Anchoring lumen-apposing metal stent with coaxial plastic stent for endoscopic ultrasound-guided drainage of pancreatic fluid collections: any benefit?

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Abstract

Background Anchoring double-pigtail plastic stents (DPSs) within lumen-apposing metal stents (LAMSs) has been proposed to prevent adverse events during endoscopic drainage of pancreatic fluid collections (PFCs). We sought to compare the outcomes of patients who received LAMSs alone and those who received both LAMSs and anchoring DPSs for drainage of PFCs.

Methods A retrospective study was conducted at the University of Kentucky. Patients with PFCs who underwent endoscopic ultrasound-guided drainage using LAMSs, with or without DPSs, between January 2016 and March 2018 were included. Categorical data were analyzed using chi-square tests, and continuous variables using 2-sample t-tests. Adverse events were defined according to the American Society for Gastrointestinal Endoscopy’s Lexicon. The primary outcome was to evaluate the efficacy (PFC resolution), and safety (adverse events) of LAMSs with or without DPSs used to drain PFCs.

Results Fifty-seven patients with PFCs were treated by 2 experienced endoscopists over 26 months. Twenty-one (37%) patients received LAMSs alone, and 36 (63%) received LAMSs plus DPSs. Forty-three patients had walled-off pancreatic necrosis, and 14 patients had pancreatic pseudocyst. Clinical success (resolution of PFCs) was achieved in 15 patients (71.4%) in the LAMSs alone group, and 21 patients (58.3%) with LAMSs plus DPSs (P=0.32). In patients with LAMSs alone, 6 patients (28.6%) had adverse events, while in those with LAMSs plus DPSs, 14 (38.9%) patients had adverse events (P=0.43).

Conclusion No significant difference was identified in fluid resolution or adverse events between patients with LAMSs alone and those with LAMSs plus DPSs.

Keywords Endosonography, pancreatic diseases, stents, metals

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Introduction

Acute pancreatitis is one of the top 3 gastrointestinal disorders requiring hospital admission in the USA [1,2]. It is a self-limited disease, but can be fatal in patients with organ failure if it persists for more than 2 days [3]. Local complications of acute pancreatitis have been classified according to the revised 2012 Atlanta international consensus [4]. Post-pancreatitis fluid collections (PFCs) that persist for more than 4 weeks and have a well-defined wall are divided into pancreatic pseudocyst (PP), which contains homogenous fluid, and walled-off pancreatic necrosis (WOPN), which is heterogeneous and contains liquid and solid material [4]. Until lately, these collections were drained by needle-knife into the cavity, without or with endoscopic ultrasound (EUS) guidance, and placing multiple double-pigtail stents (DPSSs) [5,6]. Even though the DPSs aid the drainage of liquid material, they are not effective in draining solid material from the cyst cavity and several exchanges are required over guidewires and serial balloon dilatations. In addition, the procedure is time-consuming and relatively challenging. With the introduction of EUS-guided lumen-apposing metal stents (LAMSs) in 2012 (Axios, originally Xlumena Inc, Mountain View, CA; now...
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Boston Scientific, Marlborough, MA), draining PFCs have become much more accessible. Given its large diameter (up to 15 mm), a standard or a therapeutic gastroscope can be passed through the stent into the cavity, allowing necrotic tissue debridement in the setting of WOPN [7,8]. Many centers, including our own, have been placing LAMSs for drainage of PFCs because of their ease of deployment. Although several studies have been published on the use of LAMSs for PFCs, there are limited data on the safety and efficacy of placing coaxial DPSs within LAMSs for draining PFCs [9-12]. Currently, there are no predefined criteria for patient selection as regards using the LAMSs alone or LAMSs plus DPSs during the drainage of PFCs. We sought to compare the outcomes of patients who received LAMSs alone and those who received both LAMSs and anchoring DPSs for EUS-guided drainage of PFCs.

Patients and methods

This was a retrospective cohort study conducted at a tertiary healthcare center from January 2016 until March 2018. Included were patients with symptomatic PFCs who underwent EUS-guided transmural drainage using LAMSs, with or without DPSs. The 2 approaches (LAMS vs. LAMS plus DPS) were compared. The database was retrospectively reviewed to evaluate the safety and efficacy of using the LAMSs with or without DPSs for draining PFCs, after approval had been obtained from the local institute's Internal Review Board. Patient's imaging data, including computed tomography or magnetic resonance imaging, were reviewed along with the EUS findings. PFCs were classified into PP and WOPN according to the revised 2012 Atlanta international consensus [4]. Inclusion criteria were age above 18 years, provision of informed written consent, symptomatic PFCs requiring transmural drainage, as specified by the Working Group of the International Association of Pancreatology [13], and PFCs that persisted for more than 4 weeks with a mature cavity wall on imaging. Exclusion criteria were any other type of fluid collection (fluid was sent for analysis of amylase level, cytology, and carcinoembryonic antigen level during the EUS), immature cavity wall, drainage with stents other than LAMSs, pregnancy, lack of patient stability for endoscopy, and severe coagulopathy or thrombocytopenia.

