Endoscopic Bariatric/Metabolic Surgery

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Abstract

Endoscopic approaches to the bariatric patient have increased significantly in the last two decades. Obese patients present a challenge to surgeons as they are at higher risk of having perioperative complications. Patients also are seeking less and less invasive ways to have procedures performed. The combination of these two demands has led to the development of new technologies in the bariatric arena. Now there are several new endoscopic approaches for primary weight loss. Endoscopic metabolic procedures are developing for the management of diabetes. This article reviews the latest in these technologies.

Keywords
► bariatric endoscopy
► balloon
► endoscopic sleeve
► endoscopic weight loss procedures

Obesity is a healthcare crisis that is impacting cost and patient care in the United States. The core components of any weight loss program are diet and exercise, but multiple studies have shown that this is not a durable approach for many patients, especially those in obesity Class 2 and higher. The most durable weight loss approaches described are surgical and include procedures that are restrictive, malabsorptive, or both. In between the two extremes, there is a gap in weight loss management. This is where medications and endoscopic approaches to obesity fit in. Medications for weight loss will not be discussed in this article.

Endoscopic approaches are attractive to the patient because they are less invasive and less anatomically altering than surgical procedures. These procedures are attractive to referring physicians as these are perceived as less invasive and therefore carry less risk. For the interventional endoscopist, these procedures are technically challenging but very rewarding when the patient has successful weight loss. Endoscopic procedures are perceived as better for poor surgical candidates who need staging procedure to tolerate surgery super obese or those who are at the borderline of becoming obese. Emerging treatments also may help those who are normal weight but require help with glycemic control.1

For the last several decades, many endoscopic procedures have been attempted. These procedures follow the same principles as surgical approaches in that they are restrictive or malabsorptive. In the restrictive category, options include intragastric space occupying devices and sutures/staples to limit gastric capacity. These procedures work by limiting the capacity of food, creating satiety by using stretch receptors at the antrum, or reducing gastric emptying. The malabsorptive procedures include implantable sleeves, ablative techniques, and endoscopic anastomoses created by magnets. Many of these procedures are considered metabolic procedures that may provide new ways to treat diabetes. This article will review the latest technologies in all of these arenas.

Primary Endoscopic Weight Loss Procedures

Space-Occupying Devices

One of the first endoscopic devices for weight loss was the Garren-Edwards Gastric Bubble, developed in the 1980. This was considered groundbreaking at the time until this had complications including deflations that required surgical interventions, gastric ulcers, and patient intolerance. This device was removed from the market due to these serious complications and ineffective weight loss. These devices were then absent from the US market for years while they gained momentum in Europe.2 Now there are currently three space filling devices that are US Food and Drug Administration (FDA) approved within the United States—Orbera, Reshape, and Obalon. The first two are fluid filled, while the last is air filled.

Fluid-Filled Balloons

Orbera

Orbera (Bioenterics Intragastric Balloon (BIB), Apollo Endosurgery Inc., Austin, TX) is the first of these devices to be FDA approved in August of 2015. The fluid-filled single balloon is
placed and removed endoscopically under sedation after 6 months of implantation. The device is introduced by mouth, placed within the stomach, and inflated with saline from 500 to 700 mL to a grapefruit size volume (~10 cm) under direct endoscopic visualization. The device is made of a silicone polymer and is resistant to acid. Contraindications include gastroparesis, hiatal hernia, large ulcers, or prior gastric surgery.1–6

The first large series of Orbera device placements in Spain reported an excess weight loss (EWL) of 33.9 ± 18.7%. The authors noted diabetes resolution or improvement in 86.9% and hypertension improvement or resolution in 93.7% of patients. Their complication rate was low at 2.8%, but included perforation and death.7 Datis et al demonstrated that long-term weight loss can be maintained out to 2 and 1/2 in 100 individuals who had the balloon for only 6 months. Many studies all showed successful weight loss that is maintained if the patient is compliant and behavior change is initiated in the early stages of treatment.7 One of the largest series to date of the Orbera balloon is the Brazilian consensus statement of 40,000 balloons, of which ~32,000 were the Orbera. The mean percentage of total weight loss (TWL) was 18.4 ± 2.9%. Patients lost a mean of 18.3 ± 4.4 kg. The failure rate (defined as percentage of TWL < 10%) was only 8.3 ± 6.7%. The statement provides a guide reflective of large clinical experience on how to perform the procedures, manage patient nausea symptoms, and manage ongoing care of patients. Note, the data in this study shows better weight loss than most studies with the balloon.8

