Endoscopic Ultrasound-Guided Hepaticogastrostomy: Technical Review and Tips to Prevent Adverse Events

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INTRODUCTION

Malignant biliary obstruction is usually treated under endoscopic retrograde cholangiopancreatography (ERCP) guidance. Most patients who need biliary drainage can be treated using ERCP guidance. However, ERCP can be challenging in patients with complications, such as malignant duodenal obstruction, or a surgically-altered anatomy, such as a Roux-en-Y anastomosis, which prevent advancement of the duodenoscope into the ampulla of Vater. Percutaneous transhepatic biliary drainage (PTBD) has been conventionally attempted as an alternative to bile duct drainage. Biliary drainage can also be achieved under single-balloon enteroscopy (SBE) or double-balloon enteroscopy (DBE) in patients with surgically-altered anatomy. However, although these procedures have clinical benefits, their disadvantages include self-tube removal or cosmetic issues after PTBD, and prolonged procedures and risk of perforation in SBE or DBE.

Endoscopic ultrasound-guided biliary drainage (EUS-BD) via transhepatic or transduodenal approaches has recently emerged as an alternative means of biliary drainage. Typically, EUS-guided gallbladder drainage (GBD) or choledochoduodenostomy (CDS) can be performed via both approaches. Additionally, EUS-guided hepaticogastrostomy (HGS) can also be performed. Among these approaches, EUS-HGS, because of its transgastric approach, can be performed in patients with malignant duodenal obstruction. Initial EUS-HGS procedures were technically challenging and associated with critical adverse events such as stent migration. However, along with device and technical developments, the technical tips for EUS-HGS have reached maturity due to device and technical developments. Although the technical success rates of EUS-HGS are high, the rate of adverse events is not low, with stent migration still being reported despite many preventive efforts. In this review, we described technical tips for EUS-HGS related to bile duct puncture, guidewire insertion, fistula dilation, and stent deployment, along with a literature review. Additionally, we provided technical tips to improve the technical success of EUS-HGS. (Gut Liver 2021;15:196-205)

Key Words: Endoscopic retrograde cholangiopancreatography; Endoscopic ultrasound-guided biliary drainage; Drainage; Endoscopic ultrasound
This review describes technical tips to improve the technical success of EUS-HGS, and provides a literature review of adverse event prevention.

**SUMMARY OF REPORTS RELATED TO EUS-HGS**

Various studies and meta-analyses on EUS-HGS have been published. Table 1 shows reports of EUS-HGS for malignant biliary obstruction that included over 10 cases. The reported technical and clinical success rates of EUS-HGS ranged from 65% to 100%, and from 76% to 100%, respectively. The most frequent adverse events were bleeding (n=12), followed by bile peritonitis (n=8), bile leak (n=7), and pneumoperitoneum (n=4). In addition, stent migration was observed in four patients. Patient death due to procedural related adverse events occurred in four patients.

The standard algorithm for performing EUS-HGS in patients with advanced malignant biliary obstruction at our hospital is presented in Fig. 1. Since duodenal obstruction is one of the risk factors for early biliary stent obstruction, EUS-HGS is considered as the first option in patients with duodenal obstruction. Of course, if ERCP fails, an EUS-guided rendezvous technique is attempted. EUS-HGS has several disadvantages. First, as shown in Table 1, adverse events are not infrequent. Second, stent migration is sometimes fatal, and although bile peritonitis is usually treated conservatively, this adverse event might be critical in patients with advanced malignant tumor. Therefore, endoscopists should carefully consider the indications for EUS-HGS. In patients with massive ascites, severe adverse events, such as stent migration, can occur. In addition, in patients with insufficient dilatation of the intrahepatic bile duct, bleeding can occur while puncturing the bile duct because blood vessels usually run near the intrahepatic bile duct.

**TECHNICAL TIPS FOR BILE DUCT PUNCTURING**

The first step in EUS-HGS is puncture of the intrahepatic bile duct using a 19-gauge fine needle. Two important points should be focused on here. One is the puncture site, and the other is avoidance of vessel puncture, which can be quite simply achieved using color Doppler ultrasound.

