J. S. et al.) 8 Sept ember 2011</td>
<td>1-2, 9, 15-16</td>
</tr>
<tr>
<td></td>
<td>See abst ract; paragraphs [0198H0199], [0202] - [0203]; and figures 13-16.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>See abst ract; paragraphs [0025]; [0028]; and figure 1.</td>
<td></td>
</tr>
<tr>
<td>A</td>
<td>US 2011-0092998  Al (HIRSZOWICZ, E. et al.) 21 April 2011</td>
<td>1-16</td>
</tr>
<tr>
<td></td>
<td>See abst ract; paragraphs [0194H0196], [0199], [0212], [0215H0218];</td>
<td></td>
</tr>
<tr>
<td></td>
<td>and figures 5, 8-10.</td>
<td></td>
</tr>
<tr>
<td>A</td>
<td>US 5259399    A (BROWN, A.) 9 November 1993</td>
<td>1-16</td>
</tr>
<tr>
<td></td>
<td>See abst ract; column 5, lines 9-54; column 6, lines 1-25;</td>
<td></td>
</tr>
<tr>
<td></td>
<td>column 7, lines 31-42; and figures 1-5.</td>
<td></td>
</tr>
<tr>
<td>A</td>
<td>US 7056305    B2 (GARZA ALVAREZ, J. R.) 6 June 2006</td>
<td>1-16</td>
</tr>
<tr>
<td></td>
<td>See abst ract; column 6, line 19 - column 7, line 19; and figures 1-3, 13-14.</td>
<td></td>
</tr>
</tbody>
</table>

Further documents are listed in the continuation of Box C. See patent family annex.

| Special categories of cited documents: |
| "A" document defining the general state of the art which is not considered to be of particular relevance |
| "E" earlier application or patent but published on or after the international filing date |
| "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) |
| "O" document referring to an oral disclosure, use, exhibition or other means |
| "P" document published prior to the international filing date but later than the priority date claimed |

"X" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention

"Y" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone

"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art

"&" document member of the same patent family

Date of the actual completion of the international search:
07 March 2014 (07.03.2014)

Date of mailing of the international search report:
10 March 2014 (10.03.2014)

Name and mailing address of the ISA/KR
International Application Division
Korean Intellectual Property Office
189 Cheongna-ro, Seo-gu, Daejeon Metropolitan City, 302-701,
Republic of Korea
Facsimile No. +82-42-472-7140

Authorized officer
Han, Inho
Telephone No. +82-42-481-3362

Form PCT/ISA/210 (second sheet) (July 2009)
<table>
<thead>
<tr>
<th>Patent document cited in search report</th>
<th>Publication date</th>
<th>Patent family member(s)</th>
<th>Publication date</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>EP 2004269 A4</td>
<td>04/09/2013</td>
</tr>
<tr>
<td></td>
<td></td>
<td>EP 2356956 Al</td>
<td>17/08/2011</td>
</tr>
<tr>
<td></td>
<td></td>
<td>JP 2009-331122 A</td>
<td>03/09/2009</td>
</tr>
<tr>
<td></td>
<td></td>
<td>JP 2011-147800 A</td>
<td>04/08/2011</td>
</tr>
<tr>
<td></td>
<td></td>
<td>US 2013-274789 Al</td>
<td>17/10/2013</td>
</tr>
<tr>
<td></td>
<td></td>
<td>US 8430894 B2</td>
<td>30/04/2013</td>
</tr>
<tr>
<td></td>
<td></td>
<td>US 8430895 B2</td>
<td>30/04/2013</td>
</tr>
<tr>
<td></td>
<td></td>
<td>AU 2004-317849 B2</td>
<td>31/03/2011</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CA 2569043 Al</td>
<td>13/10/2005</td>
</tr>
<tr>
<td></td>
<td></td>
<td>EP 1732635 A2</td>
<td>20/12/2006</td>
</tr>
<tr>
<td></td>
<td></td>
<td>EP 1732635 A4</td>
<td>27/05/2009</td>
</tr>
<tr>
<td></td>
<td></td>
<td>EP 1732635 Bl</td>
<td>27/07/2011</td>
</tr>
<tr>
<td></td>
<td></td>
<td>HK 1101680 Al</td>
<td>18/11/2011</td>
</tr>
<tr>
<td></td>
<td></td>
<td>US 8236023 B2</td>
<td>07/08/2012</td>
</tr>
<tr>
<td></td>
<td></td>
<td>WO 2005-094257 A2</td>
<td>17/10/2005</td>
</tr>
<tr>
<td>US 5259399 A</td>
<td>09/11/1993</td>
<td>None</td>
<td></td>
</tr>
<tr>
<td>US 7056305 B2</td>
<td>06/06/2006</td>
<td>CA 2438409 Al</td>
<td>19/09/2002</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CA 2538127 Al</td>
<td>17/03/2005</td>
</tr>
<tr>
<td></td>
<td></td>
<td>EP 1366716 Al</td>
<td>03/12/2003</td>
</tr>
<tr>
<td></td>
<td></td>
<td>EP 1366716 Bl</td>
<td>27/04/2005</td>
</tr>
<tr>
<td></td>
<td></td>
<td>EP 1671590 Al</td>
<td>21/06/2006</td>
</tr>
<tr>
<td></td>
<td></td>
<td>US 2004-0059289 Al</td>
<td>25/03/2004</td>
</tr>
<tr>
<td></td>
<td></td>
<td>WO 02-071951 Al</td>
<td>19/09/2002</td>
</tr>
<tr>
<td></td>
<td></td>
<td>WO 2005-023118 Al</td>
<td>17/03/2005</td>
</tr>
</tbody>
</table>