Endoscopic treatment of chronic pancreatitis: European Society of Gastrointestinal Endoscopy (ESGE) Guideline – Updated August 2018

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Appendix 1s, Tables 1s – 8s
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MAIN RECOMMENDATIONS

ESGE suggests endoscopic therapy and/or extracorporeal shockwave lithotripsy (ESWL) as the first-line therapy for painful uncomplicated chronic pancreatitis (CP) with an obstructed main pancreatic duct (MPD) in the head/body of the pancreas. The clinical response should be evaluated at 6–8 weeks; if it appears unsatisfactory, the patient’s case should be discussed again in a multidisciplinary team and surgical options should be considered.

Weak recommendation, low quality evidence.

ESGE suggests, for the selection of patients for initial or continued endoscopic therapy and/or ESWL, taking into consideration predictive factors associated with a good long-term outcome. These include, at initial work-up, absence of MPD stricture, a short disease duration, non-severe pain, absence or cessation of cigarette smoking and of alcohol intake, and, after initial treatment, complete removal of obstructive pancreatic stones and resolution of pancreatic duct stricture with stenting.

Weak recommendation, low quality evidence.

ESGE recommends ESWL for the clearance of radiopaque obstructive MPD stones larger than 5mm located in the head/body of the pancreas and endoscopic retrograde
cholangiopancreatography (ERCP) for MPD stones that are radiolucent or smaller than 5 mm. Strong recommendation, moderate quality evidence.

ESGE suggests restricting the use of endoscopic therapy after ESWL to patients with no spontaneous clearance of pancreatic stones after adequate fragmentation by ESWL. Weak recommendation, moderate quality evidence.

ESGE suggests treating painful dominant MPD strictures with a single 10-Fr plastic stent for one uninterrupted year if symptoms improve after initial successful MPD drainage. The stent should be exchanged if necessary, based on symptoms or signs of stent dysfunction at regular pancreas imaging at least every 6 months. ESGE suggests consideration of surgery or multiple side-by-side plastic stents for symptomatic MPD strictures persisting beyond 1 year after the initial single plastic stenting, following multidisciplinary discussion. Weak recommendation, low quality evidence.

ESGE recommends endoscopic drainage over percutaneous or surgical treatment for uncomplicated chronic pancreatitis (CP)-related pseudocysts that are within endoscopic reach. Strong recommendation, moderate quality evidence.

ESGE recommends retrieval of transmural plastic stents at least 6 weeks after pancreatic pseudocyst regression if MPD disruption has been excluded, and long-term indwelling of transmural double-pigtail plastic stents in patients with disconnected pancreatic duct syndrome. Strong recommendation, low quality evidence.

ESGE suggests the temporary insertion of multiple side-by-side plastic stents or of a fully covered self-expandable metal stent (FCSEMS) for treating CP-related benign biliary strictures. Weak recommendation, moderate quality evidence.

ESGE recommends maintaining a registry of patients with biliary stents and recalling them for stent removal or exchange. Strong recommendation, low quality evidence.

1 Introduction

The Clinical Guideline on the endoscopic treatment of chronic pancreatitis (CP) published in 2012 by the European Society of Gastrointestinal Endoscopy (ESGE) made recommendations on the indications and modalities of treatment for CP [1]. New evidence has become available since then and is discussed in the present update, and new recommendations are issued.

2 Methods

ESGE commissioned this Guideline and appointed a Guideline leader (J.M.D.) who invited the listed authors to participate in the project development. The key questions were prepared by the coordinating team (J.M.D., A.T., M.D.) and then approved by the other members. The coordinating team formed task force subgroups, each with its own leader, who was assigned key questions (see Appendix 1s, online-only Supplementary Material).

Each task force performed a systematic literature search to prepare evidence-based and well-balanced statements on their assigned key questions. The literature search was performed using MEDLINE and Embase to identify new publications since January 2012 published in English. The Grading of Recommendations Assessment, Development and Evaluation (GRADE) system was adopted to define the strength of recommendation and the quality of evidence [2]. Each task force proposed statements on their assigned key questions which were discussed during a meeting in Brussels, Belgium, in June 2017. Literature searches were re-run in August 2018. This time-point should be the starting point in the search for new evidence for future updates to this Guideline. In August 2018 a draft prepared by J.M.D. was sent to all group members for review. The draft was also reviewed by two members of the ESGE Governing Board, by external reviewers, and by the ESGE National Societies and
Individual Members. After agreement on a final version, the manuscript was submitted to the journal Endoscopy for publication. All authors agreed on the final revised version.

