Single-Phase Percutaneous Recanalization of Malignant Bile Duct Obstructions with a Covered Stent Graft

Perkutane Rekanalisation von malignen DHC Gallengangsobstruktionen mit einer gecoverten Stentprothese im einzeitigen Verfahren

Abstract

Purpose: The aim of this study was to evaluate the benefit of a percutaneous transhepatic biliary drainage (PTCD) endoprosthesis in the case of patients with malignant occlusion of the common bile duct (CBD).

Materials and Methods: 32 patients (mean age 72 ± 13 years) were treated with an endoprosthesis (VIABIL; M. L. GORE & ASSOCIATES, INC., USA) due to failed attempts of endoscopic retrograde cholangiopancreatography (ERCP) in the case of malignant occlusion of the CBD.

Results: The technical success rate was 96.9%. In one patient the probing of an intrahepatic bile duct was impossible. Two major complications (bleeding, liver abscess) were successfully treated with appropriate measures. The bilirubin level did not significantly decrease immediately after intervention (13.2 ± 6.5 mg/dl; p > 0.05). However, the follow-up displayed a highly significant decrease of bilirubin to 6.0 ± 7.4 mg/dl; p < 0.05). The endoprosthesis was extended with bare metal NITINOL stents in 9 patients. The mean survival time of the patient group was 64 ± 28 days (range 2 – 250 days).

Conclusion: The implantation of an endoprosthesis proved to be an option with high technical success, a low complication rate and good benefit in our patients with malignant bile duct obstruction in palliative therapy situations.

Key Points:

- The primary objective in the case of malignant bile duct obstruction is the treatment of jaundice.
- After failed endoscopic recanalization of the bile ducts, transhepatic biliary drainage is desirable.
- An ePTFE-FEP covered endoprosthesis is a good treatment option in palliative situations.
- A single-stage procedure shortens hospitalization time.

Zusammenfassung


Material und Methoden: Bei 32 Patienten (mittleres Alter 72 ± 13 Jahre) mit kausal maligner Okklusion des Ductus choledochus (DHC) und frustranem Versuch der endoskopischen retrograden Cholangiopancreatographie (ERCP) wurde die Indikation zur perkutanen transhepatischen Cholangiodrainage (PTCD) und Implantation einer Endoprothese (VIABIL der Firma M. L. GORE & ASSOCIATES, INC., USA) gestellt.

Ergebnisse: Die technischen Erfolgsraten lag bei 96.9%. Bei einem Patient war die Sondierung eines intrahepatischen Gallengangs nicht möglich. Zwei Majorkomplikationen (Nachblutung, Leberabszess) konnten durch entsprechende Maßnahmen erfolgreich behandelt werden. Der Gesamtbilirubin war mit 13,2 ± 6,5 mg/dl unmittelbar postinterventionell nicht signifikant erniedrigt (p > 0.05). Im Follow-up zeigte sich jedoch eine signifikante Erniedrigung des Gesamtbilirubin auf 6,0 ± 7,4 mg/dl (p < 0.05). Bei 9 Patienten wurde die Stentprothese mittels Baremetal Nitinolstents verlängert. Die mittlere
Überlebenszeit des Patientenkollektivs lag bei 62 ± 71 Tagen (range: 2 – 250 Tage).

**Schlussfolgerung:** Die interventionelle Implantation einer Endoprothese stellt bei Patienten mit maligner Gallengangsoberstruktur in einer palliativen Therapiesituation und frustranter endoskopischer Sondierung eine Option mit gutem Benefit dar.

**Introduction**

In the majority of cases, malignant bile duct obstructions represent a palliative situation with an unfavorable prognosis for the affected patient [1]. They can be caused by pancreatic carcinomas, bile duct tumors, or metastases, lymphomas, gallbladder tumors, tumors or the stomach and duodenum, and hepatocellular carcinoma [2, 3]. The primary goal of palliative treatment of malignant bile duct stenoses is to improve symptomatic jaundice and limit the risk of cholangitis and renal and hepatic collateral damage [4]. A further improvement of clinical symptoms can be achieved by supplementary palliative chemotherapy and radiation therapy [3].