No evidence has been published about which of the bile duct sites B2, or B3, is more suitable for EUS-HGS. Generally, guidewire manipulation is easier if B2 is punctured rather than B3 because the bile duct runs relatively straight. However, if B2 is punctured, a puncture site at the intraluminal portion might be from the esophagus. Therefore, severe adverse events such as mediastinitis, can occur. If puncture site is not performed via the esophagus, the intraluminal puncture site is sometimes located in the stomach directly below the esophagus. In this situation, the proximal portion of the EUS-HGS stent might turn in the oral direction during deployment. Severe adverse events such as these have been associated with EUS-HGS stents. Therefore, we recommend puncture at the B3 site during EUS-HGS. With B3 puncture, the echoendoscope angle might be helpful in preventing puncture from the esophagus. Fig. 2 shows that the angle of the echoendoscope might be approximately 170° in the esophagus due to the limited width of the esophagus, but around 90° when it is in the stomach. Therefore, the shape of the scope should be checked by fluoroscopic imaging before intrahepatic bile duct puncture. The scope shape might also be important for guidewire manipulation. In addition, the bile duct, which runs from the upper left to the lower right corner on EUS images should be punctured to advance the guidewire towards the hepatic hilum.

**TECHNICAL TIPS FOR GUIDEWIRE MANIPULATION**

Guidewire manipulation is the most limiting step during EUS-HGS, particularly when attempted by non-experts. Guidewire shearing can occur under awkward guidewire manipulation. Vila et al. have described their initial experience with interventional EUS in a multicenter, retrospective, national survey of 125 patients in Spain who underwent interventional EUS. Among them, EUS-guided puncture was achieved in 120 patients, and contrast medium was injected into the ducts of 113 patients. However, unsuccessful manipulation of the guidewire was responsible for 68.2% of the technical failures. Therefore, they stated that intraductal manipulation of the guidewire seems to be the most challenging stage of the procedure. Guidewire manipulation can fail during insertion into the intrahepatic bile duct through the fine-needle aspiration (FNA) needle and while advancing the guidewire into the hepatic hilum after inserting it into the intrahepatic bile duct.

When the guidewire is inserted into the intrahepatic bile duct, it can sometimes penetrate the hepatic parenchyma (Fig. 3A). The guidewire should be gently manipulated to prevent this. Appropriate guidewire selection is also important. A flexible tip on the guidewire might be important, and the knuckle guidewire technique might be...
Table 1. Summary of Previous Studies (including 10 over Patients)