This Guideline was issued in 2018 and will be considered for review in 2022, or sooner if new and relevant evidence becomes available. Any updates to the Guideline in the interim period will be noted on the ESGE website: https://www.esge.com/esge-guidelines.html.

3 Choice of treatment and initial work-up

**RECOMMENDATION**

ESGE suggests endoscopic therapy and/or extracorporeal shockwave lithotripsy (ESWL) as the first-line therapy for painful uncomplicated chronic pancreatitis (CP) with an obstructed main pancreatic duct (MPD) in the head/body of the pancreas. The clinical response should be evaluated at 6–8 weeks; if it appears unsatisfactory, the patient’s case should be discussed again in a multidisciplinary team and surgical options should be considered. Weak recommendation, low quality evidence.

The first step proposed to relieve pain in patients with uncomplicated CP includes lifestyle modifications plus, in selected patients, endoscopic therapy and/or ESWL [3]. If endoscopic therapy and/or ESWL provide no persistent pain relief or technically fail, or if the patient is not a good candidate for endoscopic therapy and/or ESWL, medical treatment including analgesics and adjunctive agents (e.g., pharmaceutical agents aimed to relieve neuropathic pain) are proposed, with the final step being early surgery for nonresponders. In one large prospective multicenter U.S. cohort (n = 521), medical therapy, endoscopic therapy, and pancreatic surgery were performed in 69%, 52%, and 18% of patients, respectively [4]. Similarly, in 33 series of CP patients treated with endoscopic therapy and/or ESWL, surgery was performed during long-term follow-up in a minority of patients, less frequently in those with stones as the main obstructing factor (117 of 1695 [6.9%], 13 series, Table 1a, see Supplementary Material, online-only) as compared to those with strictures (157 of 1061 patients [14.8%], 20 series, Table 2a; P<0.001).

Two trials have suggested that surgery was superior to endoscopic therapy and/or ESWL for pain relief [5–7]. In the first trial [5], pain was absent after 5 years of follow-up in 15% vs. 34% of patients treated with endoscopic therapy vs. surgery, respectively, showing that neither of these options is entirely satisfactory. Furthermore, neither ESWL nor cumulative stenting were used and the randomized design of the study is questionable. In the other trial [6,7], only 39 patients were included; all of them had advanced CP and most were opioid-dependent. For these reasons, the results cannot be extrapolated to all patients with CP. A cost–effectiveness model based on data of this randomized controlled trial (RCT) unsurprisingly concluded that surgery was more effective and less costly than endoscopic therapy in CP [8], but another RCT has shown that ESWL could provide satisfactory clinical results at a relatively low cost in patients with obstructive stones in the main pancreatic duct (MPD) (62% of patients with no pain relapse at 4-year follow-up after ESWL) [9]. Finally, a retrospective study (86 CP patients) reported similar pain relief 5.4 years after endoscopic therapy and/or ESWL vs. surgery, but surgery carried more complications and higher costs [10].

Endoscopic therapy and/or ESWL aim to relieve an obstruction in the MPD. They are proposed only to patients with marked ductal changes, mainly dilation, corresponding to the most severe grade in the Cambridge classification of pancreatitis [11]. No recent publications have reported the results of endoscopic therapy in patients with less severe changes [12–13]. In painless CP, endoscopic therapy and/or ESWL are not performed because the only potential benefit (preserving the pancreatic function) is uncertain: a single prospective nonrandomized comparative study examined this in 42 CP patients and found that the mean value of the N-benzoyl-L-tyrosyl para-aminobenzoic acid test was higher at 5-year follow-up after stenting vs. no stenting of a MPD stricture while no differences were observed for overt diabetes [14]. These results have not been confirmed and in most long-term studies the pancreatic function deteriorated during follow-up [15–19].

