Introduction

Gastroesophageal reflux disease (GERD) is one of the most prevalent gastrointestinal (GI) diseases in the US with an estimated prevalence of 18.1–27.8% (1). The socioeconomic burden of GERD is tremendous with an estimated health care cost of $12.3 billion dollars annually. It is the most common reason for an outpatient GI appointment (8.9 million outpatient visits per year) and the primary reason for an upper endoscopy (2,3). In the US, it is estimated that approximately 40% of the entire population will experience heartburn at least monthly, 14% will experience weekly episodes, and 7–10% will have daily symptoms (4).

GERD was initially defined by the Montreal Consensus as “a condition which develops when reflux of stomach contents cause troublesome symptoms and/or complications” (5). Many underlying causes of GERD exist [defective lower esophageal sphincter (LES), impaired gastric emptying, failed esophageal peristalsis], and the disease is associated with a broad spectrum of symptoms, both...
typical esophageal symptoms (heartburn, regurgitation, and dysphagia) and extra-esophageal atypical symptoms (laryngitis, cough, asthma, and dental erosions) (4). While GERD itself is benign, disease progression can lead to harmful sequela including erosive gastritis, Barrett’s esophagus (BE), and esophageal malignancy. From a surgical perspective, GERD is considered a physiologic disorder of the gastroesophageal junction (GEJ), the natural anti-reflux barrier. When functioning properly, the LES, the crural diaphragm, and the geometry of the GEJ all aid in preventing the reflux of stomach acid into the esophagus.

The most common treatment for GERD is a step-up approach starting first with lifestyle modifications (i.e., weight loss; avoiding alcohol, spicy foods, and eating before lying down; and elevating the head of the bed) (6). After failing lifestyle modifications, patients are typically started on a H2 receptor antagonists (H2RA) or proton-pump inhibitors (PPI) with symptomatic relief expected in 60% and 83% of patients, respectively (4). However, an alternative to medical therapy, and often the next approach offered by clinicians, remains anti-reflux surgery. Patients with objective evidence of reflux disease may proceed with surgical intervention if they fail medical management (persistent or inappropriately controlled esophageal or extra-esophageal symptoms), prefer surgical management despite successful medical management, or suffer complications of GERD (BE or stricture) (5). A variety of surgical modalities are available today, including laparoscopic anti-reflux surgery (LARS), robotic assisted laparoscopic fundoplication (RALF), and endoscopic anti-reflux therapy. This review will provide a comprehensive assessment of each of these procedural interventions, assess their respective advantages and disadvantages, and appraise the current data available regarding each of the modalities effectiveness to treat GERD.

**Laparoscopic treatment of reflux disease**

*Introduction of laparoscopic anti-reflux surgery*

The first fundoplication, described as a complete 360° wrap of the stomach fundus around the esophagus, was performed by Dr. Rudolf Nissen in the 1950s. Other modifications soon followed and are named after their creators: Dor, Toupet, Belsey, Hill and Collis. These open techniques remained the standard of care until 1991 when Dr. Dallemagne published his first series utilizing the relatively new minimally invasive technique, laparoscopy, to perform a 360° wrap of the stomach (7). LARS soon became the standard of care with reduced short-term morbidity, shortened postoperative length of stay, and decreased incisional hernia rates (8). The minimally invasive approach made fundoplication a more acceptable treatment option for patients and contributed to the tendency to offer surgery earlier in the disease course (9). Multiple randomized clinical trials and meta-analyses have found LARS to be safe and effective with reduced perioperative morbidity as compared to its open counterpart. In 2009, a meta-analysis of 12 randomized clinical trials comparing open anti-reflux surgery to LARS found that patients who underwent LARS had a reduction in total length of hospital stay by 2.7 days, returned to normal activity an average of 7.8 days sooner, and had a 65% reduction in their postoperative complication rate (10). Similar results were found by Catarci et al. (11). In a pooled meta-analysis of multiple randomized control trials, the authors found that LARS was associated with a significantly lower operative morbidity rate, a shorter postoperative stay, and shorter patient sick leave. Importantly, there was no significant differences in the incidence of reflux recurrence, dysphagia, bloating and reoperation rates between the two approaches and no perioperative deaths were recorded (11). Additionally, laparoscopy has several non-GERD-related long-term advantages over open surgery. Incisional hernias can have a tremendous impact on patient’s quality of life, and in a 17-year follow-up study examining open fundoplication vs. LARS, Oor et al. found a significantly decreased rate of re-intervention for incisional hernias in the patients treated with laparoscopy (14% vs. 2%) (8).

