Successful treatment of cervical esophageal leakage by endoscopic-vacuum assisted closure therapy

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AIM: To evaluate the efficacy and safety of endoscopic-vacuum assisted closure (E-VAC) therapy in the treatment of cervical esophageal leakage.

METHODS: Between May and November 2012, three male patients who developed post-operative cervical esophageal leakage were treated with E-VAC therapy. One patient had undergone surgical excision of a pharyngo-cervical liposarcoma with partial esophageal resection, and the other two patients had received surgical treatment for symptomatic Zenker’s diverticulum. Following endoscopic verification of the leakage, a trimmed polyurethane sponge was fixed to the distal end of a nasogastric silicone tube and endoscopically positioned into the wound cavity, and with decreasing cavity size the sponge was positioned intraluminally to cover the leak. Continuous suction was applied, and the vacuum drainage system was changed twice a week.

RESULTS: The initial E-VAC placement was technically successful for all three patients, and complete closure of the esophageal leak was achieved without any procedure-related complications. In all three patients, the insufficiencies were located either above or slightly below the upper esophageal sphincter. The median duration of the E-VAC drainage was 29 d (range: 19-49 d), with a median of seven sponge exchanges (range: 5-12 sponge exchanges). In addition, the E-VAC therapy reduced inflammatory markers to within normal range for all three patients. Two of the patients were immediately fitted with a percutaneous enteral gastric feeding tube with jejunal extension, and the third patient received parenteral feeding. All three patients showed normal swallow function and no evidence of stricture after completion of the E-VAC therapy.

CONCLUSION: E-VAC therapy for cervical esophageal leakage was well tolerated by patients. This safe and effective procedure may significantly reduce morbidity and mortality following cervical esophageal leakage.

Key words: Endoscopic-vacuum assisted closure therapy; Vacuum therapy; Negative pressure wound therapy; Cervical esophageal leakage; Anastomotic leakage

Core tip: Traditional methods to treat cervical esophageal leakage close to the upper esophageal sphincter are associated with high morbidity and mortality. The newly developed method of endoscopic-vacuum assisted closure (E-VAC) therapy using polyurethane sponges has been demonstrated as efficacious for treating gastrointestinal tract leakages. We applied E-VAC therapy to three patients with post-operative cervical leakage and achieved complete closure in all, without any procedure-related complications. The E-VAC therapy was well tolerated by patients with cervical esophageal leakage, and its application in this patient population may contribute to a significant reduction in morbidity.
and mortality.

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INTRODUCTION

Anastomotic leakage is a potentially life-threatening complication that may follow esophageal surgery. The leakage may range in severity from a minor anastomotic defect to a fulminant leak with systemic sepsis and multiple organ failure. Cervical anastomoses have been associated with leakage rates as high as 40% and a mortality rate of 5%.[1-3] The treatment of cervical anastomotic leakage above the upper esophageal sphincter is particularly challenging, and only limited treatment options are available. Traditionally, the repair of cervical leakage has involved surgical intervention;[4] however, re-operation is associated with high morbidity and mortality rates.[5] Placement of self-expandable metal stents in such situations is difficult or even impossible and is associated with a high rate of procedure-related complications, such as globus sensation and/or respiratory insufficiency. Therefore, the procedure is often not performed.[6-11]

Over the last decade, several endoscopic treatment options for repair of esophageal anastomotic leakages have emerged, including fibrin glue injection, endoscopic transluminal drainage and self-expanding metal stents.[12-14] Endoscopic treatment using self-expandable metal or plastic stents has become the treatment of choice for cervical anastomotic leakage, and its reported success rates are above 80%.[15-18] Most recently, endoscopically-assisted vacuum closure (E-VAC) has been suggested as an effective treatment modality for esophageal anastomotic leakage in the upper gastrointestinal tract.[19] E-VAC therapy involves placing polyurethane sponges into the wound cavity that was induced by the leak, followed by application of an external vacuum through a transnasal tube to drain the infected fluid and induce the formation of granulation tissue. Recent studies of E-VAC therapy for the treatment of leaks following esophageal anastomoses have demonstrated that the procedure is capable of achieving successful wound closure with no associated mortality.[20-23] However, these studies have mainly examined intrathoracic anastomotic leakages. Here, we report the successful application of E-VAC therapy to treat cervical anastomotic leakages in three patients.