**RECOMMENDATION**

ESGE suggests, for the selection of patients for initial or continued endoscopic therapy and/or ESWL, taking into consideration predictive factors associated with a good long-term outcome. These include, at initial work-up, absence of MPD stricture, a short disease duration, a short disease duration, non-severe pain, absence or cessation of cigarette smoking and of alcohol intake, and, after initial treatment, complete removal of obstructive pancreatic stones and resolution of pancreatic duct stricture with stenting. Weak recommendation, low quality evidence.

During the pretherapeutic evaluation of a patient, factors associated with a good long-term clinical outcome may help to select patients for endoscopic therapy and/or ESWL. These factors should be considered as orientative only as the differences in the proportions of patients with long-term success for an individual factor are small. The factors include absence of MPD stricture (see above) as well as short disease duration, non-severe pain (including low dose use of narcotics), the absence or cessation of cigarette smoking and of alcohol intake, cephalic location of stones, the absence of pancreas divisum if MPD stenting is required, and steatorrhea (4, 4, 3, 2, 1, and 1 studies, respectively) (Table 3a). Favorable prognostic factors related to endoscopic therapy and/or ESWL include complete stone removal and MPD stricture resolution after stenting (2 and 1 studies, respectively).

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Patients with an MPD obstruction located only in the tail of the pancreas are not considered candidates for ESWL and/or endoscopic therapy by some groups of authors [20].

**RECOMMENDATION**

ESGE suggests performing a high quality pancreatic computed tomography (CT) scan and/or magnetic resonance imaging with cholangiopancreatography to reasonably rule out pancreatic cancer and to plan treatment in patients with chronic pancreatitis.  
Weak recommendation, low quality evidence.

The risk of pancreatic cancer is increased in patients with CP, particularly in the first years following diagnosis [21]. A meta-analysis (52 studies, 5399 patients) found that endoscopic ultrasonography (EUS), CT scan, and magnetic resonance imaging (MRI) present similar diagnostic accuracies for the diagnosis of pancreatic cancer [22]. Imaging methods of the pancreas are constantly refined and they are often used in combination [23, 24]. In the particular context of CP, MRI with diffusion-weighted imaging has shown sensitivity and specificity for the diagnosis of malignancy of 86% and 82%, respectively, in a meta-analysis [25], while EUS-guided sampling seems to be less sensitive according to a retrospective and a prospective study (54% and 74% vs. 89% and 91% in the presence vs. the absence of CP, respectively) [26, 27]. The yield of EUS elastography and contrast-enhanced harmonic EUS as well as methods to improve the accuracy of EUS-guided sampling are discussed in dedicated ESGE Guidelines [28, 29].

Non-contrast CT scan accurately delineates calcified stones in the pancreas and allows measurement of stone density, a factor associated with the completeness of stone extraction [30]. Contrast enhancement may help to locate stones relative to the ducts [31, 32]. Magnetic resonance cholangiopancreatography (MRCP) identifies ductal abnormalities; in two retrospective studies its diagnostic accuracy for ductal abnormalities was 73.2% (41 children with CP) and 92.2% (30 adults with CP) [33, 34].

4 Pancreatic stone management

Pancreatic stones seem to arise as either direct and evenly calcified stones or as radiolucent protein plugs that may or may not become calcified during the course of the disease [35]. The vast majority of pancreatic stones are calcified and radiopaque; their prevalence increases with time to reach 50% and 100%, at 5 and 14 years after the onset of the disease, respectively [36]. In a multicenter survey (879 CP patients with a mix of newly diagnosed and long-standing disease), calcified pancreatic stones were detected in 62% of patients; they were more frequent in men, heavy drinkers (>80g/day), and heavy smokers (>20 cigarettes/day) [37]. Pancreatic stones in CP patients who undergo endoscopic therapy and/or ESWL are solitary in 10% – 62% of patients; they are most frequently located in the pancreatic head only, with a mean size of 10 mm, and they are associated with strictures in approximately 50% of patients (Table 4s).

Successful stone fragmentation following ESWL has been defined as stones broken into fragments ≤2 or 3 mm, or by the demonstration at X-ray of decreased stone density, increased stone surface, and heterogeneity of the stone which may fill the MPD and adjacent side branches [38]. Ductal clearance has been defined as complete, partial, or unsuccessful if the proportion of stones cleared was >90%, 50%–90%, or <50%, respectively [39].