Gastrointestinal obstructions usually require bypass surgery, but this is associated with a relatively high mortality rate in palliative situations [4, 5]. Endoscopic and radiological interventional procedures are the treatments of choice for malignant bile duct stenosis with indication for the procedure ideally being determined by a multidisciplinary panel of experts [2, 4]. In addition to the different interventional procedures, prosthesis materials in the form of plastic or metal stents are available [5].

The goal of this retrospective study was to evaluate the clinical outcome of a percutaneous transhepatic endoprosthesis in malignant obstruction of the common bile duct (CBD) in a palliative treatment situation.

**Materials and Methods**

The data of 32 patients (average age: 72 ± 13 years) with a malignant bile duct obstruction in a palliative treatment situation between 2009 and 2012 were included in this retrospective study. In addition to the primary disease and obstructive jaundice, the inclusion criteria were unsuccessful endoscopic recanalization attempts and the standards for performing minimally invasive image-guided interventions of the German Society of Interventional Radiology and Minimally Invasive Therapy (DeGIR) [6]. Surgical treatment of the bile duct obstruction was not indicated in any of the patients included in the study. The location of the bile duct obstruction was categorized according to the Bismuth classification [7]. The survival and hospitalization time, the serum bilirubin in the follow-up period, interventional complications, the cause of death, and the rate of restenosis were the most important examination parameters of this study. The serum bilirubin was determined in a standardized manner before and one day after intervention. In the follow-up period, the bilirubin value was recorded as a function of the hospitalization time and the ambulant follow-up examinations. The interventions were performed by three different radiologists with a high level of interventional procedure expertise.

**Treatment method**

The access for percutaneous transhepatic cholangiography (PTC) was planned preinterventionally via a plain CT scan of the liver. All patients were treated prophylactically with a single-shot i. v. antibiotic of 2 grams ceftriaxone prior to the intervention. The intervention was performed under sterile conditions and with cardiovascular monitoring of the patient after intravenous analgesic infusion with 2 – 5 mg midazolam and local anesthesia with 20 – 30 ml mepivacaine. After puncture of the bile duct with a 21G introducer needle and dilation of the access for insertion of a 4.5 French lock (AccuStick from Flexima Biliary®, Terumo), a 0.035” Terumo Stiff guidewire was carefully advanced into the duodenum and the 4.5 French lock was replaced by a 6 French lock (Radiofocus Introducer 2®, Terumo). PTC was then performed to visualize the length, the location of the bile duct occlusion, and the cystic duct (Fig. 1a, b). Pre-dilation of the bile duct stenosis was performed with a 5/80 balloon catheter (Mustang, Boston Scientific) and a change to a 10 French lock (Radiofocus Introducer 2®, Terumo) with a working length of 25 cm was made. A self-expanding 8/60 mm ePTFE-FEP (expanded polytetrafluoroethylene fluorinated ethylene propylene) covered stent (VIABL®, W. L. GORE & ASSOCIATES) was implanted in the region of the CBD with the distal end of the prosthesis extending beyond the papilla of Vater by less than 2 cm. The post-dilation of the VIABL endoprosthesis...
was performed with a balloon catheter with a diameter of 7 mm and a length of 80 mm (Mustang®, Boston Scientific) (Fig. 2a–d).

In the case of very long stenoses of the extrahepatic bile duct, a second VIABIL endoprosthesis with the same nominal diameter and same length was implanted in individual cases. If the bile duct occlusion continued in a central direction into the hepatic fork or in the case of an additional stenosis, a self-expanding nitinol stent (S. M. A. R. T.®, Cordis) of varying size was used and post-dilated accordingly (Fig. 3a, b). The flow of the contrast medium into the duodenum was documented. Remaining stenosis < 30% was evaluated as a technical success after angiographic examination. Follow-up examinations via ultrasound and/or computed tomography were only performed in the case of a recurrence of symptomatic jaundice.

**Statistical analysis**

The results of this study are provided as mean ± standard deviation. The Wilcoxon test was used as a statistical test for comparing quantitative data. The significance level was defined as < 0.05.