While a step by step technical guide is outside the scope of this review, there are several technical approaches to a laparoscopic Nissen fundoplication that have been shown to improve postoperative outcomes. Based on a consensus of 40 expert foregut surgeons and published by the SAGES Guidelines Committee in 2010, the recommended surgical technique includes some of the following elements: (I) opening the phrenoesophageal ligament in a left-to-right fashion, (II) circumferential dissection of the hiatus, (III) sufficient transhiatal mobilization to allow approximately 3 cm of intraabdominal esophagus, (IV) short gastric vessel division to allow for a tension free wrap, (V) crural closure posteriorly with non-absorbable sutures, (VI) creation of a 1.5–2.0 cm wrap incorporating the anterior muscular wall of the esophagus, and (VII) bougie placement at the time of the fundoplication if a complete wrap is formed (5). A partial fundoplication has also been described as an...
effective and accepted alternative for anti-reflux surgery (12). Another laparoscopic anti-reflux procedure option, which has gained popularity for the treatment of reflux disease is the magnetic sphincter augmentation system, LINX® (Torax Medical LLC; Ethicon US, LLC) (12,13). Further details of these procedures, preoperative diagnosis and workup, intraoperative technical considerations and postoperative care and complications are discussed in other chapters of this edition.

**Medical vs. laparoscopic anti-reflux surgery**

Medical therapy has been successful in the treatment of GERD, and maintenance therapy with PPIs can be used to afford patients high rates of symptom resolution and esophageal healing. Developed in the late 1980’s, omeprazole (the first PPI) quickly became a mainstay for GERD treatment with usage rates doubling from 1999 to 2012 (14). GERD, however, is a chronic disease, and patients are often reluctant to take medications for their entire lifetime. Furthermore, since the implementation of PPIs, understanding of potential adverse effects of long-term PPI use has grown considerably and increased effort has been placed on finding potential alternatives to PPI therapy (15). Since this time, LARS has become established as an effective alternative to long-term PPI therapy. Multiple randomized control trials and meta-analyses have compared the results of medical and surgical therapy. Most have supported surgery as an effective alternative for both the treatment of GERD in patients with good symptom control and for those with only a partial response to medication. The LOTUS trial, published in 2011, was a 5-year randomized, multicenter control trial comparing optimized PPI therapy vs. standardized LARS in patients with chronic GERD (16). While the estimated remission rate in the PPI therapy group was 92% vs. 85% in the LARS group, prevalence and severity of heartburn and acid regurgitation was lower in the LARS group (16). However, there was significantly higher rates of dysphagia, bloating and flatulence with LARS. As the trial’s primary outcome measure was time to treatment failure, the authors concluded that both drug and surgically induced acid suppression allow patients to achieve and maintain satisfactory disease remission at 5 years (16).

Many studies have demonstrated objective evidence for LARS effectiveness. Based on manometric data and impedance measurements, fundoplication results in both less esophageal acid exposure and significantly increased LES pressures when compared with medical therapy. In a large, matched randomized clinical trial, Mahon et al. compared LARS to PPI therapy. At 3 months, mean DeMeester scores significantly improved from 42.7 to 8.6 in the LARS group and from 36.9 to 17.7 in the PPI group (17). There was also a significant increase in mean gastrointestinal symptom score and general well-being scores at 12-months in the LARS group as compared to patients randomized to PPI therapy (17). Additionally, a prospective, randomized open parallel-group, multicenter trial that compared the efficacy and safety of LARS to PPI therapy demonstrated that esophageal acid exposure was significantly reduced in the LARS group both at 6 months and 5 years (18).

Data also demonstrates that surgical treatment of GERD is effective and safe in the long-term. In a long-term follow-up study of a prospective randomized trial comparing medical and surgical treatment, Spechler et al. found that statistically fewer patients in the surgical group were using anti-reflux medications (62% for surgical treatment vs. 92% for medical treatment) at follow-up (9.1 years for surgical group and 10.6 years for medical treatment) (19). However, there was no significant difference in the grade of esophagitis, the frequency of esophageal stricture treatment, and the overall satisfaction with anti-reflux therapy between the two groups (19). The longest outcome data comes from Oor et al. (8). To compare conventional open surgery vs. LARS, they evaluated a total of 111 patients (60 LARS and 51 open fundoplication) 17 years after their initial operation. Both groups showed excellent long-term outcomes. However, they found that, compared with their preoperative symptom scores, patients in the LARS group continued to report significant improvement in general health and quality of life and 90% of patients reported continued symptom relief. While, at 17 years, 42% of LARS patients were dependent on the daily use of acid suppression medications, medication usage was significantly lower as compared with their preoperative dose (8).

