Review of ERCP Techniques in Roux-en-Y Gastric Bypass Patients: Highlight on the Novel EUS-Directed Transgastric ERCP (EGDE) Technique

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Abstract
Purpose of Review Hepatobiliary complications are common in Roux-en-Y gastric bypass (RYGB) patients. Despite development of multiple surgical and endoscopic access techniques over the years, ERCP using standard duodenoscope remains challenging in these patients due to the altered anatomy.

Recent Findings Limited success with enteroscope-assisted and laparoscope-assisted ERCP led to the evolution of the novel EUS-directed transgastric ERCP (EDGE) procedure, with variations of this technique termed as Gastric Access Temporary for Endoscopy (GATE), EUS-guided TransGastric ERCP (EUS-TG-ERCP), EUS-guided GastroGastrostomy-assisted ERCP (EUS-GG-ERCP), and EUS-directed transgastric intervention (EDGI). EDGE has high technical (100%) and clinical success rates (60–100%), lower adverse event rate (1.5–7.6%), and up to 20% access stent migration rate; without any significant weight changes. EDGE has significantly shorter procedure time (73 vs 184 min), post-procedural hospital stays (0.8 vs 2.65 days) and is more cost effective compared to other modalities.

Summary EDGE technique addresses the challenges of RYGB anatomy as a minimally invasive, clinically successful, fully endoscopic, and cost-effective option. We present a literature review of the EDGE technique from its inception to current, in addition to reviewing other access techniques, their advantages, disadvantages and outcomes.

Keywords Roux-en-Y gastric bypass (RYGB) • Endoscopic retrograde Cholangiopancreatography (ERCP) • Endoscopic ultrasound directed transGastric ERCP (EDGE) • Device assisted ERCP • Laparoscope-assisted ERCP (LA-ERCP) • Gastric access temporary for endoscopy (GATE)

Introduction
Over the recent years, obesity has emerged as a pandemic in the US and worldwide, contributing to about 400,000 deaths attributable to poor diet and physical inactivity [1]. Although diet and lifestyle modifications are the initial approaches to obesity treatment, their modest outcomes have led to an increased interest in bariatric surgery [2, 3]. Multiple bariatric procedures such as gastric banding, sleeve gastrectomy, and Roux-en-Y-Gastric Bypass (RYGB) have emerged, of which, RYGB has superseded other bariatric procedures by 70%–80% [4, 5].

About 29%–36% of post bariatric patients develop gallstones, and 13% develop gallbladder sludge within 6 months to 18 months after surgery [6, 7]. Traditional endoscopic retrograde cholangiopancreatography (ERCP) using a standard
duodenoscope to treat pancreaticobiliary disorders is challenging in post RYGB patients because of the altered anatomy (Fig. 1) and the technical difficulty of maneuvering the duodenoscope down the Roux limb to the jejuno-jejunal anastomosis and then up the biliopancreatic limb (short limb 80–100 cm or long limb 100–150 cm) to reach and gain access to the papilla [8, 9].

In this article, we discuss the various endoscopic and surgical techniques that have been developed over the years to address this challenge, with a focus on the efficacy, safety, and comparative outcomes of the newly developed Endoscopic ultrasound-directed transgastric ERCP (EDGE) procedure over other techniques.

Techniques

Colonoscope and Enteroscope-Assisted ERCP

In 1988, Gostout and Bender first reported the use of a pediatric colonoscope in 3 patients to reach the papilla in Roux-en-Y anatomy [10]. Later, Elton et al. from 1994 to 1997 (n = 18) described their experience using a pediatric colonoscope and enteroscope for diagnostic and therapeutic intervention in long limb bypass patients. In this study, if the procedure with pediatric colonoscope was not successful on first attempt then an enteroscope was used to reattempt the ERCP. With both the endoscopes, they reported an overall success rate of access to

![Fig. 1 Roux-en-Y Gastric Bypass Anatomy compared to normal ERCP anatomy (inset)](image-url)
papilla or bilio-enteric and pancreatico-enteric anastomoses as 84%, cannulation rate of 94%, and performance of ERCP in 86%. Although success rates were higher, the disadvantages included the lack of side viewing orientation and an elevator with the scope as well as smaller channel size that precluded the use of large diameter stents and accessories [11].

**Duodenoscope**

In 1997 Hintze et al. studied the efficacy of conventional duodenoscope and reported a success rate of only 33% in reaching the papilla in RYGB patients, and 67% in patients with Billroth II anastomoses [12].

**Combined Duodenoscope and Colonoscope**

Later in 2002, a prospective study reported a 67% ERCP success rate after multiple attempts with the use of a forward-viewing colonoscope and the duodenoscope in long limb RYGB patients with intact papilla. Even though success was achieved on repeat attempts, the number of failed initial attempts even in experienced hands highlighted the need to develop better techniques for this procedure [13].