**Results**

In total, a VIABIL endoprosthesis was able to be successfully percutaneously transhepatically implanted in 31 patients. The primary technical success rate was 96.9%. An intrahepatic bile duct could not be probed in one patient. In 21 of 31 cases the cause of the occlusion was the primary tumor or the primary disease followed by liver metastases in 7 cases and lymph node metastases in 4 of 31 cases. De-
pending on the length of the bile duct stenosis, two VIABIL endoprostheses were implanted in individual cases using the stent-in-stent technique. Intrahepatic and proximal extrahepatic stenoses were treated with uncovered nitinol stents. The VIABIL endoprosthesis had to be extended proximally by nitinol stents in 10 cases. The median bilirubin value prior to the intervention was 14.5 ± 6.5 mg/dl. In the laboratory test one day after the intervention, no significant reduction of the total bilirubin was observed (median 13.2 ± 6.5 mg/dl; p > 0.05).

However, further follow-up examinations showed a highly significant reduction in the bilirubin value (p < 0.05) to 6.0 ± 7.4 mg/dl in 30 of 31 patients (Fig. 4). In the postinterventional observation period, two patients had significantly increased serum bilirubin values that were higher than prior to the intervention. Hemobilia was diagnosed in both patients in the follow-up examinations and was treated by temporary external percutaneous transhepatic biliary drainage (PTCD). A liver abscess developed postinterventionally in one case and was successfully treated via drainage and i.v. antibiotics. Perinterventional bleeding with a non-clinically relevant drop in hemoglobin did not progress over time. Tumor infiltration of a nitinol stent required a new percutaneous transhepatic cholangiography with probing and balloon dilation of the restenosis in one central bile duct carcinoma (Table 1). Otherwise, no additional restenosis of the stent implant was seen in the observation period. The average survival time of the patient collective was 62 ± 71 days (range: 2 – 250 days) with an average total hospitalization time of 14 ± 11 days (range: 1 – 40 days). 16 of 29 patients died within 30 days. The cause of death was unclear in 22 of 29 cases of death at home. The cause of death was cardiac in 5 of 29 patients. One patient died of a pulmonary artery embolism, one of upper gastrointestinal hemorrhage, and one of multiple organ failure due to sepsis. 2 of 31 patients survived to the end of data collection.

**Discussion**

During palliative treatment of malignant bile duct stenoses, the primary goals are to reduce symptomatic jaundice and to prevent cholangitis and liver failure [8]. The postinterventional serum bilirubin is an independent predictor for the prognosis of the patient. Brontzos et al. defined the cut-off value for a poor prognosis at 4 mg/dl [9, 10]. Compared to benign stenoses, malignant bile duct stenoses are located distal to the liver hilum or in the extrahepatic course of the bile ducts in particular [10]. Pancreatic carcinomas and distal cholangiocellular carcinomas (CCCs) are typical tumors of this bile duct segment. The group of malignant obstructive tumors of the proximal bile ducts and the hilar region includes gallbladder carcinomas, advanced gastric carcinomas, lymphomas, liver metastases, and CCCs [2].

In the case of non-resectable malignant bile duct tumors, PTCD with intrahepatic and/or extrahepatic biliary drainage is indicated in addition to endoscopic recanalization. However, due to the lower complication rate and the lower strain on the patient with successful reduction of jaundice, endoscopic recanalization is considered the primary procedure [11, 12].

For the recanalization of inoperable malignant bile duct occlusions, the use of metal stents is considered state of the art [13]. Temporary PTCD without the use of self-expanding stent systems is performed in the case of benign bile duct stenosis [12]. The advantages of self-expanding metal stents compared to plastic stents are the greater inner lumina, the option to use the stent-in-stent technique, lower reocclusion rates, shorter hospitalization times, and the lower cost [13 – 15]. Fanelli et al. came to the conclusion that ePTFE-FEP covered stents provide an effective and feasible means of inner biliary drainage in the case of malignant bile duct obstruction and are resistant to tumor ingrowth but not tumor overgrowth [13]. This complication can be counteracted by overstenting [16]. The disadvantages of covered metal stents are the greater risk of cholecystitis or pancreatitis when overstenting the cystic duct or accessory bile ducts [2]. ePTFE-FEP covered stents were only used in the extrahepatic course of the common bile duct in our study. Stenoses in a more proximal position were treated with uncovered nitinol stents. In individual cases, two VIABIL