MATERIALS AND METHODS

Patients and procedure description

Between May and November 2012, three male patients with post-operative cervical esophageal leakage were treated with E-VAC therapy at the Endoscopy Unit of the Hannover Medical School (Hannover, Germany). E-VAC placement was performed as described previously,[24] as the modified form of the VAC technique, which is an established treatment modality for chronic and infected cutaneous wounds.[25-28] Briefly, a trimmed polyurethane sponge, pore size 400-600 μm (KCI, Wiesbaden, Germany) was fixed to the distal end of a nasogastric silicone tube (Freka 15 Ch; Fresenius Kabi, Bad Homburg, Germany) and introduced into the cavity under endoscopic vision. With decreasing cavity size, the sponge was placed endoluminally to cover the entire esophageal defect. A continuous negative pressure of 125 mmHg was applied using a vacuum pump (KCI). The vacuum drainage system was endoscopically changed two times per week. All endoscopic interventions were performed either under general anesthesia or conscious sedation with propofol and midazolam. All three patients gave informed consent for publication of their case, and retrospective analysis was performed in accordance with the Declaration of Helsinki.

Descriptive statistics were used to evaluate the patients’ demographic and clinical characteristics. The data are presented as individual values, median, and ranges.

RESULTS

Characteristics of patients

We used E-VAC therapy to treat three male patients with post-operative cervical esophageal leakage. The patients were 69-, 71- and 80-year-old (Table 1). Patient 1 had undergone surgical excision of a pharyngo-cervical liposarcoma with partial esophageal resection followed by an insufficiency 3 cm below the upper esophageal sphincter (17 cm from the incisors). Patients 2 and 3 had suffered from cervical esophageal perforation following surgical treatment of a symptomatic Zenker’s diverticulum. Patient 2 had open surgery with a diverticulectomy and myotomy (Figure 1). Patient 3 suffered from recurrent Zenker’s diverticulum and was treated with transoral endoluminal mucotomy. The insufficiency in these two cases was located above the upper esophageal sphincter, at 17 cm from the incisors in patient 2 and at 19 cm from the incisors in patient 3.

Results of E-VAC therapy

All three patients had endoscopically diagnosed esophageal leakage and their initial E-VAC placement was technically successful. In all three cases, the sponge was initially placed into the extraluminal cavity (intracavitary), which was changed to intraluminal placement with decreasing cavity size. Two patients immediately received a percutaneous enteral feeding tube with jejunal extension (PEG-J tube) and the third patient received parenteral feeding (Table 2). The median duration of E-VAC therapy was 29 d (range: 19-49 d) with a median of seven sponge exchanges (range: 5-12 sponge exchanges) (Table
Median hospitalization time was 46 d (range: 42-108 d). In all three patients, complete closure of the leakage was achieved without any procedure-related complications and without the need for surgical re-intervention (Figure 2). Sponge therapy was well tolerated and there was no evidence of residual leakage either clinically or after Gastrografin swallow in patients 2 and 3. Inflammation was assessed by measuring white blood cell (WBC) counts and C-reactive protein (CRP) levels. In two patients, the WBC count was initially elevated but decreased to within the normal range following E-VAC therapy. All three patients had markedly elevated CRP levels (range: 152-296 mg/L) at the beginning of the treatment, which were reduced to almost normal (range: 3-34 mg/L) by the time of discharge (Table 3). Patients were clinically followed-up after hospital discharge and endoscopy was performed in two patients at post-discharge days 47 and 206. All three patients had normal swallow function and no evidence of stenosis after completion of the E-VAC therapy.

**DISCUSSION**

Esophageal anastomotic leakage is associated with high morbidity and mortality rates, particularly when surgical repair is required[1,4,8]. Consequently, efforts have been made to devise less invasive treatment modalities. A number of endoscopic techniques have emerged in recent years, including E-VAC therapy. Here, we report the successful use of E-VAC therapy for the treatment of post-
without any procedure-related complications. None of the patients required further surgical intervention, and all three patients displayed regular swallow function after completion of the E-VAC therapy. Follow-up endoscopy in patients 2 and 3 demonstrated complete healing of the esophagus.