LARS has been shown to be safe and effective relieving GERD symptoms and improving quality of life. However, it is still unclear if LARS is more effective than medical therapy at inducing remission of BE or esophageal dysplasia. BE occurs when the esophageal lining undergoes metaplastic change from the normal squamous cell epithelium to gastric columnar epithelium. BE significantly increases the risk of developing esophageal adenocarcinoma, and as a result, adequate treatment of BE is of utmost importance as GERD becomes more prevalent in the US. A non-randomized prospective study by Rossi et al. compared...
patients with low grade dysplasia (LGD) who were treated with high dose PPI therapy vs. LARS (20). They reported a significantly improved rate of regression of LGD to BE at 12 months in the surgical group (94%) vs. patients treated with high dose PPI therapy (63%). At 18 months, all LARS patients had continued confirmed absence of LGD (20). A matched cohort study out of Italy enrolled 33 patients with BE or LGD, 20 of whom underwent LARS and 13 who were treated medically. Not only was LARS associated with a better control of both acidic and weakly acidic reflux, it was also associated with a higher probability of LGD reversal (21). Smaller retrospective case series have also looked at the rate of regression of BE following LARS. A case series out of the Czech Republic examining 50 patients with BE demonstrated that 38% of patients following LARS had no detectable disease postoperatively. However, in their same series, 36% of patients had unchanged disease and 10% ultimately had disease progression (22). A larger retrospective case series by Morrow et al. showed a regression rate of 22% in patients who underwent LARS and a progression rate of 7% (23). The evidence of improved regression of BE following LARS or medical therapy remains inconclusive, and it remains clear that LARS does not alter the need for continued endoscopic surveillance in these patients. High-grade dysplasia should continue to be treated by endoscopic therapy to achieve complete histological eradication before anti-reflux surgery is attempted and esophageal adenocarcinoma should be treated by a multidisciplinary approach with esophagectomy, chemotherapy and radiation according to oncological standards (5).

Refractory GERD

While LARS is validated for patients who respond to PPI therapy, the data of treating PPI non-responders is mixed. Despite the high efficacy of PPI therapy, clinical failure remains common and can occur in around 20–45% of patients. Furthermore, poor response to PPI is associated with a negative impact on physical and mental-health quality of life (24). As such, treating refractory reflux can be challenging and frustrating for both the patient and clinician. For a substantial portion of these patients, anti-reflux surgery should be considered. The differential diagnosis for reflux disease is broad, however, and not all patients will be helped by surgical management. Clinicians should, therefore, carefully select patients for surgical therapy (25). Of note, while not specifically addressed in this review, there are several other indications for LARS including, but not limited to, biliary reflux disease, laryngopharyngeal reflux disease, and hiatal hernias.

Anvari and Allen were the first investigators to show that poor responders to PPI therapy can benefit from LARS and demonstrated that patients who underwent LARS had significant improvement in postoperative symptom scores and quality of life scores that correlated with lower esophageal acid exposure (26). They studied a group of 719 patients, all of whom had inadequate response to PPI therapy (<70% relief on visual analogue scale), and found a significantly improved postoperative quality of life for both physical health and mental health (26). Other groups have shown fewer promising outcomes. Wilkerson et al. found that while good and poor responders had a significant decrease in symptom score following LARS, poor responders had a lower percentage of “excellent” or “good” surgical outcomes, and many patients continued to report severe heartburn (27). In a prospective study of 370 patients who underwent LARS (296 PPI responders and 74 non-responders), Hamdy et al. found that good responders had a greater reduction of heartburn and regurgitation symptoms (24). Patient satisfaction with surgery was also significantly better in the good responder group. Postoperative assessment did not reveal any significant differences on esophageal manometric testing, LES pressures, or 24-h pH monitoring (24).

Laparoscopy challenges

The widespread acceptance of laparoscopy has offered multiple improvements for patient outcomes including shorter hospital stay, less postoperative pain, and improved long-term quality of life. Several disadvantages of laparoscopic surgery quickly became apparent including loss of 3D visualization, reduction in haptic feedback, mechanical constraint secondary to the “fulcrum effect,” instrument rigidity, tremor enhancement, and decreased range of motion (28,29). Laparoscopy has also been associated with new surgeon discomfort and ergonomic challenges. Previous studies have shown that even experienced laparoscopic surgeons suffer from significantly increased upper extremity muscle discomfort and physical work (30). Several factors contribute to the surgeon’s musculoskeletal stress, including prolonged static head and trunk posture, a greater amount of shoulder and upper arm movements while using laparoscopic instruments, and poor mechanical design of laparoscopic instruments.
Patient characteristics have also contributed to differences in ergonomic stress. For example, a patient’s increased body mass index (BMI) adds increasing difficulty to a procedure. Additionally, laparoscopy relies on the experience and the knowledge of a surgical assistant and incorrect maneuvering can lead to poor visualization, increased operating room times and intraoperative complications (31). The challenges associated with laparoscopy paved the way for the development of surgical robotic systems, which seek to resolve some of these difficulties.