**RECOMMENDATION**

ESGE recommends ESWL for the clearance of radiopaque obstructive MPD stones larger than 5 mm located in the head/body of the pancreas, and endoscopic retrograde cholangiopancreatography (ERCP) for MPD stones that are radiolucent or smaller than 5 mm.  
Strong recommendation, moderate quality evidence.

Endoscopy alone, using pancreatic sphincterotomy and a basket or a balloon, allows stone extraction in a minority of CP patients: 9% of 1041 patients in two retrospective series [40–41] and 14% of 1834 patients in a survey of 125 hospitals [42]. Failed stone extraction using these techniques is associated with stones >10 mm, diffuse location, stone impaction, and location upstream from a stricture [41, 43]. Furthermore, pancreatic mechanical lithotripsy carries a complication rate threefold higher compared with biliary mechanical lithotripsy according to a retrospective study of 712 patients [44]. Complications in the 69 patients with pancreatic stones included trapped or broken basket, traction wire fracture, and one pancreatic ductal leak which required surgery [44]. In one of the above-mentioned series, ESWL allowed the endoscopic extraction of pancreatic stones in >80% of the patients after failed stone extraction at primary endoscopy [40]. Similarly, a retrospective study (70 patients) found that performance of ESWL prior to the endoscopic attempt at stone extraction was the only independent factor associated with successful stone clearance [45]. Therefore, a primary endoscopic attempt at pancreatic stone extraction is reserved to selected patients, based on a reasonable expectation of success or on technical difficulty in performing ESWL as with radiolucent stones or stones <5 mm that are difficult to target using X-rays.

A meta-analysis (27 studies including 6 prospective ones, in total 3189 patients with pancreatic stones >5mm) reported that pancreatic ESWL allowed complete/partial MPD clearance in 70%/22% of patients, respectively, that pain was absent or mild-moderate during the 2 years following treatment in 52.7% and 33.4% of patients, respectively, and that quality of life improved after ESWL in 88.2% of patients [39]. ERCP was combined with ESWL in most studies. Table 1 summarizes the outcomes of ESWL alone or combined with endoscopic stone extraction. Pain relapsed in 30% – 50% of patients dur-
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Pancreatic stone fragmentation is obtained after ESWL in approximately 90% of patients [50]; this may require multiple ESWL sessions (up to 8 in a large series with a high rate of successful fragmentation) [20]. More shockwaves may be required for stones that are larger [51], multiple [52], or associated with a MPD stricture [53], while pancreatic stenting prior to ESWL seems to decrease the number of shockwaves and of ESWL sessions required [51]. Multicenter surveys have suggested that stone fragmentation is less frequently successful in low case volume centers while the role of the type of lithotripter has been controversial [41, 42, 49].

After ESWL, endoscopic clearance of stone fragments has been more frequently successful with solitary stones [17, 20, 30, 45, 53], stones located in the pancreatic head [20], stones with a density at CT scan of <820.5 Hounsfield units [30], if a pancreatic stent had been inserted prior to ESWL [54, 55], if secretin had been administered at the beginning of ESWL [55], and if ERCP was delayed by more than 2 days after ESWL [56]. Pancreatic pseudocysts (PPCs) did not affect stone clearance or adverse events in a prospective series of 849 patients (59 with a PPC) [57].

The most frequent complication of ESWL is pancreatitis; it has been reported in 4.2% of the patients in a meta-analysis, but most of the included studies were retrospective and did not allow the attribution of complications to either endoscopic therapy or ESWL as both were performed in most patients [39]. In a prospective study (634 patients, 1470 ESWL sessions), transient adverse events (asymptomatic hyperamylasemia, hematuria, gastrointestinal mucosal injury) and complications were detected in 21.2% and 6.7% of the procedures, respectively [58]. Complications included pancreatitis, infection, steinstrass (acute stone incarceration in the papilla), bleeding, and perforation; they were classified as moderate or severe in 1.1% of the cases. Skin erythema and tenderness in the region in contact with the shockwave head were noted in most patients [58].

Contraindications to ESWL include non-correctable coagulation disorders, pregnancy, and presence in the shockwave path of bone, calcified vessels, or lung tissue [59]. Specific precautions should be taken for patients with implantable defibrillators and pacemakers [60].