**Advanced Endoscopy Techniques**

The development of advanced endoscopic techniques is broadly classified into two categories: device-assisted enteroscopy and alternative access point creation for use of duodenoscope.

**Device Assisted Enteroscopy (DAE)**

Enteroscopes were designed and widely used for the diagnosis and treatment of small bowel diseases. Recently, balloon tip overtube or rotational overtube-assisted procedures such as Double Balloon Enteroscopy (DBE), Single Balloon Enteroscopy (SBE) and Spiral Enteroscopy (SE) have been utilized to perform ERCP in RYGB patients [14].

**Double Balloon Enteroscopy and Single Balloon Enteroscopy**

DBE was first described in 2001 by Yamamoto et al. as a means of deep exploration of the small bowel [15]. Five years later it was used to perform ERCP in RYGB patients [16]. There is a long and short length DBE scope with lengths of...
200 cm and 155 cm respectively, and a working channel of 3.2 mm. The shorter length of the latter scope allows for the use of standard ERCP devices (Fig. 2). Subsequently, the feasibility of performing ERCP in RYGB patients using the single balloon tip overtube was first reported in 2008 [17]. The SBE length is similar to the long DBE scope at 200 cm but with a working channel of 2.8 mm (Fig. 3). In RYGB patients, a systematic review showed that DBE was able to reach the papilla or the anastomosis in 89%, cannulation was successful in 93% with a therapeutic success rate of 82%. Whereas with SBE, papilla or anastomosis was reached in 82%, cannulation was successful in 86% of cases with an overall therapeutic success rate of 68% [18]. Although DBE and SBE demonstrated higher success rate when compared to standard endoscopes, the success rates were more attributed to patients with short Roux limb with bilioenteric anastomosis and intact papilla (80%), compared to 58% with long Roux limb with intact papilla ($p = 0.040$) [19].

Spiral Enteroscopy

SE was introduced as an alternative to balloon assisted enteroscopy by Akerman and Cantero for the management of small bowel disorders [20]. Two studies so far have described the use of SE to perform ERCP in RYGB patients (Fig. 4). In both the studies, SE was able to reach the papilla in 76.2% to 86% of patients. Once the papilla was reached, cannulation and therapeutic intervention was successful in 92.3% to 100% of patients [21, 22].

DAE-ERCP Comparative Studies

Although the reported data with SE alone has shown higher efficacy rates, a large multicenter comparative study of all the three techniques such as DBE, SBE and SE in RYGB reported ERCP technical success rates of 74%, 69% and 72% and clinical success rates of 63%, 60% and 65%, respectively [23••].

The reasons for the limited success rates with forward viewing enteroscopes were the 1) inability to maneuver the endoscope to reach the native papilla due to the long length of the Roux limbs, internal hernias, and/or adhesions leading to sharp angulations; 2) forward viewing nature of the scope makes cannulation of the ampulla difficult due to the caudal approach; 3) lack of an elevator; 4) the long durations of the device assisted procedures; and 5) limited compatible
accessories to fit the length and diameter of the scope channel. Even though short overtubes were introduced to overcome some of these limitations, the small working channel remains a challenge for large diameter stent insertion and use of standard biliary accessories. Also, the success rate is dependent on the available expertise at select tertiary care centers and thus difficult to generalize for the community practices.

**Alternative Access**

To achieve higher efficacy and success rates, a second technique called alternative access ERCP, which includes Laparoscope-assisted ERCP (LA-ERCP), Percutaneous Assisted Trans prosthetic Endoscopic Therapy (PATENT), and EDGE procedure, was developed to provide the ability to use a standard duodenoscope and thereby the available standard ERCP accessories.

**Laparoscope-Assisted ERCP**

LA-ERCP was first described in 2002 [24]. This procedure entails a laparoscope-assisted surgical port placement into the excluded stomach, followed by percutaneous passage of the duodenoscope via the lap port into the duodenum. This facilitates the use of standard accessories via the side viewing duodenoscope (Fig. 5). A systematic review of 509 cases from 26 studies described the feasibility, safety and outcomes of LA-ERCP in patients with RYGB. The study reported 100% successful gastric access and 98.5% successful ductal cannulation [25]. A large multicenter evaluation of 579 patients reported a median procedure time for LA-ERCP to be 152 mins, with median length of hospital stay of 2 days [26]. In addition to the ERCP success rates, laparoscopic examination facilitates the diagnosis and treatment of adhesions and internal hernias which is a potential morbid complication seen with Roux-en-Y reconstruction [27].

**LA-ERCP Comparative Studies**

LA-ERCP, with more than 95% technical success rates, has surpassed the DAE techniques which has 60–70% technical success rates in the treatment of pancreaticobiliary diseases in RYGB patients [25]. Desai et al. showed that LA-ERCP has a higher success rate (100%) as compared to SE (57%) (p = 0.005) [28]. However, the complication rate was 11% higher with LA-
ERCP when compared to DAE, and 80% of these complications were related to the gastrostomy site [25, 29, 30].