**Robotic surgical treatment of reflux disease**

**History of robotic surgery**

Robotic surgery was developed to address many of the previously identified challenges with laparoscopy, and its use is now becoming widespread in anti-reflux surgery. The robotic platform confers full control of the camera and instruments to the primary surgeon, improving surgeon ergonomics, visualization, and autonomy (31). While a number of early systems were developed, the predominant system for abdominal surgery at the time of this writing is the da Vinci Surgical System (dVSS) (Intuitive, Incorporated; Sunnyvale, CA) (Figure 1). Robotic technology has become pervasive throughout modern surgical practice and has been rapidly incorporated into many specialties such as gynecology, urology, colorectal, and bariatrics, and now ever increasingly, foregut surgery (32). In the year 2000, there were only 18 robotic surgery systems worldwide. As of March 2019, the number of available systems worldwide has skyrocketed to 5,114 and the total number of robotic-assisted procedures surpassed 5 million (33).

Commonly-used robotic surgical platforms are equipped with a variety of technologies that improve the precision and efficiency of complex surgical task performance compared to traditional laparoscopic surgery (34-38). Robotic platforms employ wristed instruments that provide surgeons with seven total degrees of freedom when performing surgical tasks, an increase compared to the four degrees of freedom provided by traditional rigid laparoscopic instrumentation (39). Commonly-used robotic platforms are paired with a high-definition three-dimensional camera that allows for improved precision and visualization of surgical targets when compared to the two-dimensional camera platforms used in laparoscopy (34-36). In addition, the surgeon is always in control of the camera, able to obtain the visualization required without added communication and guidance from a surgical assistant. The surgeon’s console allows for scalability of a robotic surgeon’s movements up to 3:1 to allow for finer movement at the instrument tip than traditional laparoscopic instrumentation and provides the added benefit of tremor elimination (39).

The laparoscopic operating room environment has many ergonomic challenges that robotic surgical systems seek to address (40-42): suboptimal operating table and monitor positioning, non-ergonomic instrument handles,
and maintenance of awkward body positioning for extended periods of time (43,44). While using a robotic surgical platform, the operating surgeon is in a seated position viewing the procedure through a viewfinder on the console and manipulating instruments using lightweight masters. These features provide ergonomic benefits to operating surgeons and subjective improvements in ergonomic stress (45-47). In a quantitative comparison of ergonomic stress associated with laparoscopic and robotic assisted laparoscopic procedures, robotic platforms were associated with significantly decreased activation of multiple muscle groups: biceps, triceps, and deltoids bilaterally (48).

However, the robotic platform has several practical and technical limitations that affect their use. Robotic platforms provide operating surgeons with significantly less haptic feedback than is provided during conventional laparoscopy. Additionally, most robot-assisted surgical platforms restrict surgical activity to only part of the body cavity (although, the latest robotic platforms aim to address this restriction and are equipped for a wider field of surgery) (37). Furthermore, robotic surgery is associated with somewhat of a learning curve, such that surgeons less familiar with robotic surgery may initially experience a temporary slowing of their skills that can subsequently impact their clinical performance (49). Lastly, there is generally an increased upfront equipment cost associated with the robotic platform compared to conventional laparoscopic procedures, with a variable impact on length of hospital stay or operative time (50).

**Robotic surgery vs. laparoscopic surgery—patient outcomes**

Robotic-assisted laparoscopic fundoplication (RALF) has been demonstrated to be safe and feasible with similar short-term outcomes to LARS (51-55). In a randomized clinical trial by Draaisma et al. comparing 50 cases of RALF and LARS, the authors found no difference in postoperative pain scores, hospital stay, complication rates, reoperation rates, and self-rated quality of life improvement (55). A longer postoperative period was analyzed by Nakadi et al. who found that, after the 1st, 6th, and 12th postoperative months, patients reported a similar amount of postoperative complaints. Both groups had similar lengths of postoperative hospital stays (53). Yet another randomized control trial by Muller-Stich et al. found no difference in conversion to an open approach, mean length of hospital stay, or symptomatic outcomes at 30 days postoperatively between the two procedures (56). In a large analysis of anti-reflux procedures specifically examining readmission rates and patient outcomes, Owen et al. found a lower 30-day readmission rate after LARS, but no significant difference in mortality, morbidity, and length of stay between LARS and RALF (57).

RALF has also been shown to significantly decrease postoperative esophageal acid exposure time (EAET). Frazzoni et al. retrospectively examined postoperative manometric and acid reflux parameters between RALF and LARS and found that while there were no significant differences in manometric parameters, the median EAET was significantly lower for RALF than LARS. Additionally, they also found that abnormal EAET values were found in 0% (0/44) of patients who underwent RALF vs. 14% (6/44) of patients who underwent LARS. Furthermore, normal EAET was observed significantly less frequently after LARS as compared to RALF (86% vs. 100%) (58). This small, but significant, improvement can be clinically relevant in patients with challenging PPI-refractory reflux disease where any demonstrated therapeutic gain can be meaningful (58).