| Author (year) | No. of patients | Technical success rate | Clinical success rate | Dilation devices | Type of stent | Adverse events |
|---------------|-----------------|------------------------|----------------------|------------------|---------------|---------------|
| Bories et al. (2007) | 14 | 91 / 10 | 100 / 10 | Cystotome (6 or 8 F) | PS (7, 8.5, or 10 F), CSEMS (10 mm) | Ileus [1], biloma [1], stent migration [1], cholangitis [1] |
| Park et al. (2011) | 15 | 100 / 15 | 100 / 15 | ERCP catheter (4 F), dilator (6 and 7 F), needle knife | PS (7 F, 6-8 cm), FCSEMS (8-10 mm, 4-10 cm) | Pneumoperitoneum [4], bleeding [2] |
| Vila et al. (2012) | 16 | 65 / 22 | NA | NA | NA | Bleeding [3], biloma [3], perforation [2], liver hematoma [2], abscess [1] |
| Park et al. (2013) | 17 | 93 / 14 | 100 / 14 | ERCP catheter (4 F), dilator (6 and 7 F), needle knife | PS (7 F, 6-8 cm), FCSEMS (8-10 mm, 4-10 cm) | Biloma [1], intraperitoneal stent migration [1] |
| Kawakubo et al. (2014) | 18 | 95 / 20 | NA | Dilation catheter, balloon catheter, stent retriever, diathermic dilator | Plastic stent (8 F, Type IT) | Bile leak [2], stent misplacement [2], bleeding [1], cholangitis [1], biloma [1] |
| Paik et al. (2014) | 19 | 96 / 27 | 87 / 27 | ERCP catheter (4 F), dilator (6 and 7 F), needle knife | PCSEMS (8 mm, 10 cm) | Bacteremia [1], biloma [2], bleeding [3] |
| Artifon et al. (2015) | 20 | 96 / 25 | 89 / 25 | Needle-knife, dilating catheters | PCSEMS (8 mm, 10 cm) | Abdominal pain [3], bleeding [1] |
| Umeda et al. (2015) | 21 | 100 / 23 | 100 / 23 | Standard or tapered catheter, cautery dilator, dilation catheter (8 F), balloon (4 mm) | Plastic stent (8 F, Type IT) | None |
| Poincloux et al. (2015) | 22 | 98 / 66 | 94 / 66 | Needle-knife, dilator (6 or 7 F) | Plastic stent (10 F), PCSEMS (10 mm, 8-10 cm) | Bile leak [5], pneumoperitoneum [2], liver hematoma [1], severe sepsis and death [2] |
| Khashab et al. (2016) | 23 | 92 / 56 | 89 / 56 | Balloon, dilator, cautery dilator | Metal stent | None |
| Ogura et al. (2016) | 24 | 100 / 26 | 92 / 26 | ERCP catheter, balloon (4 mm) | PCSEMS (10 mm, 10, 12 cm) | None |
| Nakai et al. (2016) | 25 | 100 / 33 | 100 / 33 | Cautery dilator, bougie dilator (9 or 10 F) | PCSEMS | Bleeding [1], abscess [1], cholangitis [1] |
| Paik et al. (2017) | 26 | 100 / 20 | 100 / 20 | ERCP catheter, balloon (4 mm) | PCSEMS (10 mm, 10, 12 cm) | Intraprostatic stent migration [1], cholecystitis [1] |
| Minaga et al. (2017) | 27 | 97 / 29 | 76 / 29 | Dilator (6 or 7 F), balloon (4 mm) | Plastic stent, CSEMS | Bile peritonitis [1] |
| Ogura et al. (2017) | 28 | 100 / 10 | 100 / 10 | ERCP catheter, balloon (4 mm) | PCSEMS (10 mm, 8-10 cm) | None |
| Cho et al. (2017) | 29 | 100 / 21 | 86 / 21 | Needle-knife, balloon (4 mm) | Hybrid metal stent | Pneumoperitoneum [2], bleeding [1] |
| Sports et al. (2017) | 30 | 100 / 26 | 92 / 26 | Cystotome | PCSEMS | Severe sepsis [2], bile leak [2], bleeding and death [1] |
| Moroysset et al. (2017) | 31 | 94 / 17 | 76 / 17 | Cystotome | PCSEMS (10 mm) | Bleeding and death [1] |
| Oh et al. (2017) | 32 | 96 / 12 | 88 / 12 | Cannula (6 F), dilator (6 or 7 F), needle knife | Plastic stent (7-10 F, 6-10 cm), FCSEMS (6-10 mm, 6-10 cm) | Bacteremia [6], bleeding [5], bile peritonitis [4], pneumoperitoneum [4], intrahepatic stent migration [3] |
| Honjo et al. (2018) | 33 | 100 / 49 | 100 / 49 | Tapered mechanical dilator, cystotome, balloon (4 mm) | PCSEMS (6, 8 mm, 10, 12 cm), plastic stent (Type IT) | Abdominal pain [6], bleeding [5] |
| Okuno et al. (2018) | 34 | 100 / 20 | 95 / 20 | Dilator, ERCP catheter | PCSEMS (6 mm, 12, 15 cm) | Cholangitis [3] |
| Miyao et al. (2018) | 35 | 100 / 41 | 100 / 41 | ERCP catheter, balloon (4 mm) | PCSEMS (10 mm, 10, 12 cm) | Bile peritonitis [4], cholangitis [1], stent migration [1] |
| Paik et al. (2018) | 36 | 97 / 32 | 84 / 32 | None | PCSEMS (DEUS) | Cholangitis [1] |
| Minaga et al. (2019) | 37 | 87.5 / 20 | 100 / 20 | Bougie, balloon, cautery dilator | PCSEMS (8 mm, 10 cm) | Pancreatitis [1], bile peritonitis [1] |
| Ogura et al. (2019) | 38 | 96 / 29 | 89.7 / 29 | ERCP catheter, balloon (4 mm) | PCSEMS (8 mm, 10 cm) | Bile peritonitis [3] |