Although LA-ERCP reported higher technical and clinical success rates when compared with DAE, its limitations include the need for higher technical expertise, more resource utilization due to operating room use as opposed to endoscopy suite, need for sterilizing the scope, and coordination of the surgical and endoscopist schedules [27, 31, 32]. The expertise of the surgeon working alongside the endoscopist is also very important. In patients with high BMI, multiple adhesions, prior surgeries, and the ability to access to the bypassed stomach can be technically challenging and time consuming. The endoscopist also must be experienced in navigating the duodenoscope through the trocar and positioning it in the duodenum through the bypassed scope.

To mitigate the complications associated with laparoscopic creation of a gastrostomy tract, some institutions have reported ERCP via gastrostomy tract created by interventional radiologists, but this can only be performed in a non-emergent setting [33]. However, the above studies highlighted the need of a complete endoscopic approach which overcomes the above disadvantages, leading to the evolution of another alternative access ERCP procedure such as PATENT and EDGE techniques.

**Percutaneous Assisted Trans Prosthetic Endoscopic Therapy (PATENT)**

**DAE-Guided PATENT**

PATENT technique was first described by Baron et al. in 2012 [34]. This technique was designed with an intent to develop a complete endoscopic approach by placing percutaneous gastrostomy (PEG) tube and subsequent performance of ERCP via the PEG in RYGB patients. This technique was demonstrated in 9 pigs and 1 human case in 2012 [35••]. The technical success rate of PEG and stent placement was 100%, but cholangiography was successful only in three animals. Stent migration and peristomal infection were the two adverse events reported in the animals.
Later, in 2013 a retrospective case series by the same group demonstrated the use of PATENT technique in 5 patients [36]. All patients underwent transoral DAE-assisted (DBE n = 4; SBE n = 1) gastrostomy creation in the excluded stomach with the use of three T-tags in a triangular configuration around the intended PEG site to secure apposition of the gastric and abdominal walls. After sequential dilation of the PEG site, a fully covered self-expanding esophageal metal stent (FCSEMS) was deployed within the gastrostomy tract. The SEMS was then maximally expanded and a standard duodenoscope was advanced through the percutaneous SEMS and the distal stomach to perform antegrade ERCP (Fig. 6). After the ERCP was completed, a 26-Fr balloon bumper PEG tube was placed at the gastrostomy site, which was subsequently removed no sooner than 4 weeks after the procedure to allow for tract maturation. The median procedure time of intubation of the enteroscope to PEG placement was reported as 97 min (IQR 76–186). Sphincterotomy induced perforation in one patient was the only reported complication in the study.

**EUS-Guided PATENT**

PATENT technique was further modified by Law et al. in 2015 when endoscopic ultrasound (EUS) guidance was used instead of DBE for the placement of PEG tube access [37]. The approach of using EUS for placement of PEG in RYGB patients was previously described in 2011 by Attam et al., wherein it was primarily done for feeding tube placement but they also used it to perform ERCP via the PEG site in one patient [38].

Although the PATENT technique is an endoscopic approach, it still involves creation of a gastrostomy site with significant complications of the PEG site access and longer procedure times. Thus, there was still a need for the development of a completely endoscopic and minimally invasive technique without the need for percutaneous access or gastrostomy creation.
EUS-Directed Transgastric ERCP (EDGE)

Initial Two-Stage EDGE Technique

EDGE procedure was first described by Kedia et al. in 2013 as a two-stage procedure. In the first stage, a 16 Fr gastrostomy tube was placed percutaneously into the excluded stomach using EUS guidance to identify and distend the excluded stomach via the gastric pouch. In the second stage, a FCSEMS was exchanged at the gastrostomy site, and an antegrade ERCP was performed using a side viewing duodenoscope that passed through the stent to reach the area of the papilla, very similar to the previously described EUS-guided PATENT technique. This study included 6 patients who underwent a two-stage EDGE procedure. Initial access was successful in 5 of 6 patients (83%). In one patient, first attempt was unsuccessful due to loss of wire access to the bypassed stomach. A mean wait time between the two stages was 5.8 days (± 2.2 days). Antegrade ERCP (second stage) was successfully performed in all 6 patients (100%). Though the procedure had higher success rate, it cannot be performed in emergent situations such as cholangitis because of the two-stage approach. In addition, PEG site infection was noted in 2 of 6 patients (33%), thus revealing the same limitations of the PATENT procedure [39].