**Robotic surgery vs. laparoscopic surgery—operative time and cost**

While robotic platforms have slowly been introduced as an acceptable and safe option in anti-reflux surgery, operative times and the cost of the robotic platform are often cited as potential drawbacks. Most randomized clinical trials have reported a significant increase in operative times with RALF, especially in the earlier phases of adoption (51-53,59). Morino et al. examined 50 consecutive patients undergoing LARS or RALF and demonstrated that RALF required significantly longer total operative times (131 minutes for RALF vs. 91 minutes for LARS) (52). Similarly, in a retrospective cohort study, Jensen et al. compared 103 patients who underwent RALF or LARS and found significantly longer operative durations for robotic surgery (135 minutes for RALF vs. 86 minutes for LARS). While early experience with robotic surgery was associated with longer operating times, patients who are operated on by a single surgeon and a highly experienced operating room team have greatly decreased operative times (59). In a randomized prospective cohort trial, Muller-Stich et al. showed a shorter total operative time for RALF compared to conventional LARS (88 minutes for RALF vs. 103 minutes for LARS) and the authors attributed the results to the experience of the one operating surgeon and the experienced robotics team (56).

It is no surprise that there is often a higher upfront
of 33 publications on anti-reflux
et al. Figure 2 (57) compared 9,572 LARSs and 339
found that total operative
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et al. et al.

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many seek to reduce some of the most common side-effects
of LARS: inability to belch/vomit, increased postoperative
dysphagia, and increased flatulence. Nevertheless, at this
time, most endoscopic therapies have failed to consistently
deliver adequate acid suppression in the majority of patients
and more advanced alternatives would be needed before
replacing LARS as the gold standard (9).

Today, there are four EART currently in use: three of
which are executed with proprietary devices (Figure 2),
and the fourth utilizing a traditional endoscopic set-up
[anti-reflux Mucosectomy (ARMS)]. The three currently
available proprietary options include the transoral incisionless fundoplication (TIF) (EsophyX device; EndoGastric Solutions, Redmond, Wash, USA), Stretta (Mederi Therapeutics, Restech, Houston, TX, USA), and Medigus Ultrasonic Surgical Endostapler (MUSE) (Medigus Ltd, Omer, Israel) (62). Several previous devices have been
developed but have since been removed from the market
either because of safety concerns or lack of efficacy (62). As
the currently available devices are conceived now, EART
should not be considered an alternative to traditional
laparoscopic or robotic anti-reflux surgical therapy (62).
These methods cannot adequately address the problem of
a hiatal or paraesophageal hernia greater than 2 cm and
should not be used in patients with significant anatomic
abnormalities (9). These devices also have not been
systematically studied in patients with active esophagitis,
BE, and esophageal motility disorders and limited data
exists in regards to treating patients with laryngopharyngeal
reflux disease (62).

As with LARS and RALF, selecting the proper patient
for EART is important. Many disease processes can mimic
GERD and may not be relieved with traditional anti-reflux
procedures. If a patient’s pH/impedance study is negative,
the surgeon must consider other possible diagnoses,
such as achalasia, ineffective motility, spastic esophageal
dysmotility, gastroparesis, and eosinophilic esophagitis, and
should counsel patients away from anti-reflux intervention.
Data has shown that up to 44% of patients with esophageal
dysmotility/achalasia, 57% of patients with eosinophilic
esophagitis, and 73% of patients with gastroparesis may
experience pathologic reflux symptoms that mimic reflux
experienced by patients with GERD (63). As mentioned
previously, patients with a hiatal hernia greater than
2 cm are not good candidates for endoscopic therapy
and require a traditional surgical procedure. Lastly, in
patients with morbid obesity, endoscopic therapy may not

Endoscopic therapy for the treatment of reflux disease

Introduction

The technological advancement of minimally invasive surgery is often driven by the desire to complete an operation with as few incisions as possible. Endoscopic anti-reflux therapy (EART) realizes this vision to become the holy grail of minimally invasive surgery: incisionless surgery through a natural orifice. Endoscopic anti-reflux therapy was developed in an attempt to further increase the number of patients with GERD who can be treated with surgical therapy. Lifestyle modifications and medical therapy remain the first line treatments, but around 20–40% of patients will experience only partial relief in their GERD symptoms. While LARS is the gold-standard for the surgical treatment of GERD, only a small percentage of patients with severe GERD opt for traditional surgical intervention. Minimally invasive endoscopic therapies have been developed to address many of the challenges faced with laparoscopy and robotics and to offer an alternative to medical therapy and major surgical intervention (62). While the currently available products all function differently, the main goal of the therapy is to endoscopically reduce lower esophageal sphincter compliance. They offer the potential for decreased