These case series indicate that E-VAC therapy has clinical utility in the repair of cervical esophageal leakage. These data justify conducting further studies to examine the potential of E-VAC therapy for treating other iatrogenic cervical esophageal perforations, such as perforations after transesophageal echocardiography, foreign body impaction, or endoscopic and surgical procedures. Compared to the previous studies of E-VAC therapy for treating thoracic esophageal leakage\cite{20,26}, our case studies of E-VAC therapy for treating cervical esophageal leakage required longer treatment times and a higher number of sponge changes. Therefore, we recommend early PEG placement for enteral feeding. However, despite the longer treatment times, the E-VAC therapy was well tolerated by all of our patients.

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Our case studies suggest that use of E-VAC therapy allows for rapid removal of infected tissue. Prior to E-VAC therapy, all three patients displayed high levels of inflammatory markers that were indicative of systemic inflammatory complications from the esophageal leakage. Notably, a considerable reduction in the levels of these inflammatory markers was observed following treatment, without any procedure-related complications. None of the patients required further surgical intervention, and all three patients displayed regular swallow function after completion of the E-VAC therapy. Follow-up endoscopy in patients 2 and 3 demonstrated complete healing of the esophagus.

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which suggests that the E-VAC therapy resulted in rapid drainage of the infected wound cavity and control of inflammation.

In summary, we report that E-VAC therapy is a safe and efficacious treatment option for cervical esophageal leakage. E-VAC therapy appears to provide adequate wound drainage, promotion of tissue granulation within the wound cavity, and closure of the cervical esophageal defect. Despite the high localization of the vacuum placement, sponge therapy was well tolerated by our patients. Application of this therapy may contribute a significant improvement in morbidity and mortality. A multidisciplinary approach, involving the coordinated efforts of abdominal and/or ear-nose-throat surgeons, may further enhance E-VAC therapy as a treatment modality for cervical esophageal leakage.

COMMENTS

Background
Traditionally, the repair of cervical esophageal leakage has involved surgical intervention, as placement of self-expandable metal stents in this situation is difficult or even impossible. Most recently, endoscopic-vacuum assisted closure (E-VAC) has been suggested as an effective treatment modality for esophageal leakage. Therefore, the authors investigated the efficacy of E-VAC therapy for cervical leakage above or slightly below the upper esophageal sphincter.

Research frontiers
Cervical esophageal leakage is associated with high morbidity and mortality rates, particularly when surgical repair is required. Therefore, this study evaluated the effectiveness and safety of a non-invasive endoscopic treatment using E-VAC therapy for treating cervical esophageal leakage.

Innovations and breakthroughs
This study demonstrates that E-VAC therapy is an efficacious and safe treatment option for treating cervical esophageal leakage. Despite the high localization of the vacuum placement, the sponge therapy is well tolerated.

Applications
E-VAC therapy can be used as an alternative treatment option for cervical esophageal leakages above or slightly below the upper esophageal sphincter. These findings indicate the benefit of future studies addressing whether E-VAC therapy may also be useful for treatment of other iatrogenic cervical esophageal perforations, such as perforations after transesophageal echocardiography, foreign body impaction, or endoscopic and surgical procedures.

Terminology
The VAC technique is an established treatment modality for chronic and infected cutaneous wounds. Recently, the endoscopic placement of a vacuum-assisted closure system (endoscopic-vacuum assisted closure, E-VAC) in the gastrointestinal tract has been shown to be an effective treatment option for anastomotic leaks. The trimmed polyurethane foam with an open-cell structure (sponge) is fixed to the distal end of a silicone duodenal tube and endoscopically introduced into the necrotic cavity of the upper or the lower gastrointestinal tract. A continuous negative pressure of 125 mmHg is applied using a vacuum-assisted closure system (endoscopic-vacuum assisted closure, E-VAC) in the upper or the lower gastrointestinal tract. A continuous negative pressure of 125 mmHg is applied using a vacuum-assisted closure system (endoscopic-vacuum assisted closure, E-VAC) in the upper or the lower gastrointestinal tract. A continuous negative pressure of 125 mmHg is applied using a vacuum-assisted closure system (endoscopic-vacuum assisted closure, E-VAC) in the upper or the lower gastrointestinal tract.

Peer review
The authors conclude that E-VAC therapy is a safe and effective treatment option for cervical esophageal leakage.

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