Novel Single-Stage EDGE Technique

With the advent of the lumen apposing metal stent (LAMS), Kedia et al. in 2014 described a case of a single stage EDGE procedure by creating an EUS-directed gastro-gastric fistula using the LAMS in RYGB patient to perform antegrade ERCP. This was the first report of an entirely endoscopic internal EDGE procedure that could be performed by a single team in a minimally invasive fashion at a single session [40]. A follow up single-center case series by Kedia et al. on this internal EDGE technique showed successful EUS-directed jejuno gastric (n = 4) and EUS-directed jejuno gastric (n = 1) access in all 5 patients (technical success 100%) with the 15 mm diameter LAMS. ERCP was successfully performed during the index procedure in 3 of 5 (60%) patients, but in 2 of 5 (40%) patients ERCP was postponed due to difficulty in passing the duodenoscope through the LAMS on the initial procedure. No adverse events such as bleeding, perforation, peritonitis or pancreatitis were reported. Stent dislodgement was seen in 3 of 5 cases, 2 of which required a second LAMS and in 1 patient the LAMS was readjusted back into position. Removal of the LAMS and fistula closure with endoscopic suturing was confirmed in 2 of 5 patients, and LAMS was left in place in 3 patients for continued biliary access. No weight gain was reported in these patients on follow up. The mean time of procedure was 68 min [41•].

EDGE Safety & Efficacy

A multicenter study by Ngamruengphong et al. (n = 13) evaluated the safety and efficacy of EDGE but coined the name as EUS-guided transgastric (EUS-TG) ERCP. Technical success rate for placement of 15 mm LAMS and clinical success rate of ERCP through LAMS was 100%. The median wait time to perform ERCP after the LAMS placement was 11 days. Stent dislodgement was seen in 33% in whom therapeutic duodenoscope was used, but none with the slim duodenoscope. Similarly, an interim analysis by Tyberg et al. (n = 16) reported technical success of 100% and clinical success of 91%. Unlike the prior study they did not comment on the wait period between the LAMS placement and ERCP, but stent dislodgement was seen in 19% in whom a FCSEMS was used for replacement of LAMS. In both the studies, interventions such as over the scope clip (OTSC), endoscopic suturing, and argon plasma coagulation (APC) were used for closure of the fistulous tract, whereas some patients were left to heal by secondary intention. On follow up there was a mean weight change of −2.85 kg to −3.6 kg [42, 43].

Some studies have assessed the fistula status in addition to technical and clinical success rates of the procedure. A retrospective analysis of 19 patients from 2018 by James et al. aimed to assess the fistula closure rate after the LAMS removal and describe the associated signs and symptoms of persistent fistula and methods of closure. Technical success of LAMS placement was 100%. ERCP was performed during the index procedure in 4 patients while the remaining 15 patients had ERCP (n = 11) and EUS (n = 4) after a mean wait period of 48 days from LAMS placement. Interestingly this is the first study wherein they used LAMS to perform EUS guided diagnostic biopsy in RYGB patients. Similar to the study by Tyberg et al., this study also managed the stent dislodgement with FCSEMS. LAMS removal was performed at a mean of 182 days ± 158 days. APC was routinely performed on 12 patients to close the fistulous tract, except in 7 others who required repeat pancreaticobiliary access via the fistula tract. On mean follow up at 281 days, upper GI series was obtained in 11 patients to assess the fistula status. One of 11 (9%) had persistent fistula and gained about 5.6 kg, successfully closed with APC followed by OTSC placement at the jejuno gastric access site, leading to subsequent weight loss of 2.8 kg. Mean cohort weight change was +1.7 kg [44•].

A multicenter study presented by Runge et al. at DDW 2019 assessed the success, long-term complications and implications following the EDGE technique. Total of 166 patients were included in the study from 12 centers. Technical success was 98%, gastro gastric access was 52% and jejuno gastric in 48% of cases. LAMS was anchored in 21% (35/166) of patients (with suturing in 25, plastic double pigtail stents in 7, hemoclip in 2, and OTSC in 1). EDGE was performed in a single session in 51% and in two sessions in 49%
of cases. On mean follow up time of 47 days, LAMS was removed in all patients; and fistula closure was performed in 73% of patients; whereas 27% were left undisturbed following LAMS removal. Upper GI series was obtained in 51% (85/166) of patients, of which 10 patients (12%) had persistent gastrogastric fistula (GGF) and endoscopic closure was performed in 7 of 10 patients with a mean of 1.2 attempts. Intraprocedural and delayed complications were reported in 17% (28/166) of patients [45].