All EUS-guided drainage procedures of WOPN and PP were performed by 2 interventional endoscopists. The decision to deploy LAMSs alone or with DPSs was at the discretion of the endoscopist. All drainage procedures were performed under general anesthesia with orotracheal intubation. PFCs were evaluated and drained using a linear echo-endoscope (180T; Olympus America, Center Valley, Pa). LAMSs were placed in all patients (AXIOS or Hot AXIOS; 10 or 15 mm × 10 mm; Boston Scientific, Marlborough, Massachusetts, USA) (Fig. 1 A,B,C). Depending on the endoscopist's decision, DPSs (10 Fr × 5 cm; Cook Medical, Bloomington, Indiana, USA) were placed coaxially through the LAMSs (Fig. 2). All patients received intravenous broad-spectrum antibiotics before the procedure. Follow-up imaging studies were performed every 3 weeks. Once the PFCs were entirely resolved, the stents were removed by a repeat endoscopy using grasping forceps (Boston Scientific).

The primary endpoint of this study was to evaluate the efficacy (PFCs resolution) and safety (adverse events) of LAMSs, with or without DPSs, used to drain WOPN and PP. The secondary endpoint was the technical success in placing LAMSs and DPSs. We defined and graded the adverse events according to the American Society for Gastrointestinal Endoscopy's Lexicon [14]. The timing of adverse events was classified as early if it happened during or immediately after the procedure, post-procedure up to 14 days after, and late after 14 days. We defined technical success as the ability to place the stents successfully, and clinical success as the resolution of PFCs on cross-sectional imaging to <2 cm with no further interventions needed, together with an improvement in the patient's symptoms.

Statistical analysis

Categorical data were analyzed using chi-square tests, and continuous variables were analyzed using 2-sample t-tests. A statistical significance level of 0.05 was used, and all analyses were completed in SAS 9.4 (SAS Institute Inc., Cary, NC, USA).

Results

Fifty-seven patients with PFCs (23 female and 34 male; average age 47 years) were treated by 2 advanced endoscopists.
over 26 months. Twenty-one (37%) patients received LAMSs alone, while 36 (63%) patients received LAMSs plus DPSs. Forty-three patients had WOPN, and 14 patients had PP. We classified the size of the PFCs into 3 groups: group 1 (0-50 mm), group 2 (51-100 mm), and group 3 (>100 mm). The average PFCs size in the LAMSs and LAMSs plus DPSs groups were 90 mm and 100.5 mm, respectively (Table 1).

The technical success rate in placing LAMS and DPS was 100%. The drainage site was transgastric in 37 patients, transduodenal in 13 patients, while 7 patients had both (2 multigated and 5 sequential). Clinical success (resolution of PFCs) was achieved in 15 patients (71.4%) in the LAMSs alone group, and 21 patients (58.3%) with LAMSs plus DPSs (P=0.32). The median stent duration was 7 weeks in LAMSs group and 8 weeks in LAMSs + DPSs (Fig. 3, Table 2).

In patients with LAMSs alone, 6 (28.6%) patients had adverse events: 1 (16.6%) bleeding, 1 (16.6%) stent migration, and 4 (66.6%) stent obstruction (P=0.43). In those with LAMSs