Reports of intracorporeal lithotripsy using electrohydraulic or laser lithotripsy under peroral pancreatoscopy are sparse. A systematic review (10 studies, 87 patients) reported successful MPD clearance in 43% – 100% of patients [61]. Results may be biased as the reports included selected patients with anatomical features thought to permit passage of the pancreatoscope to the target stone in a stable position. The largest study reported complete and partial stone clearance in 24 (63%) and 10 (26%) of 38 patients, respectively, after a total of 280 endoscopic therapy sessions, including 88 with pancreatoscopy; complications (post-ERCP pancreatitis and one perforation) were reported for 20 procedures and the overall clinical success rate was 74% [62].

RECOMMENDATION
ESGE suggests restricting the use of endoscopic therapy after ESWL to patients with no spontaneous clearance of pancreatic stones after adequate fragmentation by ESWL. Weak recommendation, moderate quality evidence.

ESWL: technical factors, complications and contraindications

Pancreatic stone fragmentation is obtained after ESWL in a follow-up of 1–14 years and surgery was performed in 6.9% of patients. Of note, the studies that reported the timing of pain relapse showed that patients with no pain relapse at 2-year follow-up rarely experience pain relapse thereafter [9, 16, 45], in particular if stone clearance has been complete [30]. Approximately half of patients with relapsing pain present with stone recurrence [46].

Weak recommendation, low quality evidence.
5 Pancreatic strictures

Since the previous publication of this Guideline no new definitions of the types of MPD strictures in CP have been reported. Besides benign vs. malignant and single vs. multiple, strictures may be classified as either non-dominant or dominant [63]. Dominant MPD strictures are defined by the presence of at least one of the following characteristics: upstream MPD dilatation ≥ 6 mm in diameter, prevention of contrast medium outflow alongside a 6-Fr catheter inserted upstream from the stricture, or abdominal pain during continuous infusion of a nasopancreatic catheter inserted upstream from the stricture with 1 L saline for 12–24 h.

Stent insertion across a dominant MPD stricture (or the most proximal [tail] one in the case of multiple strictures) defines technical success. It aims to: (i) decompress the MPD, thereby ameliorating pain, and (ii) persistently dilate the stricture(s). Less frequent indications include facilitation of MPD stone clearance in association with ESWL as detailed above, and to bypass an obstruction in the ventral duct by inserting a stent through the minor papilla into the MPD [64]. A prospective non-randomized study showed in 42 patients with a dominant MPD stricture that pain recurred less frequently in patients who had received a temporary pancreatic stenting vs. those who had not (15 % vs. 50 % during a 5-year follow-up) [14]. Before stent dilation therapy is embarked upon, malignancy should be reasonably excluded, for example by brush cytology and cross-sectional imaging (see Section 3).

Refractory MPD strictures are defined as symptomatic dominant strictures that persist or relapse after 1 year of single pancreatic stent placement. A validated short-term definition for clinical success is still lacking. For long-term evaluation, the absence of pain during the year following stent removal still seems a reasonable and workable definition.

RECOMMENDATION

ESGE suggests treating painful dominant MPD strictures with a single 10-Fr plastic stent for one uninterrupted year if symptoms improve after initial successful MPD drainage. The stent should be exchanged if necessary, based on symptoms or signs of stent dysfunction at regular pancreas imaging at least every 6 months. ESGE suggests consideration of surgery or multiple side-by-side plastic stents for symptomatic MPD strictures persisting beyond 1 year after the initial single plastic stenting, following multidisciplinary discussion.

Weak recommendation, low quality evidence.

Insertion of a single plastic stent has been used as the initial endoscopic therapy for MPD strictures (Table 2s); these strictures were single in >80 % of the patients [65–66], and some studies explicitly excluded patients with multiple strictures [67]. After temporary insertion of a single plastic stent in the MPD, stricture resolution was achieved in 9 % [68] to 50 % [6] of 145 patients in five studies [6, 67–70] but this is not required for long-term pain relief [67]. Long-term pain relief was reported in 67.5 % of 536 patients (95 % confidence interval [CI] 51.5 %–80.2 %) in a meta-analysis of 9 studies [71]. The follow-up duration after stent removal was not calculated but in most studies it was ≥24 months, the period during which almost all pain relapses occur [6, 14, 66, 70, 72–74].