The technological advancement of minimally invasive surgery is often driven by the desire to complete an operation with as few incisions as possible. Endoscopic anti-reflux therapy (EART) realizes this vision to become the holy grail of minimally invasive surgery: incisionless surgery through a natural orifice. Endoscopic anti-reflux therapy was developed in an attempt to further increase the number of patients with GERD who can be treated with surgical therapy. Lifestyle modifications and medical therapy remain the first line treatments, but around 20–40% of patients will experience only partial relief in their GERD symptoms. While LARS is the gold-standard for the surgical treatment of GERD, only a small percentage of patients with severe GERD opt for traditional surgical intervention. Minimally invasive endoscopic therapies have been developed to address many of the challenges faced with laparoscopy and robotics and to offer an alternative to medical therapy and major surgical intervention (62). While the currently available products all function differently, the main goal of the therapy is to endoscopically reduce lower esophageal sphincter compliance. They offer the potential for decreased postoperative pain and shorter hospital stay. Additionally, many seek to reduce some of the most common side-effects of LARS: inability to belch/vomit, increased postoperative dysphagia, and increased flatulence. Nevertheless, at this time, most endoscopic therapies have failed to consistently deliver adequate acid suppression in the majority of patients and more advanced alternatives would be needed before replacing LARS as the gold standard (9).

Today, there are four EART currently in use: three of which are executed with proprietary devices (Figure 2), and the fourth utilizing a traditional endoscopic set-up [anti-reflux Mucosectomy (ARMS)]. The three currently available proprietary options include the transoral incisionless fundoplication (TIF) (EsophyX device; EndoGastric Solutions, Redmond, Wash, USA), Stretta (Mederi Therapeutics, Restech, Houston, TX, USA), and Medigus Ultrasonic Surgical Endostapler (MUSE) (Medigus Ltd, Omer, Israel) (62). Several previous devices have been developed but have since been removed from the market either because of safety concerns or lack of efficacy (62). As the currently available devices are conceived now, EART should not be considered an alternative to traditional laparoscopic or robotic anti-reflux surgical therapy (62). These methods cannot adequately address the problem of a hiatal or paraesophageal hernia greater than 2 cm and should not be used in patients with significant anatomic abnormalities (9). These devices also have not been systematically studied in patients with active esophagitis, BE, and esophageal motility disorders and limited data exists in regards to treating patients with laryngopharyngeal reflux disease (62).

As with LARS and RALF, selecting the proper patient for EART is important. Many disease processes can mimic GERD and may not be relieved with traditional anti-reflux procedures. If a patient’s pH/impedance study is negative, the surgeon must consider other possible diagnoses, such as achalasia, ineffective motility, spastic esophageal dysmotility, gastroparesis, and eosinophilic esophagitis, and should counsel patients away from anti-reflux intervention.

Data has shown that up to 44% of patients with esophageal dysmotility/achalasia, 57% of patients with eosinophilic esophagitis, and 73% of patients with gastroparesis may experience pathologic reflux symptoms that mimic reflux experienced by patients with GERD (63). As mentioned previously, patients with a hiatal hernia greater than 2 cm are not good candidates for endoscopic therapy and require a traditional surgical procedure. Lastly, in patients with morbid obesity, endoscopic therapy may not
be the ideal option; the gold standard remains bariatric surgery, with Roux-en-Y gastric bypass favored amongst the many options. Of note, patients with a previous sleeve gastrectomy and new or recurrent reflux symptoms not responding to medical therapy may respond to endoscopic radiofrequency ablation with the Stretta (64). This could potentially be offered prior to the traditional therapy of conversion to Roux-en-Y gastric bypass.

**Transoral incisionless fundoplication**

Initially approved in September 2007 by the FDA, the TIF using the EsophyX device endoscopically reconstructs the LES and restores the angle of HIS. This procedure requires general anesthesia and offers the ability to reduce small hiatal hernias (<2 cm). It can create a 2–4 cm long valve with a >270° fundoplication (62). Two endoscopists are required for the procedure: one to operate the gastroscope and the second to manipulate the EsophyX device. The device is loaded over the shaft of a compatible gastroscope and both are advanced into the stomach. With the gastroscope retroflexed to view the gastric cardia, the EsophyX device creates the fundoplication. At the end of the procedure, 20 fasteners (placed 1–3 cm above the GEJ) create the fusion of the esophagus and fundus (62). A second generation of the device, EsophyX 2, is now available and can be used to perform a slightly modified TIF 2.0 procedure.