Table 1 Demographic data and characteristics of PFCs

| Parameter                      | LAMSs alone (n=21) | LAMSs+DPSs (n=36) | P-value |
|--------------------------------|--------------------|-------------------|---------|
| Age (years) Mean               | 48.95              | 46.61             | 0.59    |
| Sex                            |                    |                   |         |
| Female                         | 10 (47.6%)         | 13 (36.1%)        | 0.39    |
| Male                           | 11 (52.4%)         | 23 (63.9%)        |         |
| Pancreatitis etiology          |                    |                   |         |
| Alcohol                        | 5 (23.8%)          | 6 (16.7%)         | 0.78    |
| Gallstones                     | 8 (38.1%)          | 14 (38.9%)        |         |
| Others                         | 8 (38.1%)          | 16 (44.4%)        |         |
| PPI use                        |                    |                   |         |
| No                             | 12 (57.1%)         | 23 (63.9%)        | 0.61    |
| Yes                            | 9 (42.9%)          | 13 (36.1%)        |         |
| Drainage site                  |                    |                   |         |
| Both                           | 0 (0.0%)           | 7 (19.4%)         | 0.09    |
| Transduodenal                  | 5 (23.8%)          | 8 (22.2%)         |         |
| Transgastric                   | 16 (76.2%)         | 21 (58.3%)        |         |
| PFCs type                      |                    |                   |         |
| Pancreatic pseudocyst          | 7 (33.3%)          | 7 (19.4%)         | 0.24    |
| Walled-off pancreatic necrosis| 14 (66.7%)         | 29 (80.6%)        |         |
| PFCs size (mm)                 |                    |                   |         |
| <50                            | 1 (4.8%)           | 2 (5.6%)          | 0.25    |
| 51-100                         | 14 (66.7%)         | 16 (44.4%)        |         |
| >101                           | 6 (28.6%)          | 18 (50.0%)        |         |
| Nutrition type                 |                    |                   |         |
| Parenteral nutrition           | 0 (0.0%)           | 1 (2.8%)          | 0.34    |
| Oral                           | 12 (57.1%)         | 14 (38.9%)        |         |
| PEG-J                          | 9 (42.9%)          | 21 (58.3%)        |         |
| The average number of necrosectomies | 1.14              | 1.56              | 0.39    |
| Percutaneous drainage          |                    |                   |         |
| No                             | 21 (100.0%)        | 30 (83.3%)        | 0.04    |
| Yes                            | 0 (0.0%)           | 6 (16.7%)         |         |
| Median stent duration (weeks)  | 7.38               | 7.86              | 0.35    |

LAMSs, lumen-apposing metal stents; DPSs, double-pigtail plastic stents; PFCs, pancreatic fluid collections; PPI, proton pump inhibitor; PEG-J, percutaneous endoscopic transgastric jejunostomy
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plus DPSs, 14 (38.9%) patients had adverse events: 2 (14.2%) bleeding, 6 (42.8%) stent migration, and 6 (42.8%) stent obstruction (P=0.43) (Table 3). The most common adverse event was stent obstruction, followed by stent migration and bleeding (Fig. 4). Most of the complications occurred post-procedure, in the first 14 days: 3 (50%) patients in the LAMSs alone and 9 (64.2%) patients in the LAMSs plus DPSs (P=0.28). Infection was not reported among any of the study participants (Table 3).

Discussion

Patients with non-resolving symptomatic PFCs may have a prolonged hospital course with high morbidity and mortality. EUS-guided drainage has become the first method of drainage for symptomatic PFCs, with technical and clinical success rates more than 90% and 75%, respectively [15-18]. The use of LAMSs in EUS-guided drainage of PFCs has been reported to be associated with a higher risk of adverse events compared with other types of stent, and this risk can be reduced by placing a coaxial DPSs with the LAMSs [8,19,20]. Currently, there are no predefined selection criteria for using the LAMSs alone or LAMSs plus DPSs during the drainage of PFCs. In this study, we found that anchoring coaxial DPSs to LAMSs was not associated with a lower rate of adverse events or a higher rate of cyst resolution in EUS-guided drainage of PFCs. Since the introduction of LAMSs, drainage of PFCs has become much easier, because a LAMS’s internal diameter is large enough for the endoscope to pass through, allowing endoscopic

Table 2 Cyst resolution and adverse events

| Parameter          | LAMS alone (n=21) | LAMSs+DPSs (n=36) | P-value |
|--------------------|-------------------|-------------------|---------|
| Cyst resolution    |                   |                   |         |
| (No)               | 6 (28.6%)         | 15 (41.7%)        | 0.32    |
| (Yes)              | 15 (71.4%)        | 21 (58.3%)        |         |
| Adverse events     |                   |                   |         |
| (No)               | 15 (71.4%)        | 22 (61.1%)        | 0.43    |
| (Yes)              | 6 (28.6%)         | 14 (38.9%)        |         |