Data are presented as percent (number/number).
PS, plastic stent; CSEMS, covered self-expandable metal stent; ERCP, endoscopic retrograde cholangiopancreatography; FCSEMS, fully CSEMS; NA, not available; PCSEMS, partially CSEMS; ND, not discussed.
Therefore, guidewire selection should be decided based on flexibility of the tip of the guidewire. However, endoscopists should also consider stiffness, because several devices, such as dilatation devices and the stent delivery system need to be inserted after guidewire deployment.

If the guidewire is advanced into the hepatic parenchyma (Fig. 3B) or into the periphery of the bile duct (Fig. 3C), the guidewire should be manipulated towards the hepatic hilum again. Guidewire shearing can also occur during this procedure. Fig. 4 shows that while guidewire shearing can easily occur if the angle between the FNA needle and the guidewire is acute, it can even occur when it is obtuse. The echoendoscope angle might help to achieve successful guidewire insertion. If the scope angle is obtuse, the angle between the bile duct and FNA needle might be acute, and vice versa. Therefore, the intrahepatic bile duct should be punctured after checking the scope shape on fluoroscopic images. However, despite these techniques, the guidewire can still advance to the periphery of the bile duct. In this situation, the guidewire should be withdrawn a little and redirected towards the hepatic hilum. The liver impaction technique might be helpful during this step, as shown in Fig. 5. When the FNA needle is pulled into the hepatic parenchyma, the guidewire can be easily manipulated due to the following reasons: a tamponade effect around the tip of the FNA needle, and fact that the angle between the FNA needle and the guidewire might prevent guidewire shearing. If the FNA needle is pulled, the angle between the guidewire will increase.

A novel steerable access device for EUS-BD has recently become available. Ryou et al. reported their initial clinical experience with this device for EUS-BD. The device has a sharp stylet to allow for puncture. After stylet removal, the blunt-tipped access catheter (18.5 gauge) assumes a pre-determined curvature (90° or 135°) and is fully rotatable. The curvature is predetermined by fenestrations at the catheter tip. Guidewire sharing is theoretically avoided because of the blunt tip and the coaxial orientation of the wire relative to the catheter tip. Although, so far, only one patient has undergone EUS-HGS using this device, it shows promise in preventing guidewire shearing.

**TECHNICAL TIPS FOR FISTULA DILATION**

To fit the axis of the puncture route to insert another device might be technically important. Fig. 6 shows that accurate fitting of the axis allows clear visualization of all devices, and that the echoendoscope angle should be essentially the same as that before puncture and insertion of

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**Fig. 1.** Drainage algorithm used at our hospital. CBD, common bile duct; CDS, choledochoduodenostomy; ERCP, endoscopic retrograde cholangiopancreatography; E-ERCP, enteroscopy-guided ERCP; EUS, endoscopic ultrasound; HGS, hepaticogastrostomy; PTBD, percutaneous transhepatic biliary drainage.

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**Fig. 2.** [A] The angle of the echoendoscope in the esophagus is approximately 170° due to the limited width of the esophagus. [B] The angle of the echoendoscope is approximately 90° if it is in the stomach.

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**Fig. 3.** [A] The guidewire penetrates the hepatic parenchyma. [B] The guidewire is inserted into the branch bile duct. [C] The guidewire is advanced into the periphery of the bile duct.
the stent delivery system. If the device fits the axis, it would facilitate devices with a relatively large diameter to be easily inserted into the bile duct.