<table>
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Tab. 1 Complications after PTC and stent implantation.
endoprotheses were implanted prophylactically using the stent-in-stent technique to prevent possible tumor overgrowth. We prefer the single-phase approach with stent implantation and internal biliary drainage since the strain on the patient in a palliative situation and the hospitalization time, which was 14 days on average in this study, can be reduced. Fanelli et al. used a two-phase approach in most cases when implanting ePTFE-FEP covered stents to reduce the risk of hemorrhage and to better determine the location of the bile duct stenosis [13]. External PTCD was performed in two of our patients with hemobilia in order to drain the blood clots. At 97%, the technical success rate of the single-phase approach was within the range presented in the literature which was between 97% and 100% [3, 13, 17, 18]. The bilirubin value that was only minimally reduced on average one day after the intervention was not interpreted as stent occlusion but as an expression of the stimulation of the bile duct system. The bilirubin value fluctuations within the first week after stent implantation are not to be interpreted as an indication of stent blockage [3]. The total bilirubin value of the patients, which was determined on average 40 days after the intervention during follow-up, can be discussed as an indication for restenosis. However, this was the last value before death and the most common cause of death was the advanced tumor disease. According to the literature, the patency rates of ePTFE-FEP covered stents are between 28.5% and 100% in a period between 3 and 12 months after intervention [13, 16, 19]. In light of the average survival time of 62 days in our patient collective, the development of restenosis is not to be expected. The mortality rate within the first 30 days after percutaneous implantation of metal stents is between 11.5% and 36% according to the literature [13, 20, 21]. The complication rate of percutaneously implanted metal stents ranges between 6% and 61% [17, 21–23]. Endoscopically implanted stent prostheses have a complication rate of 10% to 42% [21, 24]. In particular, acute (<24 h) and early (<30 days) complications include sepsis, hemobilia, bile leakage, hemorrhage, cholecystitis, and liver abscess. Late complications (>30 days) primarily include stent occlusions [3]. Fanelli et al. administered antibiotic prophylaxis before and for 5–6 days after the intervention and did not describe any infection-related complications at a complication rate of 6.5% [13]. In the majority of cases, the complications of percutaneously implanted bile duct prostheses can be treated conservatively [2]. Of the 4 observed complications that could be attributed to PTCD in the patient collective examined here, only one case of sepsis was lethal. In addition to the progress of the tumor disease, cardiovascular causes of death that could not be linked to the interventional procedure were seen in particular. Restenosis due to tumor overgrowth is a general problem of all stents [25, 26] that is minimized by the ePTFE-FEP covered endoprosthesis as a result of its special design with an anchoring system consisting of a loop-shaped metal wire [13]. However, these stents are also not immune to tumor overgrowth and intervention may be required in individual cases. In addition to prophylactic overstenting using the stent-in-stent technique, recanalization via the laser technique is possible in individual cases [27]. Another possible application of covered stent prostheses in addition to the recanalization of malignant bile duct obstructions could be the occlusion of CBD fistulas or bile duct leaks as a supplement to percutaneous onyx application [28].

> **Conclusion**
>
> Single-phase percutaneous implantation of an ePTFE-FEP covered endoprosthesis in inoperable patients with malignant cholestasis in a palliative treatment situation represents a safe and effective alternative method to ERCP. Indication should be determined by interdisciplinary consensus of gastroenterologists, surgeons, and interventional radiologists.

> **Clinical relevance of the study**
>
> - Percutaneous implantation of an ePTFE-FEP covered endoprosthesis is a good and safe treatment option for malignant bile duct obstruction.
> - In palliative treatment situations, the single-phase procedure shortens hospitalization time and improves quality of life.
> - In a palliative situation, restenosis is rare in the case of stent prostheses.

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**References**


Dinkel HP, Triller J. Primary and long-term success of percutaneous biliary metallic endoprotheses (Wallstents) in malignant obstructive jaundice. Fortschr Röntgenstr 2001; 173: 1072 – 1078