**Reshape**
The Reshape Duo uses two balloons attached in tandem allowing for higher filling volumes from 750 to 900 mL of saline. It takes up more of the stomach volume compared with a single balloon, avoids over distention, and conforms to the natural curve of the stomach for patient comfort and tolerability. This balloon is implanted endoscopically and removed the same way 6 months later. The other advantage stated is that if one balloon deflates, the other will remain inflated, thereby reducing migration. The REDUCE trial concluded that 55% of patients (mean body mass index [BMI] 45.4 kg/m2) with the balloon lost an average of 25.1% excess body weight within 6 months compared with only 6% with diet and exercise alone. Adverse events reported were balloon deflation in 6% of patients with no migrations, and gastric ulcers in 10% of patients.9 Other series with this balloon are limited.

Currently, there are several fluid-filled balloons in development and trials.

**Spatz Balloon**
The Spatz Adjustable Balloon System (Spatz Medical, NY) has an attached anchor and a valve to adjust the volume. This allows the physician to manage early intolerance by adjusting the volume of the balloon between 400 and 800 mL, and then to increase volume as weight loss plateaus. The Spatz balloon is intended for 12 months implantation and is currently being used outside the United States. Issues with the Spatz balloon include deflation leading to the need for surgical extraction.4 The adjustable nature of this balloon and the longer implant time make this device more attractive than the other balloons. It is not yet FDA approved but is in multicenter trial in the United States.

**Devices No Longer Available**

The Silimed is a smooth transparent silicon shell with saline that is placed by endoscopic traction under direct visualization and rolled up inside a thin silicone sheath anchored to the tip of the endoscope with a snare. This is removed as an entire system held in an overtube device. The literature identifies this as being a safer technique done with direct visualization under conscious sedation.10 The Medsil balloon is saline filled with maximum volume of 700 mL placed similarly to the Orbera and Silimed.1,3 It is not offered in the United States but is available in Europe.

**Air- or Gas-Filled Balloons**

**Obalon**
The Obalon is the only FDA-approved gas-filled balloon. This device can be placed in the office as it is swallowed as a gel that sits at the end of thin tubing. The device is filled with a proprietary gas and checked for placement using fluoroscopy. Therefore, only one endoscopic procedure is required for removal. However, fluoroscopy is needed to verify placement, and therefore is an extra radiation exposure for the patient. Each balloon is 250 mL in volume of gas and up to three balloons are placed over a 6 to 12 weeks period if weight loss plateaus. This sequential addition of volume is thought to be better tolerated by patients with less nausea. The reported weight loss is ~5 kg for patients.11

**Heliosphere**
The Heliosphere BAG is a double bag polymer balloon covered with silicon envelope placed for 6 months. It is filled with air, and is stated that it may reduce the risk of digestive intolerance as compared with fluid-filled balloons. The balloon has a high rate of system failure at positioning, spontaneous deflation, absence of marker, and large size with balloon that isn’t compliant, and causes intolerance to patients.1,12

**Elipse**
The Elipse balloon (Elipse Balloon, Allurion Technologies, Natick, MA) is a very exciting new technology. This device does not require procedures for placement or removal. The Elipse balloon is swallowed in the office and the valve dissolves over time. Once the balloon deflates, it passes through the bowels and is not thought to be associated with bowel obstruction. Currently, this device is in clinical trials. A device such as this may revolutionize the nonsurgical approach to weight loss. Because no procedures are required, this allows several providers interested in weight management to have a new tool they may be able to offer their patients. The data so far is promising. An Italian study showed early results with mean %EWL of 26% at 4 months. No problems with passage of the balloon were seen in these 38 patients.13 Another balloon in development is the Endoball. This device can be filled with both fluid and air between 500 and 800 mL.1,2 No data are available at this time.
Complications with the Balloons