Various dilation techniques have been described to achieve this. Honjo et al.\(^3\) compared dilation with an ultra-tapered mechanical (n=26) or electrocautery (n=23) dilator (control) during EUS-HGS. The technical success rate of tract dilation did not differ significantly between the groups (electrocautery vs mechanical dilation: 23/23 [100%] vs 24/26 [92.3%], p=0.52), although the procedure duration was shorter in the electrocautery group (17.5±5 minutes vs 21.5±6.5 minutes, p=0.09). However, the adverse event of bleeding was significantly more frequent in the electrocautery group (5 vs 0, p=0.04). Therefore, they concluded that the ultra-tapered mechanical dilator designed for interventional EUS was safe and useful as it reduced post-procedural bleeding, together with having a high technical success rate compared with conventional electrocautery. We previously described the clinical benefits of dilation using a novel fine gauge balloon catheter for EUS-BD\(^4\) in a prospective, single-center, single-arm study of 20 patients, among whom nine underwent EUS-HGS. Technical success was defined as the insertion of a stent delivery system into the intrahepatic bile duct using...
a novel fine gauge balloon catheter without additional dilation. Consequently, all nine patients underwent EUS-HGS within a median duration of 14 minutes. Therefore, we concluded that our technique might facilitate EUS-BD. However, data from more patients and a randomized controlled comparison with other dilation techniques, such as graded or cystotome dilation, are needed.

On the other hand, electrocautery dilation might be perceived as risky, according to previous reports. Park et al. evaluated risk factors for adverse events associated with EUS-BD in 57 consecutive patients with malignant or benign biliary obstruction treated by EUS-BD after failed ERCP. That study significantly associated post-procedure adverse events with fistula dilation using a needle-knife compared with graded dilation (9/27 [33%] vs 2/28 [7%], p=0.02). Therefore, they recommended avoiding this practice in EUS-BD. The adverse event of bleeding might occur due to the "burn effect." Indeed, a comparison of electrocautery and mechanical dilation found no significant difference in terms of technical success rates and the number of additional dilations of the needle tract required to accomplish stent placement. In addition, procedure-related bleeding developed in six (18%) patients with electrocautery dilation, and in none during mechanical dilation (p=0.04). These results showed that the mechanical dilator allows safe tract dilation without the "burn effect" regardless of procedure. However, intrahepatic bile duct or stomach wall dilation can be difficult with non-electrocautery dilation devices in clinical practice, especially in patients with frequent cholangitis. Electrocautery dilation might be necessary in such patients. A novel fine gauge electrocautery dilator has recently become available in Japan. The distal end of the outer dilator has a metal tip that is only 3 F at the top, and the dilator is guided with a coaxial 0.025-inch guidewire. Therefore, the prevalence of the "burn effect" might be lower with this device than with conventional electrocautery dilation. In that study, EUS-BD, including EUS-GBD, CDS and HGS without dilation devices, were successful in all patients. In addition, severe adverse events such as bleeding did not occur. Although more patients and a prospective evaluation are essential to confirm these results, the fine gauge electrocautery dilator has the potential to improve the outcomes of EUS-BD.

Bile can leak from the intrahepatic bile duct after fistula dilation. Therefore, this step should be straightforward, and the procedural duration should be shortened or even omitted to decrease the likelihood of adverse events. According to this viewpoint, one-step stent deployment might be ideal. Park et al. evaluated the feasibility and safety of a novel dedicated device for one-step EUS-BD in a randomized trial of 32 patients with malignant biliary obstruction and failed ERCP. The patients were assigned to undergo the procedure in which a dedicated stent introducer was used with a modified hybrid metal stent (DH group, n=16) or a conventional 8.5-F biliary metal stent introducer was used with a fully covered metal stent (FC group, n=16). Technical success was defined as stent deployment without a dilation device in the DH group. One-step technical success was achieved in 14 (88%) of 16 patients, and the procedure was significantly shorter in the DH group (10 minutes vs 15 minutes, p=0.007). The rate of early adverse events was lower in the DH group (6.3% vs 31.3%, p=0.172), although the difference did not reach statistical significance. Hence, although this stent might confer benefits, since the one-step technical success rate was not 100%, further improvement of the one-step stent delivery system is needed.