EDGE as GATE

A single center case series from 2019 by Wang et al. proposed a new management algorithm for EDGE cases, also coined as Gastric Access Temporary for Endoscopy (GATE). The technical success for LAMS placement as well as clinical success rate was 100% in 10 patients, 3 gastrogastric and 7 jejuno gastric, for 9 ERCP cases, and 1 case of EUS followed by endoscopic submucosal dissection (ESD) of a duodenal mass. In 7 of 9 patients ERCP was done during the index procedure and in the remaining 2 patients ERCP was performed in 2-3 weeks, after fistula tract maturation. In 3 of 7 patients in whom ERCP was done at the time of index procedure, LAMS was exchanged with a double pigtail plastic stent immediately after the procedure as their gastric remnant access site was transgastric. In the remaining 4 patients, 3 had stent exchanged later as their access site was transjejunal and 1 had LAMS left in place with an intent of an additional follow up ERCP. Three patients were lost to follow up, and of the remaining 7 patients, all had LAMS exchanged for double pigtail plastic stents. Of these 7, 5 cases had the plastic stent spontaneously expelled and the tract had closed, and the remaining 2 had plastic stent removed manually. All 7 of 7 cases (100%) had confirmed access tract closure. Two patients (20%) had adverse events such as bleeding and stent dislodgment both of whom were transjejunal access [46*].

EDGE as EDGI

Prior studies demonstrated the use of EDGE in performing ERCP in RYGB anatomy, but a multicenter study by Kraft et al. from 2019 coined a new term called EUS-directed transgastric intervention (EDGI), discussing the use of this technique in evaluation of various luminal and extraluminal conditions such as pancreatic mass, inflammatory pancreatic fluid collection, suspected cholangiocarcinoma, idiopathic recurrent pancreatitis, common bile duct dilation, abnormal liver function tests, duodenal mass, duodenal stricture, duodenal ulcer perforation, abnormal gastric imaging on CT scan, etc. Extraluminal interventions included EUS-guided drainage pancreatic fluid collection, EUS-guided fine needle aspiration of suspected cholangiocarcinoma, EUS-guided liver biopsy and EUS FNA of pancreatic cystic neoplasm. Luminal interventions included gastro-duodenal luminal biopsies and closure of perforated duodenal ulcer [47].

Edge Comparative Studies

EDGE Vs Enteroscopy-Assisted ERCP

A multicenter study by Bukhari et al. published in 2018 compared the outcomes and adverse events between EUS-guided gastrogastrostomy-assisted ERCP (EUS-GG-ERCP) and enteroscopy-assisted ERCP (e-ERCP) in RYGB patients. Out of 60 patients, 30 underwent EUS-GG-ERCP and remaining 30 underwent e-ERCP (DBE in 19 and SBE in 11). Technical success was higher with EUS-GG-ERCP when compared to e-ERCP (100% vs 60%, p < .001). Total procedure time and median length of hospitalization was significantly shorter with EUS-GG ERCP group (49.9 min vs 90.7 min, p < .001; and 1 vs 10.5 days, p = .02). However, adverse event rate was similar in both the groups. (6.7% vs 10.0%, p = 1). No weight change was reported after EUS-GG-ERCP at mean follow up of 209 days [48*].

EDGE Vs LA-ERCP

A multicenter retrospective study published in 2018 by Kedia et al. compared the outcomes between EDGE and LA-ERCP. A total of 72 patients were included in the study (29 in EDGE group and 43 in LA-ERCP). Technical (96.5% vs 100%, p = 0.40) and clinical (96.5% vs 97.7%, p = 1.0) success rates were similar in the EDGE and LA-ERCP groups. In LA-ERCP, 21 patients had gastrosomy tube closure during the same session, whereas in 22 it was closed later. There was no significant difference in the adverse event rates between the groups (24% vs 19%, p = 0.57). EDGE had significantly shorter procedure time and length of stay compared to LA-ERCP (73 min vs 184 min p < 0.0001; and 0.8 d vs 2.65 d p < 0.00008). The overall weight change after EDGE at mean follow up of 28 weeks was - 6.6 lbs. [49].

A meta-analysis presented by Khan et al. at DDW 2018 comparing LA-ERCP to EDGE included 22 observational studies (18 LA-ERCP and 4 EDGE) with 941 patients (843 LAERCP and 98 EDGE). Technical and clinical success rates were similar in both the groups (98% vs 96% p = 0.07 and 96% vs 96% p = 0.84) without any significant difference in the adverse event rate (13% vs 10%, p = 0.32). However, pooled mean length of stay and procedure time were shorter with EDGE (1.1 vs 3.1 days and 43 min vs 166 min) [50].
The Geisinger Experience

We presented our own experience from the Geisinger Medical Center comparing outcomes of EDGE vs LA-ERCP at The American College of Gastroenterology’s Annual Scientific Meeting, held in October 2019. A total of 76 RYGB patients who underwent ERCP (59 LA-ERCP and 17 EDGE) were analyzed. All cases of LA-ERCP and EDGE were performed in a single step setting. Technical and clinical success rates were 100% in both the groups. Adverse event rate and length of hospital stay were also similar (17% vs 6%, 2.7 vs 2.6 days, \( p = 0.94 \)), however EDGE had significantly shorter procedure time when compared to LA-ERCP (103 min vs 208 min, \( p < 0.001 \)). The median time for lumen-apposing metal stent removal was 22 days (range 0–111). There was no significant weight gain (−6.33 lbs.) at median follow up of 35 days in the EDGE group [51*].