Balloons have recently come under fire as complications are coming to light. The FDA has cautioned against their use due to deaths that have been reported with the balloons.\(^\text{14}\) Pancreatitis, perforation, ulcer formation, rupture/deflation of device possibly leading to outlet obstruction or bowel obstruction, and bleeding are reported complications. The deaths have occurred primarily with fluid-filled balloons. A recent statement by Tate and Geliebter notes 33 deaths related to the Orbera and Reshape balloons since 2006.\(^\text{15}\) The FDA Web site can be visited for further information. Overall none of the deaths have resulted in removal of the devices from the market. However, they do caution the interventionalist to stay within the guidelines with regard to implantation time, attention to prior gastric procedures, compliance of the patient, large hiatal hernia, volume placed in the balloons, and BMI of the patient. Both ASMBS (American Society for Metabolic and Bariatric Surgery) and SAGES (Society of American Gastrointestinal and Endoscopic Surgeons) have issued statements stating the devices are safe if used within the guidelines.\(^\text{16}\)

Advantages of the Balloons

The balloon can not only be used for primary weight loss procedures but also can be used as a bridge procedure to other weight loss surgeries. One of the uses is to decrease anesthetic risk in severely obese patients planning to undergo bariatric surgery by getting their BMI to a more operable weight.\(^\text{17}\) Zerrweck et al. looked at patients with BMI greater than 60 kg/m\(^2\) who received balloons. These endoscopic procedures were associated with shorter operative time and lower overall risk of adverse outcomes as compared with surgery in this population.\(^\text{6}\)

Unfortunately, this advantage of the balloon cannot be realized in the United States as it is only FDA approved for BMI 30 to 40. The balloons may also be helpful in those patients who need to lose weight but are poor surgical candidates due to prior surgeries or high anesthesia risks. Again in the United States, this would not be possible for BMIs over 40. In light of the recent FDA warnings, these off-label uses should only be done in a study that is monitored by an institutional review board.

**Table 1** Intragastric balloons: Basic facts regarding implantation time, method of placement, and removal and availability in the United States

<table>
<thead>
<tr>
<th>Balloon and composition</th>
<th>FDA approval/Indications</th>
<th>Approved typical time for implantation</th>
<th>Placement and retrieval method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Orbera (BIB) Apollo Endosurgery Silicone sphere filled with 400–700 mL of saline with methylene blue solution</td>
<td>FDA approved BMI 30–40 km/m(^2) Age ≥ 20</td>
<td>6 months</td>
<td>Placed and removed with endoscopy</td>
</tr>
<tr>
<td>Reshape-integrated dual balloon system ReShape Medical Two silicone spheres attached in tandem, each filled with 375–450 mL of saline with methylene blue solution</td>
<td>FDA approved BMI 30–40 km/m(^2) and single obesity related comorbidit Age 22–60</td>
<td>6 months</td>
<td>Endoscopic placement and removal</td>
</tr>
<tr>
<td>Obalon Balloon System Obalon Therapeutics Up to 3 separate sphere plastic polymer filled with 250 mL of inert gas</td>
<td>FDA approved BMI 30–40 km/m(^2) Age ≥ 20</td>
<td>6 months</td>
<td>3 balloons placed in 1 month increment (within 12 weeks) by swallowing capsule under fluoroscopy. Removed with endoscopy</td>
</tr>
<tr>
<td>Eclipse Balloon System Allurion Technologies Balloon sphere made of proprietary film, filled with 550 mL of saline</td>
<td>Not FDA approved</td>
<td>4 months</td>
<td>Single balloon swallowed under fluoroscopy. Valve release in 4 months in situ and passed via GI tract when deflated</td>
</tr>
<tr>
<td>Spatz Balloon System Spatz FGIA Saline filled silicone balloon sphere with an attached catheter to adjust fill volume after implantation</td>
<td>Not FDA approved</td>
<td>12 months</td>
<td>Endoscopic placement, adjustment, and removal</td>
</tr>
</tbody>
</table>

Abbreviations: BMI, body mass index; FDA, US Food and Drug Administration; GI, gastrointestinal.
Table 2 Expected weight loss based on FDA-approved primary procedure