Refractory strictures may be treated by surgery, multiple side-by-side plastic stents (Table 5s), or self-expandable metal stents (SEMSs) (Table 6s).

The temporary insertion of multiple side-by-side plastic stents in 48 patients yielded stricture resolution and pain relief at 9.5-year follow-up in 89.5 % and 77.1 % of the patients, respectively [75–76].

With respect to SEMSs, uncovered and partially covered types have provided disappointing results [77] but temporary placement of a fully covered SEMS (FCSEMS) has provided pain improvement in 85 % of patients according to a systematic review of four prospective series (total 61 patients) [78]. These studies were limited by a very short follow-up, and three more recent studies (n = 41) have reported pain improvement in 37 %–88 % of patients during a follow-up of 3–4 years [79–81]. Pancreatic FCSEMS need further evaluation in the setting of clinical trials because of potential complications as listed below.

Pancreatic stenting: technical factors and complications

Whether or not a pancreatic sphincterotomy should be performed before pancreatic stent insertion has not been addressed in any study, but both methods have been reported for the insertion of a single plastic stent as well as for a SEMS [18, 65, 79, 82–85]. With respect to the performance of a biliary sphincterotomy prior to pancreatic sphincterotomy, this should only be performed in selected cases, according to a small RCT, mostly if biliary drainage is indicated or to facilitate access to the MPD [86].

In many but not all studies [51, 54, 55], pancreatic stenting was performed after stone fragmentation and removal. In prospective series, technical success was reported in 92 % of attempted insertions of a first stent [6, 14, 67, 87]. The stenting duration averaged 10.6 months (range 3.2–23 months) in 18 series totaling 811 patients [5, 6, 14, 64–67, 70, 72–74, 82, 87–92].

Multiple stent designs have been proposed, including straight, S-shaped, and winged stents, and stents with or without sideholes [93–94]. Few comparative studies have been reported; in a prospective study, stents with large sideholes have been suggested to occlude less frequently compared to other types, but only a minority of patients had CP [95]. With respect to stent diameter, CP patients treated with stents ≤ 8.5-Fr were 3.2 times more likely to be hospitalized for abdominal pain than those who had received 10-Fr stents in a retrospective study of 163 CP patients [96].

“On-demand” stent exchange consists of exchanging pancreatic stents when deemed necessary, based on patient symptoms and/or additional investigations at 1–6-month intervals (i.e., secretin-enhanced MRCP [S-MRCP] [66], abdominal ultra-
sound alone [68] or supplemented either with abdominal plain film [66] or with blood/urinary amylase measurements [69]). With this stent exchange policy, sepsis of pancreatic origin was reported in 15 (5.2%) of 288 patients in four series [66,68–69,72] and surgery was required in two patients for pancreatic abscesses; this was reported in the only series in which no additional investigations at regular intervals were performed [72]. On the other hand, in 12 series (521 patients) with stent exchange scheduled at shorter intervals, usually 3 months, septic complications have not been reported [5,14,65,67,73,74,88,90,92,97,98].

Compared with surgery, hospital stays and medical expenses were similar for patients who had pancreatic stenting for less than 1 year (n=19) but higher for those who required longer pancreatic stenting (n=15), in a retrospective study [97]. In that study, a single plastic stent was re-inserted at 3 months. However stent obstruction does not correlate with symptoms [82,89,100]. Stent-induced ductal lesions were reported at short term (average 6.2%, range 4%–39%) followed by sepsis, cholangitis, and post-sphincterotomy bleeding (average, 2.6%, 2.3%, and 1.5%, respectively) (Table 2). Severe pancreatitis or worsening of pancreatic pain were most commonly reported at short term (average 6.2%, range 4%–39%) followed by sepsis, cholangitis, and post-sphincterotomy bleeding (average, 2.6%, 2.3%, and 1.5%, respectively) (Table 2s). Severe pancreatitis has been rarely reported [73]. During follow-up, proximal and distal stent migration is reported in 2.7% and 3.6% of cases respectively, and bench tests using a column of water at a pressure lower than that observed in patients with CP [89,100] have shown that almost all stents become obstructed at 3 months. However stent obstruction does not correlate with symptoms [82,89,100]. Stent-induced ductal lesions were described in 18% of patients (range 0–26%) and mortality was reported in 0.4% (7/1620) (Table 2).