Based on our above experience, we now prefer to do LA-ERCP only when concomitant cholecystectomy needs to be performed. If RYGB patients are already post cholecystectomy and need an ERCP or access to excluded GI tract, the EDGE procedure is preferred. We perform all our EDGE cases as a single session procedure. Using a therapeutic linear echoendoscope, the excluded stomach is identified under endosonographic guidance looking for the “sand dollar sign” [52], preferentially as a gastro-gastric view when technically feasible, making sure that the distance between the two lumen is less than 10 mm. The excluded stomach is then punctured and injected with contrast under fluoroscopic guidance using a 19 g EUS-FNA needle. The excluded stomach is then distended using 250–400 ml of water mixed with indigo carmine solution via the EUS-FNA needle to create a safe target for LAMS placement. The EUS needle is then exchanged of, and the now available wider 20 mm electrocautery enhanced LAMS is placed freehand under EUS and fluoroscopic guidance to create a gastro-gastric access tract, without anchoring the stent. The LAMS lumen is then dilated using a through-the-scope balloon dilator to 20 mm after confirming reflux of the blue stained water. A diagnostic duodenoscope is then passed via the newly created gastro-gastrostomy for ampullary access or endoscopic intervention (Fig. 7). We have noticed almost no risk of stent migration with this technique. On occasion, we have created a jejuno-gastric access site using the 20 mm LAMS (Fig. 8);
when a gastro-gastric access was not technically feasible either due to very small pouch size or lack of a safe gastro-gastric access window for the LAMS deployment. A follow up procedure is performed for LAMS removal usually within 2–3 weeks and the LAMS access site is actively closed using Endosuture with which no persistent fistula cases noted at our center. This more proactive approach for closure is partly influenced by our patients traveling long distances for their care and thereby we hope to reduce the need for reintervention or loss of follow up.

**Cost-Effectiveness**

In this day and age of healthcare economics, decreasing the cost and length of stay are very important factors to be kept in mind. In a cost-analysis model comparing laparoscopic-assisted, enteroscopy-assisted, and EDGE-assisted ERCP approaches, EDGE was found to be more cost effective when compared to DAE-ERCP and LA-ERCP ($1431 vs $3147 and $9312) [53]. This was thought to be driven by the lack of need for the operating room and surgical supplies, and the associated costs. EDGE was also found to have the high total quality-adjusted life-years (QALY).

**Conclusions**

In patients with altered Roux-en-Y anatomy, traditional ERCP using a duodenoscope is technically challenging and clinically ineffective for ampullary access. Over the decades multiple trials and techniques have been devised to reach the ampulla to perform therapeutic intervention for various pancreaticobiliary diseases. The combined techniques of using pediatric colonoscope and duodenoscope to even reach the ampulla have shown only 33%–67% success rate despite multiple attempts [12, 13]. With the development of DAE for small bowel evaluation and modifying its use to achieve ampullary access in RYGB anatomy, the biliary access rates improved to only 60%–65% [23]. In 2002 when the LA-ERCP technique was developed, it surpassed the other advanced endoscopic techniques with higher technical and clinical success rates [25–27]. However, limitations remained as it is resource intensive, requiring collaboration of endoscopy and operating room schedules with the need for surgical access, and has higher complication rates and cost [30, 53]. To overcome these surgical limitations and to achieve an all-endoscopic alternative access, techniques such as PATENT were developed wherein gastrostomy tract was created endoscopically with the help of an enteroscope or EUS, without the need of...
surgical access. However, this still needed a percutaneous access which had its own set of complications from PEG site infections and need for PEG tract maturation prior to closure, delaying recovery and affecting quality of life [34–38].

In 2014, with the advent of LAMS, Kedia et al. first described an all-endoscopic “Internal EUS-Directed Transgastric ERCP (EDGE)” procedure and aptly termed it “Game Over”. This procedure has been called different names in different publications, such as EDGE, GATE, EDGI, EUS-TG-ERC, and EUS-GG-ERC. In essence, the procedure entails EUS-guided transgastric or transjejunal access into the remnant stomach followed by placement of a LAMS to facilitate the use of a standard duodenoscope for ERCP or other luminal or extraluminal endoscopic interventions.