<table>
<thead>
<tr>
<th>Type of primary procedure</th>
<th>Weight loss</th>
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<tbody>
<tr>
<td>Roux en Y gastric bypass</td>
<td>70% excess body weight</td>
</tr>
<tr>
<td>Vertical sleeve gastrectomy</td>
<td>50% excess body weight</td>
</tr>
<tr>
<td>Lap band</td>
<td>Up to 30% excess body weight</td>
</tr>
<tr>
<td>Balloon</td>
<td>10% excess body weight</td>
</tr>
<tr>
<td>Endoscopic sleeve gastoplasty</td>
<td>20% excess body weight</td>
</tr>
<tr>
<td>Aspire assist</td>
<td>25–30% excess body weight</td>
</tr>
</tbody>
</table>

Abbreviation: FDA, US Food and Drug Administration.

raised some controversy regarding the results.\textsuperscript{20} This brings up the question of expectation. Should a balloon that is implanted for only 6 months be expected to have long-term efficacy, or should it be considered as an addition to the other weight management programs out there such as Jenny Craig and Weight Watchers? (\textit{\textsuperscript{-Table 2}}) The debate remains as to whether these devices will result in long-term weight loss. However, the short-term results are promising.

Other Types of Space-Occupying Devices

The concept for a semistationary antral balloon was first published in 2008. This is a pear-shaped silicone balloon 30 cm duodenal stem and 7 g metallic counter weight at the distal end. The conical end sits in the antrum and the distal weight utilizes peristalsis to come across the pylorus and anchor this device into place. The goal is to occlude the pylorus, intermittently prolong gastric emptying, and stimulate the satiety receptor.\textsuperscript{21} This has since evolved into the transpyloric shuttle which is endoscopically placed. It is a larger balloon, \( \sim 56 \) mm in size, and sits in the stomach. The smaller balloon, \( \) mm \( \times 96 \) mm, is tethered to a small weight which sits in the duodenum. This allows for a normal peristalsis to help occlude the pylorus. Using this balloon, the Endobesity I trial showed weight loss without plateau in 3 months. It showed in the 3 months Endobesity I trial showed weight loss without plateau on peristalsis to help occlude the pylorus. Using this balloon, the weight which sits in the duodenum. This allows for a normal removal.

The Restore Suturing System by Davol/BARD has the ability to place deeper sutures without having to withdraw device for suture reloading. Transoral gastric volume reduction as an Intervention for Weight Management trial showed 27.7 \( \pm 21.9 \) EWL at 12 months follow-up. It requires an overtube and therefore intubation and general anesthesia. The procedure time is \( \sim 45 \) minutes. The EWL was 50% and less obese had a better response.\textsuperscript{24} This device is not yet FDA approved. Yet another independent platform is used for the Primary Obesity Surgery Endoluminal for tissue approximation and an endoscopic sleeved and is made by USGI Medical. This is still being studied.

The Overstitch device is the most successful device utilized for sutured gastropasty and its use is increasing. The device is able to take full-thickness sutures and therefore is considered to place sutures that are more durable than the other devices mentioned. This device is FDA approved and is in widespread use among surgical endoscopists. A single center pilot study showed feasibility for endoscopic sleeve gastropasty (ESG) with a running suture. Six to 12 stitches are placed in a triangular fashion at the anterior wall, greater curve, and posterior wall and started at the fundus. Since that study, most follow the modification by Kumar et al with suturing initiated at the antrum and running proximally toward the fundus. Interrupted sutures can be used to reinforce the sleeve.\textsuperscript{25} The BMI decrease was 34.2 kg/m\(^2\) to 29.4 kg/m\(^2\) with mean weight loss of 13.1 \( \pm 1.3 \) kg after 1 year. More and more data are coming forth from around the world surrounding the ESG. A recent study combining data from the United States and Australia showed respectable weight loss. In total, 112 consecutive patients (male 31%, age 45.1 \( \pm 11.7 \) years, baseline BMI 37.9 \( \pm 6.7 \) kg/m\(^2\)) underwent ESG. At 1, 3, and 6 months, change in weight was 9.0 \( \pm \) 4.6 kg (total body weight loss [TBWL] 8.4 \( \pm \) 4.1%), 12.9 \( \pm \) 6.4 kg (TBWL 11.9 \( \pm \) 4.5%), and 16.4 \( \pm \) 10.7 kg (TBWL 14.9 \( \pm \) 6.1%), respectively. Adverse events were noted in 2.7%.\textsuperscript{26}