LAMSs, lumen-apposing metal stents; DPSs, double-pigtail plastic stents

Table 3 Rate and timing of adverse events in patients with LAMSs alone vs. LAMSs plus DPSs

| Parameter          | LAMSs (n = 6) | LAMSs + DPSs (n = 14) | P-value |
|--------------------|---------------|-----------------------|---------|
| Adverse event rate | 6 (28.6%)     | 14 (38.9%)            | 0.43    |
| Stent obstruction  | 4 (66.6%)     | 6 (42.8%)             |         |
| Stent migration    | 1 (16.6%)     | 6 (42.8%)             |         |
| Bleeding           | 1 (16.6%)     | 2 (14.2%)             |         |

Timing of adverse events

| Intra-procedure    | 1 (16.6%) | 0          | 0.28    |
| Post-procedure     | 3 (50%)   | 9 (64.2%)  |
| Late               | 2 (33.3%) | 5 (35.7%)  |

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necrosectomy with a success rate of up to 90% [7,11,15]. LAMSs are used mainly in patients with WOPN where necrosectomy is required, but their role in the drainage of PP is still not entirely clear, as a PP can be effectively drained by DPSs [21]. Endoscopic drainage using DPSs is very effective in PP, with a clinical success rate of 88-98%, but in the case of WOPN the drainage is not optimal, with a clinical success rate of 63-70% [21-23]. According to Siddiqui et al in a recent report, the mean number of procedures needed for resolution of WOPN at 6-months follow up was lower in patients with LAMSs compared with fully covered self-expandable metal stents (FCSEMs) and DPSs 2.2 vs. 3 vs. 3.6, respectively; P = 0.04) [8]. Also, rate of resolution of WOPN at 6-month follow up was lower in patients with DPSs compared with FCSEMs and LAMSs (81% vs. 95% vs. 90%; P=0.001) [8]. FCSEMs have a larger diameter lumen compared with DPSs, and this can allow more efficient drainage with less risk of stent occlusion and superimposed infection. Therefore, they have been used for the drainage of PFCs [24,25]. However, the main disadvantage of FCSEMs was the high migration rate, leading to inefficient drainage and leak [26]. The technical success rate of FCSEMs was 78-100% and the clinical success rate is above 80% [8,18,22,23,27]. According to Sharaiha et al [15], 16 patients with DPSs were about 2.9 times more likely to have adverse events than those patients with FCSEMs.

LAMSs are designed with a bi-flanged shape that allows for tissue apposition and minimizes the risk of stent migration. They have a wide diameter lumen that provides a non-compressible fistulous tract between the gut lumen and the pancreatic fluid cavity. In addition, it can provide the channel for endoscopic necrosectomy and the need for repeated endoscopies. The clinical and technical success rates of LAMSs have been reported to be 93-100% and 89-100%, respectively [7,11,23,25,28-30]. LAMSs have become the stent of choice for endoscopic drainage of PFCs by many gastroenterologists, because of their easy deployment and direct debridement access. According to Siddiqui et al [7], in a multicenter retrospective study that assessed EUS-guided drainage of symptomatic PFCs in 82 patients, the adverse event rate was 9.8%. The adverse events among these patients were 2 maldeployed stents, 6 episodes of bleeding, 5 episodes of PFC infection, and 4 stent occlusions. Also, per Siddiqui et al [8] compared FCSEMs vs. LAMSs vs. DPSs for the drainage of WOPN. The results showed that LAMSs were more likely to be associated with early adverse events compared with FCSEMs (P=0.02) and bleeding was the most frequently reported complication. Recently, a pilot study by Aburajab et al [31] reported that EUS-guided drainage of PP via LAMSs is associated with higher rates of infection and nonresolution, but these adverse events can be minimized by placing DPSs across the LAMSs.

In our study, bleeding was not the most frequent event. It was reported in only one patient in the LAMSs group, during the procedure, and in 2 patients with LAMSs plus DPSs, with 1 episode occurring post-procedure and the other late. The most reported adverse event in both groups was stent obstruction by necrotic debris. There was no significant difference between LAMSs and LAMSs plus DPSs in terms of adverse events, while the median stent duration was 7 weeks in the LAMSs group and 8 weeks in the LAMSs plus DPSs group.

The limitations of this study are its retrospective design with a small sample size, a single-center experience, selection bias, no well-known selection criteria for the LAMSs alone or LAMSs plus DPSs approaches (the endoscopist made the decision on which approach to use), and generalizability limitations.

In conclusion, this study showed that anchoring coaxial DPSs to LAMSs was not associated with a higher rate of fluid resolution during endoscopic ultrasound-guided drainage of PFCs. Further studies with larger sample sizes are needed to validate the results.
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