This procedure has been a game changer in RYGB patients as it eliminates the need for surgical or percutaneous access, gives the option of repeat intervention for stent removal or exchange, and eliminates the need to keep a PEG tube in place for tract maturation, thereby reducing complication rates. All EDGE-related studies have demonstrated higher technical (100%) and clinical success rates (60%–100%) [41–47, 54]. EDGE has similar success rates (96.5% vs 100% p = 0.40) as compared to LA-ERC in achieving therapeutic ERCP, but appears superior when compared to DAE (100% vs 60%, p < 0.001) [48–50]. Although LA-ERC and EDGE have similar success rates, EDGE has shorter procedure time (73 vs 184 min, p < 0.00001) and post procedural hospital stay (0.8 vs 2.65 d, p < 0.00008) [49, 50].

In all the initial studies assessing the efficacy and safety of EDGE technique, a 15 mm diameter LAMS was used and a common problem encountered during the procedure was stent dislodgement (15.4%–20%) [42, 43]. In addition to stent dislodgement, other adverse events such as bleeding (7.6%), LAMS malposition (4.5%), migration (4.5%), perforation (1.5%) and pancreatitis (1.5%) were also reported [55]. Some studies have noticed that the gastrogastric access had a higher complication rate than jejuno gastric, but stent dislodgment was higher with the jejunogastric access.

To decrease the risk of stent dislodgment, EDGE was performed in two-sessions by the endoscopists with a mean wait time of 11–48 days to allow fistula tract maturation between LAMS placement and ERCP [42, 44]. To date, there is one prior report where the procedure was delayed only 72 h rather than a wait of 11–48 days when larger size (20 mm) LAMS was used [56]. Another study at DDW 2019 demonstrated the use of 20 mm LAMS. A total of nine patients were included in the study; technical and clinical success was 100%. The mean wait time was 2 days between the stent placement and the ERCP without any reported stent dislodgment [54]. In addition to the size of LAMS, no stent dislodgment was noticed when slim duodenoscope was used instead of therapeutic duodenoscope [42]. This evidence was supported with the research study at our own institution where stent dislodgement occurred only while using a therapeutic duodenoscope in a single case, and we have been able to perform all our EDGE cases as a same session ERCP without any wait time after LAMS deployment with the combination of gastrogastric access with 20 mm LAMS and use of the slim duodenoscope, making it a true immediate access single session fully endoscopic procedure [51•].

The outcomes of LAMS site fistula closure are also of concern as it may be associated with the risk of weight regain in these high-risk patients. No significant weight changes were reported from the time of LAMS insertions to removal and fistula closure while LAMS was left in place after ERCP for an average of 20–82 days [41–46]. Many modalities like APC, OTSC, endoscopic suturing, and double pigtail plastic stent have been described to close the EDGE access tract after ERCP. Double pigtail plastic stent was considered more advantageous compared to other techniques in terms of its cost, endoscopic approach and minimal technical support in one recent study [46•] (Table 1).

Our review has shown strong data in support of EDGE as an all-endoscopic, efficacious, safer and superior alternative in terms of cost and time and that can be performed as a single-session procedure using minimal resources. However, LA-ERC can be considered in patients who need simultaneous cholecystectomy. In order for EDGE to evolve into a more effective standardized procedure across the board, prospective randomized studies are needed to compare the size of LAMS used, type of duodenoscope used, one vs two session procedure to allow for tract maturity, need for anchoring the stent with endosuture or OTSC, assessing its value for ERCP vs other endoscopic interventions in the bypassed GI tract, and comparing the modalities of fistula closure vs spontaneous closure with respective response. Dedicated procedure billing codes are also needed to better code and bill for this procedure, taking into consideration all the morbidity benefits, patient convenience, and cost savings as compared to surgical alternatives.

A systematic approach is necessary in managing these patients with pancreaticobiliary disease with underlying RYGB anatomy, with close collaborations between GI, radiology, interventional radiology and surgery. A multidisciplinary approach is key in deciding the most optimal method in managing these patients based on available expertise, resources, surgical and radiological back up, experience of the endoscopist and the staff in handling these patients and the associated complications, as well as patient’s comorbidities and preference. Further prospective studies will help guide, standardize practices and management approaches for this population.
Table 1  Summary of all EDGE papers reported to date