Endoscopic Suturing Devices

Endoscopic Sleeve Gastropasty

Conceptually, these procedures have been in development for a long while. Endoscopically, sutures are placed so that the remaining lumen of the stomach emulates a vertical sleeve gastrectomy. One of the first endoscopic suturing devices was the Endocinch (Bard). This device was initially developed to treat gastroesophageal reflux disease and then was extended for suturing for weight loss. Fogel first showed weight loss results comparable to traditional bariatric procedures with up to 21.1% EWL at 1 month, 39.6% EWL at 3 months, and 58.1% EWL at 12 months of follow-up. The durability of the plications was in question due to plications not being full thickness.\textsuperscript{23}

The EndoCinch device was the first endoscopic suturing device approved by the FDA in 2008. The device was able to place deeper sutures without having to withdraw device for suture reloading. Transoral gastric volume reduction as an Intervention for Weight Management trial showed 27.7 \( \pm 21.9 \) EWL at 12 months follow-up. It requires an overtube and therefore intubation and general anesthesia. The procedure time is \( \sim 45 \) minutes. The EWL was 50% and less obese had a better response.\textsuperscript{24} This device is not yet FDA approved. Yet another independent platform is used for the Primary Obesity Surgery Endoluminal for tissue approximation and an endoscopic sleeved and is made by USGI Medical. This is still being studied.

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When compared directly to sleeve gastrectomy in a case-matched study, the sleeve showed 6% more EWL with a higher incidence of new onset reflux.\textsuperscript{27}

**Transoral Gastroplasty**
A device that is no longer available is the stapler for transoral gastroplasty (TOGA). A set of transoral flexible endoscopically guided staplers are used to create a gastric sleeve along the lesser curve of the stomach. The procedure requires two operators and a 60 Fr. Savory dilator. A flexible stapler is passed peroral in an over the wire technique. A 7 to 8 cm stapled apposition is made from the cardia to lesser curve and then followed distally. Adverse events within the first week included nausea, vomiting, pain, and transient dysphagia. The reported mean % EWL were 16.2% at 1 month, 22.6% at 3 months, and 24.4% at 6 months. Gaps were noted in the staple line in 13 patients at 6 months follow-up on endoscopy. A second human trial with closer staple apposition and use of steroid and diclofenac had better results with a % EWL at 19.2% at 1 month, 33.7% at 3 months, and 46% at 6 months. The TOGA trial by Familiari et al demonstrated safety and efficacy at 1 year. The mean starting BMI was 41.5 kg/m\textsuperscript{2} and had a % EWL of 29.3% at 3 months, 36.8% at 6 months, and 38.7% at 12 months.\textsuperscript{28} Minimal complications, such as respiratory insufficiency and an asymptomatic pneumoperitoneum, were reported and were treated conservatively. Staple line gaps incidence remained a problem; however, the gaps were reported to be small. Another limitation of the study was the variable weight loss counseling provided to the patients at the different sites.\textsuperscript{1,29} The technology was novel; however, the implementation of the procedure, surrounding counseling, and the expectation of equivalence to surgery may have been miscalculated.

**TransEndoscopic Restrictive Implants System**
TransEndoscopic Restrictive Implants System mimics the gastric band. With this device, a prosthetic diaphragm with a 10 mm orifice is stapled into the gastric cardia. This is designed to be permanent but can be easily removed or modified. The trial resulted in one patient with gastric perforation requiring surgery and removal of device, and two with pneumoperitoneum that could be managed non-operatively. The % EWL was 12.3% at 1 month and 22.2% at 3 months.\textsuperscript{1,2} The high complication rate resulted in the device being returned to the design phase.