Four randomized clinical trials have evaluated TIF 2.0 procedure with the EsophyX 2 device for the treatment of GERD. The RESPECT (Randomized EsophyX 2 vs. Sham, Placebo-Controlled Transoral Fundoplication) study was a multicenter, blinded, randomized control trial that compared the TIF 2.0 procedure plus placebo medication vs. a sham operation and optimal PPI therapy for patients with >6 months of GERD and regurgitation symptoms despite PPI treatment. By an intention-to-treat analysis, TIF 2.0 eliminated troublesome regurgitation in 67% of patients vs. only 47% of patients with PPI therapy alone. Esophageal pH improved but did not normalize after TIF 2.0 and both groups had similar improvements in GERD symptom scores (65). A large systematic review and network meta-analysis comprising 1,128 patients by Richter et al.

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**Figure 2** This figure depicts the three proprietary endoscopic devices used for the treatment of GERD. (A) Transoral incisionless fundoplication (TIF® 2.0) procedure to rebuild the GEJ (EsophyX® Z+ device; EndoGastric Solutions, Redmond, Wash, USA) (B) Medigus Ultrasonic Surgical Endostapler (MUSE) (Medigus Ltd, Omer, Israel) (C) Stretta catheter with four electrode needle sheaths with resultant tightening of the GEJ (Mederi Therapeutics, Restech, Houston, TX, USA).
compared the efficacy of TIF, LARS, PPI therapy and a sham procedure. The authors found that while TIF had the highest probability of increasing patient’s health related quality of life (followed by LARS, sham procedure, and PPI), LARS was the best at controlling esophageal pH (followed by PPI, TIF, and sham) (66). Interestingly, other than the sham procedure, the TIF was associated with the greatest percentage of patients with persistent esophagitis (66). Ultimately, the authors concluded that TIF was not an adequate long-term alternative to LARS or PPI therapy (66).

**Stretta**

Originally cleared by the FDA in 2000, updated and cleared again in 2011, Stretta acts by delivering radiofrequency (RF) current to ablate the muscles of the LES (62). While the exact mechanism remains unclear, RF reduces the number of transient lower esophageal sphincter relaxations and decreases LES compliance (62). Stretta RF does not require general anesthesia and can be performed in a routine outpatient setting under sedation or monitored anesthesia care. Low power RF energy is delivered to the muscularis propria of the lower esophagus (5W per channel at 460 kHz frequency). Proper placement of the needle electrodes in the muscle is confirmed by impedance measurements (62). After performing an upper endoscopy to measure the distance from the bite block to the Z-line, a guidewire is introduced, the endoscope is retracted, and Stretta is advanced over the guidewire to about 1 cm proximal to the Z-line. Without endoscopic guidance, RF ablation is applied to a total of 56 treatment sites at 6 different treatment levels (4 antegrade and 2 retrograde in the proximal stomach) (62). Safety has been a major factor for the implementation of Stretta. Lipka et al. performed a Manufacturer and User Facility Device Experience search to assess the risks associated with the Stretta procedure. Since cleared by the FDA, multiple reported serious adverse events have occurred including pneumonia, gastroparesis, esophageal perforation, cardiac arrest and four reported deaths (67).

In total there have been four randomized clinical trials that have evaluated the Stretta device. The largest of which was a randomized control trial of 64 patients that compared Stretta (35 patients) to a sham procedure (29 patients) (68). At 6 months, there was no improvement in PPI usage or reduction in median esophageal acid exposure time. However, the group found a significant improvement in symptomatic relief (61% of Stretta patients vs. 33% in the sham group) and GERD-HRQL scores (68). A smaller randomized control trial of 22 patients randomized patients to a sham or Stretta procedure (69). Similarly to Corley et al., they found no difference in esophageal acid pH exposure or reduction in PPI use, but that Stretta significantly improved quality of life scores for bodily pain (69). Additionally, the authors examined the distensibility of the LES by using a barostat bag: a decrease in LES distensibility was noted after the Stretta procedure, but was found to be reversible on local administration of sildenafil (69). The authors concluded that the decreased compliance of the LES following the Stretta procedure is likely secondary to altered LES neuromuscular motility rather than LES fibrosis as was previously thought (69). A recent meta-analysis and systematic review by Fass et al. found more favorable results (70). In a review of 28 studies (4 randomized control trials) representing 2,468 patients, the authors found that Stretta was associated with improved health related quality of life scores and pooled heartburn standardized scores at a mean follow up of 25 months (70). Additionally, Stretta was associated with a significant reduction in the rate of erosive esophagitis by 24% (70). However, no change in LES basal pressure was found (70).

**Medigus ultrasonic surgical endostapler**

The Medigus Ultrasonic Surgical Endostapler (MUSE™), cleared for use by the FDA in January 2015, is an endoscopic stapling system that creates a partial fundoplication. The MUSE endoscope is advanced into the stomach through a previously placed overtube, retroflexed and pulled back to the correct level above the GEJ (usually around 3 cm) using a built-in ultrasonographic gap finder (62). The anvil engages with the rigid section of the endoscope shaft to clamp the fundus against the distal esophagus. A staple is delivered and the procedure is repeated to form a 180º fundoplication.