With SEMSs, stent migration (15%–46%) and de novo strictures (16%–27%) have also been reported and specific complications include severe pain (7%–20%) leading to cholestasis and FCSEMS removal (15%) (Table 6s).

### RECOMMENDATION

ESGE recommends performance of endosonography-guided access and drainage of the MPD only in tertiary centers after multidisciplinary discussion and preferably in a research setting.

Strong recommendation, low quality evidence.

Potential indications for endosonography-guided access and drainage of the MPD include patients with symptomatic MPD obstruction and failed conventional transpapillary drainage. Briefly, the technique consists of puncturing the MPD through the gastric or duodenal wall and advancing a guidewire into the MPD to proceed with transpapillary (rendezvous technique) or transmural drainage using a plastic stent [50], or more recently a FCSEMS [101]. It is recognized as one of the most difficult techniques of EUS-guided therapy [102].

Endosonography-guided access and drainage of the MPD has been reported in retrospective, small, single-center studies [103–107] or larger multicenter studies (36 to 80 patients) [108–110] with a follow-up ranging from a few weeks up to 55 months (median 1 year). In all these series, the annual inclusion rate per center was always below 4, illustrating the rarity of the indications.

Immediate pain relief after successful endosonography-guided access and drainage of the MPD has been reported in a majority of patients with obstructive CP (range 50%–100%). In the two series to date with available long-term follow-up, complete or major pain relief was achieved in 70%–90% of patients but the probability of remaining free of pain dropped sharply over time [108,109].

Failed endosonography-guided access and drainage of the MPD occurs in approximately 10% of cases and the incidence of moderate to severe complications also averages 10% in the largest series, including severe pancreatitis, perforation, bleeding, and hematoma [103–110]. No procedure-related mortality has been reported. Migration and occlusion of stents necessitating endoscopic re-intervention frequently occur (20%–55% of patients).

### 6 Pseudocyst management

Approximately one third of CP patients develop PPC during the course of their disease [36]. PPCs should be differentiated from cystic neoplasms such as potentially malignant mucinous neoplasms, particularly when they present for the first time.

Endoscopic therapy of PPCs consists of inserting a drain from the digestive lumen into the PPC, through the digestive wall ("transmural drainage"), through the papilla ("transpapillary drainage"), or using a combination of these routes. Transpapillary PPC drainage is feasible only if the PPC communicates with the MPD, a situation detected in approximately half of PPCs [111]. Technical and clinical success are usually defined, respectively, as the insertion of at least one stent between the PPC and the digestive lumen (plus removal if indicated) [112], and disappearance of symptoms with complete resolution of the PPC or a decrease in size to less than 2 cm [113].
Spontaneous regression of chronic PPCs is infrequent (0 to 27%) and occurs most commonly for PPCs smaller than 4 cm and/or located within the pancreas [114–115]. The indications for treatment listed above are commonly accepted. In asymptomatic patients with a PPC compressing a major vessel, the risk–benefit ratio of any intervention should be thoroughly analyzed; progressively enlarging collections are considered a valid indication by some authors while others suggest that such patients be followed until symptoms develop [116–117].

A meta-analysis of 7 retrospective studies (490 patients with various types of pancreatic fluid collections [PFCs]) found that, compared with percutaneous drainage, endoscopic drainage was associated with a higher clinical success rate, fewer re-interventions, shorter hospital stay, and similar morbidity and recurrence rates [118]. Although percutaneous drainage has mostly been abandoned for the definitive treatment of CP-related pseudocysts because it often results in an external fistula [119], it may be useful as an emergency measure (e.g., for infected PPC not accessible to endoscopic drainage in a frail patient).

A meta-analysis (5 comparative studies including one RCT, 255 patients) found that, compared with endoscopic therapy, surgery has a higher success rate (odds ratio [OR] 0.43, 95% CI 0.20–0.95), but is associated with a longer length of hospital stay and higher hospital costs as well as similar rates of morbidity (18.0% vs. 11.5%) and recurrence (3.2% vs. 3.1%) [120]. A more recent multicenter prospective cohort study (71 patients) reported a similar overall success rate and a shorter hospital stay for endoscopic therapy vs. surgery [121].