**Malabsorptive Endoluminal Procedures**

**EndoBarrier**
The duodenal jejunal bypass sleeve, The EndoBarrier (GI Dynamics, Lexington, MA), is one of the first endoluminal malabsorptive procedures introduced. The device was created to have malabsorption similar to that of the Roux-en-Y gastric bypass, with the primary goal of improving diabetes. A nitinol anchor sits in the duodenal bulb with a 60 cm fluoropolymer sleeve that prevents mixing of bile and pancreatic fluids with food, thereby limiting absorption in the proximal intestine. The theory is that the accelerated delivery of chyme to the distal gastrointestinal tract allows for early release of GLP-1 and incretins which have roles in glycemic control and energy homeostasis. In theory, this should help with weight loss and control of type 2 diabetes mellitus. The first human trial involved 12 patients who were left implanted for 12 weeks. Mean % EWL at 12 weeks was 23.6%. Complications were two reports of a mucosal tear and an oropharyngeal mucosal tear not requiring intervention. Three of the four patients with diabetes mellitus were weaned off medications within 24 hours of implantation. He subsequently followed up his study looking closer at the hormonal effects and concluded the GLP-1 is not affected.\textsuperscript{30}

A second trial by Tarnoff et al compared the device (25 patients implanted) versus diet control (15 patients). Eighty percent of patients kept their implant in place for 12 weeks, and adverse events included gastrointestinal bleed, anchor migration, and stent obstruction. %EWL at 12 weeks was 22% for device versus 5% for control.\textsuperscript{31,32} The early European experience has similar results.\textsuperscript{33} Multiple complications were noted in all of these trials including slippage (due to sharp angulation of the duodenum), mucosal tears, bleeding, liver abscess, cholangitis, cholecystitis, and esophageal perforation.\textsuperscript{33} Ultimately, the high incidence of liver abscesses in the US trial led to early termination of the trial and the device remains in redesign.

**ValenTx**
Gastroduodenojugal bypass sleeve (ValenTx, Inc., Hopkins, MN) is a sleeve anchored from gastroesophageal junction and extends from the proximal stomach into 120 cm of small bowel, thereby blocking absorption of food from the stomach to the jejunum. It mimics the malabsorptive properties of a gastric bypass. Post-procedural dysphagia is a common complication. The effectiveness of this procedure was first evaluated in 13 patients (mean BMI of 42 kg/m\textsuperscript{2}). The mean % EWL was 54% for 10 patients who kept the implant for 1 year. Four patients with partial cuff detachment had less weight loss. Comorbidity improvement was noted with diabetes mellitus (DM), hypertension, hyperlipidemia, and use of DM medications.\textsuperscript{34}

The device is currently in trials in the United States. The device has been placed mostly with a hybrid endoscopic and laparoscopic approach. A fully endoscopic approach is in development.

**Mucosal Resurfacing**
This technology is truly metabolic and involves ablation of the duodenal mucosa. The Revita DMR catheter (Fractyl, Lexington, MA) is passed into the duodenum under endoscopic visualization, and after a circumferential saline lift distal to the ampulla, a 2 cm balloon is inflated in the lumen. A burst of energy in the coils of the balloon ablated the surface of the duodenal mucosa. Ablation is performed to the desired length. Improved glycemic control is noted almost immediately with the theory of ablaing diseased mucosa that will hopefully regenerate with normal mucosa. Adverse events include duodenal stenosis and abdominal pain.\textsuperscript{2,35} This technology is promising and is providing more insight into mechanisms of diabetes and obesity.
Aspiration Therapy

Aspiration therapy is extremely controversial and has been likened to controlled bulimia. However, when the device and its success are examined closely, one can conclude that it is a viable option for the right patient for weight management that can be controlled, not disruptive to a patient’s chemistries and effective for ongoing weight loss. As opposed to other endoscopic weight loss techniques, this device is considered more durable as it can remain implanted in the stomach for an indefinite period of time. A gastrostomy tube