STENT SELECTION AND TECHNICAL TIPS FOR STENT DEPLOYMENT

Critical adverse events, such as stent migration, can occur during this step. Previous reports indicate that almost all endoscopists use fully (FCSEMS) or partially (PCSEMS) covered self-expandable metal stents. The latter might have several advantages over the former. Since the site at the distal side is uncovered, branch bile duct obstruction might be prevented. If branch bile duct obstruction does occur as a complication, focal cholangitis can occur at the obstruction site. Uncovered sites of the PCSEMS might also play a role in anchoring the stent, which might prevent stent dislocation into the stomach. However, this needs to be confirmed in a prospective randomized controlled study. New metal stents dedicated to EUS-HGS have recently been developed. Park et al. investigated the feasibility of a new device for one-step EUS-BD, in which the stent has a dedicated introducer with a 3-F catheter and 4-F tapered metal tip. Therefore, simple puncture of the intestinal wall and hepatic parenchyma can be achieved without the need for additional dilation. This stent has an uncovered, funnel-shaped proximal end, a covered body, and a distal portion covered with a silicone membrane. The distal end is equipped with four flaps to prevent inward stent migration. Indeed, none of the stents migrated in that study. Cho et al. described the long-term outcomes of a new hybrid metal stent specifically for EUS-BD which has a 3.5-cm distal portion comprising a silicone-covered nitinol wire to prevent bile leakage. In addition, the covered portion has proximal and distal anchoring flaps to immobilize the stent and prevent proximal and distal migration after placement. The proximal portion, which ranges from 1.5 to 6.5 cm in length, is uncovered to prevent intrahepatic dilatation.
patic bile duct obstruction. The term “hybrid” refers to the two main features of the stent: anchoring flaps that reduce stent migration, and an uncovered portion that decreases the likelihood of obstructive cholangitis. The diameter and length of the hybrid stent are 8–10 mm and 5–10 cm, respectively and the delivery system has a diameter of 8 F. The technical success rate of EUS-HGS, using this stent in 21 patients was 100%, and none of the stents migrated during the procedure or over a median follow-up period of 148.5 days. While these novel stents might positively impact clinical practice, stent migration has recently been reported. Therefore, the length of stents might also impact their migration. We recently evaluated factors associated with stent patency in EUS-HGS in 51 patients. Median stent patency was significantly shorter when the length of the luminal portion was <3 compared with ≥3 cm (52 days vs 195 days, p<0.01). In contrast, median stent patency did not differ significantly between luminal portions ≥4 and <4 cm (194 days vs 127 days, p=0.1726). A luminal portion ≥3 cm in length (hazard ratio [HR], 9.242; 95% confidence interval [CI], 3.255 to 26.244; p<0.05) and the performance of chemotherapy (HR, 3.022; 95% CI, 1.448 to 6.304, p<0.05) were also associated with prolonged stent patency in Cox proportional hazards models. In addition, none of the stents migrated after EUS-HGS. Nakai et al. also described the safety and efficacy of EUS-HGS using a long PCSEMS (LP-CMS) for malignant biliary obstruction in 33 patients. Their study reported a 100% technical success rate with stent lengths of 10 and 12 cm in 76%, and 24% of the patients, respectively. Although the adverse events of bleeding, abscess formation and cholangitis occurred in one patient each, none of the stents migrated either during the procedure or during follow-up.

While the type of metal stent selected is important to prevent migration, technical tips for stent deployment are also important. Technically inappropriate stent deployment can lead to stent migration during deployment. Intra-scope channel release is reportedly clinically useful. We previously used computed tomography to measure distances between the hepatic parenchyma and the stomach wall in patient groups that underwent EUS-HGS with extra-scope (n=20) and intra-scope (n=21) channel release. The distance (mean±standard deviation) between the hepatic parenchyma and stomach wall was significantly shorter after intra-scope than extra-scope channel release (0.66±1.25 cm vs 2.52±0.97 cm). Therefore, although EUS-HGS stents and stent deployment techniques require further improvement, intra-scope channel release and long PCSEMS might be important in preventing stent migration, especially when the procedure is being performed by non-expert endoscopists.