The clinical efficacy of the MUSE was assessed in a multicenter, prospective trial. Sixty-nine (69) patients underwent MUSE endoscopic stapling: GERD-HRQL scores improved by >50% off PPI therapy in 73% of patients and 65% of patients were no longer using daily PPIs at 6 months (71). Long-term follow-up of the data showed that at four years, 69% were off their daily PPIs and their GERD-HRQL score had significantly decreased from baseline (71). Another smaller, non-randomized study was performed to assess the product’s safety in which endoscopic stapling (11 patients) was compared with laparoscopic
fundoplication (16 patients) (72). The authors did not find a significant difference in GERD-HRQL scores: scores decreased by 87% in the LARS group vs. 64% in the endoscopic stapling subset. PPI use was found to be higher in the endoscopic stapling group, but the results were not significant (72).

Safety has been the main concern with the MUSE device: 8 serious adverse events, including pain, fever, viral infection and mediastinitis, were reported in the first 24 patients in a multicenter trial. In the same series, two severe adverse events were also reported: esophageal leak (resulting in pneumothorax, empyema and a 22 day hospital stay) and severe upper GI bleeding (requiring two units of blood) (72). These early adverse events led to changes in the procedure protocol (prophylactic antiemetic therapy and an additional stapling site) and the routine use of a post-procedure chest radiograph to exclude a postoperative esophageal leak (62,72). Limited adverse events have been reported following these changes.

**Anti-reflux mucosectomy**

ARMS is the newest endoscopic technique designed to treat reflux disease. It is unique amongst the endoscopic options as it does not require special or proprietary equipment. The procedure consists of a hemi-circumferential endoscopic mucosal resection of the gastric cardia around the GEJ (73). With the scope in a retroflexed position, the mucosa is marked with the snare 270° around the GE valve. The mucosa of the cardia is raised with a combination of saline, methylene blue and epinephrine; the tissue is banded; and subsequently transected with forced coagulation (73). This process is repeated, rotating the scope around the GEJ. As the mucosectomy bed heals and scars, the tissue contracts and tightens to augment the natural anti-reflux valve of GEJ (73).

Limited long-term data exists on the ARMS procedure. An early pilot study to assess ARMS’ efficacy to treat reflux disease followed 10 patients after undergoing the procedure. The authors found significantly decreased total DeMeester scores, mean regurgitation scores, and mean heartburn scores at 2 months (74). 24-hour esophageal pH monitoring revealed that the fraction of time at pH <4 improved from 29% to 3%, but was ultimately not statistically significant. In all 10 patients, PPI therapy was discontinued. However, in the cohort, two patients required repeat balloon dilation to control post-procedural esophageal stenosis (74). A second case series by Hedberg et al. detailed 19 patients who underwent ARMS. In this series, 68% of patients showed significant symptom improvement scores and significant rates of PPI discontinuation. However, three patients, post-procedure, had troubling dysphagia and required balloon dilation (73). Additionally, of the six patients who did not have symptom relief following ARMS, 3 (50%) ultimately underwent LARS. More data and future randomized studies are needed to accurately compare the efficacy and safety of ARMS to mainstay PPI therapy and LARS.

**Endoscopic anti-reflux therapy conclusion**

Endoscopic anti-reflux therapy offers a truly minimally invasive option for select patients. While randomized trials and case series have reported mixed efficacy with these techniques, EART potentially expands the treatment options available for troubling reflux disease. Although most trials have not shown consistent improvement in objective measurements, such as normalization of pH values and augmentation of LES pressures, many patients do report subjective clinical improvement following EART and many can ultimately wean from PPI therapy. The complexity of GERD and the many underlying causes of the disease dictates that a thorough, multidisciplinary diagnostic evaluation occur prior to decision making and that patient selection must be done carefully. Each of the aforementioned EART modalities have unique features and their selection needs to be tailored to the individual patient’s history and clinical presentation. As these and other minimally invasive techniques arrive on the market and long-term data becomes available, these devices should be limited to centers specializing in reflux disease (65,75). Future work should be geared to testing these devices against both laparoscopic and robotic anti-reflux procedures and should examine their long-term efficacy.

**Conclusions**

Today, patients and physicians have a multitude of anti-reflux therapy options for treating GERD. As with any interventional procedure, proper patient selection is key. Each of the three main modalities, LARS, RALF, and EART, come with their own set of advantages and disadvantages. Foregut surgeons and gastroenterologists should be familiar with each of these options to be able to provide their patients with customizable therapy to best treat their disease. Further work with large well-designed studies are needed to continue to evaluate the effectiveness
of each of these three therapies and determine which patients may benefit most from each specific procedure.

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Footnote

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