Fig. 1 (A–I) Gastric endoscopic bariatric therapies.
is placed traversing the abdominal wall from the lumen of the stomach to the surface of the skin. Patients are counseled on mindful eating. Then 20 minutes after eating, the patient syphons 30% of the stomach contents, leading to less caloric absorption and weight loss. Placement and removal are under conscious sedation with closure similar to a PEG tube. In a randomized trial for the treatment of Class 11 and Class 111 obesity, aspiration therapy was noted to be significantly more successful than lifestyle changes alone: AspireAssist group with 31.5 ± 26.7 of their excess body weight (12.1 ± 9.6 total bodyweight), whereas those in the lifestyle counseling group had lost only a mean of 9.8 ± 15.5 of their excess body weight (3.5 ± 6 total body weight) (p < 0.001).\(^{36}\) One of the longest longitudinal studies had 201 participants with 4 years follow-up. The starting BMI was 43.6 ± 7.2 kg/m\(^2\) and the mean %TWL at 1, 2, 3, and 4 years, respectively, was 18.2 ± 9.4, 19.8 ± 11.3, 21.3 ± 9.6, and 19.2 ± 13.1. Clinically, significant reductions in glycated hemoglobin A1c (HbA1c), triglycerides, and blood pressure were observed. For participants with diabetes, HbA1c decreased by 1% (p < 0.0001) from 7.8% at baseline to 6.8% at 1 year.\(^{37}\) Although FDA approved for years, the adoption of this therapy has been slow.

**Magnamosis**

Magnets are utilized to create intestinal anastomoses so that a gastric bypass can theoretically be performed endoscopically. These are semi-assembling magnets that are placed using upper and lower endoscopy with fluoroscopic guidance for dual path enteral bypass in the small bowel (incisionless magnetic anastomosis system). Once the magnets connect, necrosis creates a fistula opening. Transient nausea and diarrhea were reported and resolved. A small study reported 10.6% EWL at 6 months.\(^{2,38}\) This is currently undergoing evaluation in the United States but is not FDA approved.

**Full Sense Device**

The Full Sense Device is placed via endoscopy in the distal esophagus and proximal stomach. By applying pressure to this area, it causes satiety without the need for the presence of food. At this time, it is not available for commercial use anywhere but is in preclinical trials.

**Conclusion**

Endoscopic approaches to weight loss are gaining popularity. Many technologies are currently being used such as the space filling devices and ESG (►Fig. 1). Novel approaches to primary bariatric surgery are being developed and are very promising. Some technologies have not withstood the test of time. In general, it appears these endoscopic approaches are here to stay. What is not clear is the expected success and durability of these procedures. There is a gap in therapy between diet/exercise and surgery for weight loss. These endoscopic procedures can provide an alternative to those patients with Class 1 and 2 obesity. They can even be used for higher class obesity for high-risk patients, those with multiple abdominal surgeries and as a bridge to other weight loss surgery. Currently, these devices are in greater use outside the United States than within it. A large barrier to these procedures in the United States is cost. In the United States and in many countries, these endoscopic approaches are not covered by insurance. Many of these approaches also require expensive devices, general anesthesia, and skilled endoscopists. These requirements are barriers to their widespread use.

What is known is that obesity has reached epidemic proportions worldwide. Level 1 evidence for many of the endoscopic procedures for weight loss is lacking. However, several case series are in the literature and speak to weight loss that is greater than lifestyle changes alone but may be less than surgery with a lower risk profile. Ongoing studies are needed with large numbers and comparative trials.

Another debatable topic with these endoluminal approaches is the definition of success. Should these endoscopic techniques be expected to be as durable and effective as surgery? Or should these approaches be considered another tool in the toolbox for weight loss? Can they be put in a category that allows more pounds lost than programs such as WeightWatchers and Jenny Craig, but less than surgery? These technologies can fit the void between diet/exercise and surgery. Both SAGES and ASMBS support the development of these procedures and recognize their need in the fight against obesity.\(^{39}\)

We know obesity is a multidisciplinary disease and multiple approaches need to be available to our patients to be successful. All of these endoscopic approaches should be offered in a comprehensive weight loss program that includes dietary and lifestyle counseling. Adequate behavioral support is also helpful. As the epidemic of obesity grows, endoluminal therapies for metabolic surgery will provide a pivotal role both as primary and metabolic modalities in the bariatric surgeon/endoscopists armamentarium.

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