A new plastic stent has become available in Japan. It is a single-pigtail bile duct stent (total length, 20 cm; effective length, 15 cm; four flanges with apertures and side holes; total of 12 holes; distal straight part; four holes and a pigtail site; eight holes) with a tapered tip. Umeda et al. conducted a prospective preliminary feasibility study of EUS-HGS using this stent in 33 patients. The technical success rate was 100% without stent migration during a median follow-up period of 5.0 months, although several adverse events such as self-limiting abdominal pain, occurred. Therefore, they concluded that this new EUS-HGS-dedicated plastic stent was technically feasible and might be applicable in selected patients.

However, plastic stents have several disadvantages as compared to metal stents for EUS-HGS. The diameter of plastic stent is smaller, than that of metal stents, which might shorten the duration of stent patency. this would require discontinuation chemotherapy to treat recurrent biliary obstruction, which, in turn, would influence the survival or quality of life of patients. In addition, if a stent becomes occluded before fistula creation between the hepatic parenchyma and the stomach wall, reintervention might be challenging. Next, the tamponade effect is not obtained because plastic stents are not self-expandable. This function might play important roles in the prevention of bleeding or bile leakage from puncture sites or fistulae. Although these theories should be confirmed by

![Fig. 7.](image-url)
a randomized comparison of metal and plastic stents, we recommend EUS-HGS using metal stents. However, the risk of stent migration, which can lead to critical outcomes in patients, might be higher with these stents.

To overcome this adverse feature of metal stents, EUS-HGS has been combined with antegrade stent (EUS-HGAS) deployment, which has several advantages. Double stent deployment might prolong stent patency compared with EUS-HGS. This is because if one of the stents becomes occluded, recurrent biliary obstruction does not occur due to patency of the second stent. Next, because the antegrade stent is deployed before the EUS-HGS, bile leakage through the fistula is unlikely. However, this feature is also seen if the antegrade stent is deployed without fistula dilation. Therefore, a metal stent with a fine gauge stent delivery system should be selected to realize this benefit. Additionally, if the stent migrates during EUS-HGS deployment, pressure on the biliary tract might be decreased due to the presence of the EUS-guided antegrade stenting (EUS-AS), and hence, conservative treatment might be enough. Indeed, we have experienced case of EUS-HGS stent migration in which conservative treatment alone was required because of the EUS-AS (Fig. 7).

TECHNICAL TIPS FOR REINTERVENTION FOR OCCLUDED EUS-HGS STENTS

Recently, overall survival in patients with hepatobiliary malignancy has been prolonged due to improvement of chemotherapy with drugs such as Folfirinox. Therefore, reintervention for EUS-HGS should be considered in cases with EUS-HGS stent occlusion. Reintervention for occluded EUS-HGS stents depends on the type of stents. If plastic stents are deployed, stent exchange might not be difficult. However, use of long SEMS for EUS-HGS might prevent stent migration. EUS-HGS stents might become occluded due to mucosal hyperplasia at the distal end of the EUS-HGS stent or by the presence of sludge. However, owing to the long length of the stent in the gastric lumen, reintervention might be sometimes challenging. Several authors described their efforts with reintervention techniques. We previously described a simplified reintervention method for EUS-HGS stent obstruction. In our method, the covered site of the FCSEMS is first penetrated by use of a diathermic dilator. Next, an ERCP catheter is inserted and easily advanced into the intestine. Stent placement for the occluded HGS stent is also performed by use of an uncovered metal stent. Minaga et al. also described a similar technique using a precut needle knife. The most suitable technique requires evaluation by further clinical study.

CONCLUSIONS

The rate of adverse events during EUS-HGS is still high, and such events can sometimes be fatal, as with stent migration. Also, bile peritonitis might occur during fistula dilation. Therefore, one-step stent deployment without device exchanges is most ideal. Additionally, to prevent stent migration, improvements in stents, such as lumen apposing formation, is also required. Finally, endoscopists should pay attention not only to technical success, but also preventing adverse events, to improve the clinical benefits of EUS-HGS in the selected patients in whom it is performed.

CONFLICTS OF INTEREST

No potential conflict of interest relevant to this article was reported.

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