| Study Publication        | Type of LAMS: Cautery enhanced (CE), Non cautery enhanced (NCE) | Size of LAMS | Type of Fistula: Gastrogastric (GG), Jejunogastric (JG) | Technical success | Clinical success | No. of sessions | LAMS removal median/mean time | Type of closure: Endoscopic suturing (ES), Over the scope clip (OTSC), Argon plasma coagulation (APC), Through the scope clip (TTS) | Persistent fistula | Adverse events | Weight changes |
|--------------------------|---------------------------------------------------------------|--------------|---------------------------------------------------------|-------------------|------------------|-----------------|-------------------------------|---------------------------------------------------------------------------------------------------------------------------------|------------------|-----------------|-----------------|
| Kedia et al. 2015 (n=5)  | NR                                                             | 15 mm        | GG=4 JG=1                                              | 100%              | 60%              | one session=3; two session=2 | 3 weeks                      | ES=2 none stent dislodgement (60%), jejunal perforation in one patient (19%) | negative 2.85 kg | none            |                 |
| Tyberg et al. 2016 (n=16)| NR                                                             | 15 mm        | GG=6 JG=10                                             | 100%              | 91%              | one session=4; two session=6  | NR                           | ES=7 OTSC=2 secondary intention =1 1 out of 8                |                  |                 | negative 3.6±4.8 kg |
| Namomungpong et al. 2017(n=13) | CE=5 NCE=8                                                       | 15 mm        | GG=6 JG=7                                              | 100%              | 100%             | one session=2; two session=11 | 20 days                      | ES=6 OTSC=3 APC=3 1 out of 12 stent dislodgement (33%) |                   |                 |                 |
| James et al. 2018 (n=19) | CE=14 NCE=5                                                     | 15 mm        | GG=8 JG=11                                             | 100%              | 100%             | one session=4; two session=15 | 182 days                     | APC=12 None=5 1 out of 11 stent malposition (619)                      |                   |                 | weight gain of 1.7 kg |
| Bukhari et al. 2018 (n=30)| CE=14 NCE=16                                                    | 15 mm        | GG=17 JG=13                                            | 100%              | 100%             | one session=8; two session=22 | 26 days                      | ES=8 OTSC=7 APC=15 LAMS migration (67%), bleeding (33%), LAMS malposition (4.5%), LAMS migration (4.5%), perforation (15%), pancreatitis (15%) |                   |                 | neg 1.1±6.1 kg |
| Chiang et al. 2018 (n=66)| CE=33                                                           | NR           | GG=30 JG=34                                             | 92.40%            | NR               | one session=43                | NR                           | NR NR NR bleeding (7.6%), LAMS malposition (4.5%), LAMS migration (4.5%), perforation (1), pancreatitis (2) stent dislodgement (3) bleeding (1) stent dislodgement (20%), bleeding in one patient (none) |                   |                 |                 |
| Kedia et al. 2018 (n=29) | NR                                                             | 15 mm        | GG=28                                                   | 96.50%            | 96.50%           | NR                           | NR                           | negative 6.6 lbs |                 |                 |
| Wang et al. 2019 (n=10)  | NR                                                             | 15 mm        | GG=3 JG=7                                              | 100%              | 100%             | one session=7; two session=2  | 14–217 days                   | exchanged with Plastic double pigtail stent None | NR |                 |                 |
| Hsueh et al. 2019 (n=9)  | NR                                                             | 20 mm        | GG=5 JG=4                                              | 100%              | 100%             | one session=2; two session=7  | NR                           | NR NR NR |                  |                 |
| Rung et al. 2019 (n=166) | NR                                                             | NR           | GG=86 JG=80                                            | 98%               | NR               | one session=85; two session=81 | 47 days                      | ES=51 APC=46 OTSC=15 TTS=6 None=44 perforation (42%), stent migration (06%), bleeding, pneumoperitoneum (13%), post ERCP pancreatitis (2%), sepsis (0.6%), GGF |                 |                 |                 |
Abbreviations  RYGB, Roux-en-Y Gastric Bypass; ERCP, Endoscopic Retrograde Cholangiopancreatography; EDGE, Endoscopic ultrasound Directed transGastric ERCP; DAE, Device Assisted Enteroscopy; DBE, Double Balloon Enteroscopy; SBE, Single Balloon Enteroscopy; SE, Spiral Enteroscopy; LA-ERCP, Laparoscope-Assisted ERCP; PATENT, Percutaneous Assisted Transprosthetic Endoscopic Therapy; PEG, Percutaneous Endoscopic Gastrostomy; FCSEMS, Full Covered Self-Expandable Metal Stent; SEMS, Self-Expandable Metal Stent; EUS, Endoscopic Ultrasonography; LAMS, Lumen Apposing Metal Stent; EUS-TG-ERCP, EUS-guided TransGastroduodenal ERCP; OTSC, Over The Scope Clip; APC, Argon Plasma Coagulation; GATE, Gastric Access Temporary for Endoscopy; EDGI, EUS-Directed transGastric Intervention; ESD, Endoscopic Submucosal Dissection; DDW, Digestive Disease Week; GGF, Gastro-Gastric Fistula; EUS-GG-ERCP, EUS-guided GastroGastrostomy-assisted ERCP; e-ERCP, Enteroscopy-assisted ERCP; FNA, Fine needle aspiration

Declarations

Conflict of Interest  Harshit S. Khara, Swetha Parvataneni, Steven Park, Jihye Choi, Truptesh H. Kothari, and Shivangi T. Kothari declare that they have no conflict of interest.

Human and Animal Rights and Informed Consent  This article does not contain any studies with human or animal subjects performed by any of the authors.

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•• Of major importance

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