CT scan, MRI, and EUS allow the characterization of PFCs but the assessment of their solid content is less precise with CT scan [122–124]; this is important only in subacute PFCs where necrotic debris may impede endoscopic drainage. S-MRCP also allows diagnosis of MPD rupture. This has important consequences for treatment planning: (i) in the absence of MPD rupture, endoscopic drainage can be transmural only; (ii) if a partial MPD rupture is present, insertion of a stent bridging the rupture (as opposed to below it) is associated with treatment success [63,64]; and (iii) in the case of a complete MPD rupture (disconnected pancreatic duct syndrome), removal of transmural stents is associated with PFC recurrence so that long-term indwelling of transmural double-pigtail plastic stents should be considered [125,126]. Therefore, some centers perform imaging of the MPD by S-MRCP and/or ERCP prior to drainage of and/or stent removal from PFCs.

Although ERCP is still considered to be the gold standard for the diagnosis of MPD disruption, it presents limitations including an accuracy rate of approximately 75% and adverse events such as infection of a sterile PFC [127,128]. In small series, S-MRCP showed an accuracy of >90% for diagnosing MPD disruption in patients with PFCs [123,129].

These imaging modalities have not been compared for the detection of pseudoaneurysms close to pseudocysts, which is another important consideration when planning treatment.

Compared with transmural drainage, transpapillary drainage provides similar success with a similar morbidity rate but fatal or surgical complications are less frequent (1/176 vs. 15/283; P = 0.007); however, transpapillary drainage as the only endoscopic therapy has been performed for relatively smaller collections (generally ≤50 mm) than those managed by transmural drainage alone or combined transpapillary and transmural drainage (Table 7s). If transmural drainage is performed, the addition of transpapillary drainage seems to add no benefit according to a meta-analysis of 9 non-randomized comparative studies (7 including PPCs exclusively, 604 drainage procedures).
No definitive conclusion can be drawn as the proportion of patients in whom a transpapillary stent was inserted across as opposed to below a partial MPD disruption, a predictor of success following transpapillary drainage [131,132], was not known. However, this factor may be of marginal importance as the insertion of a stent across a partial MPD rupture succeeds in only 33%–67% of the patients [131,132].

**RECOMMENDATION**

ESGE recommends endosonography-guided over conventional access for the transmural drainage of CP-related pseudocysts.

Strong recommendation, moderate quality evidence.

For the transmural drainage of PPCs, a systematic review (four studies, 229 patients) found a higher technical success rate for EUS vs. conventional approach (relative risk [RR] 12.38, 95%CI 1.39–110.22) and no other significant differences (complications, short and long-term clinical success) [133]. The difference in technical success was due to the presence of non-bulging collections which account for approximately half of PFCs [111]; EUS guidance is the only option for transmural drainage in these cases.

**RECOMMENDATION**

ESGE suggests the use of double-pigtail plastic stents for the transmural drainage of CP-related pseudocysts; a fully covered biliary SEMS can be considered if disconnected pancreatic duct syndrome has been excluded and indwelling duration is expected to be less than 6 weeks.

Weak recommendation, low quality evidence.

Plastic stents are generally used for the transmural drainage of PPCs. Three retrospective studies examined the role of the number or diameter of plastic stents in a total of 307 patients; all studies included biliary patients with various types of PFCs [134–136]. Double-pigtail stents of 7–10 Fr were used in the two most recent series as straight stents may migrate and erode large vessels [135]. One study found that the insertion of a single stent was associated with failure of endoscopic therapy (defined as severe procedure-related complication or need for another treatment modality) [135] while two studies found no differences according to the number and diameter of plastic stents [134,136].

Plastic stents and FCSEMSs have been compared for the transmural drainage of PPCs in three meta-analyses [113,137,138]. The two most recent meta-analyses included comparative studies exclusively but only approximately 10% of patients had CP. These two meta-analyses reported: (i) a similar success and a lower morbidity rate with FCSEMSs vs. plastic stents (OR 0.4, 95%CI 0.21–0.73) (three studies, 301 patients) [138]; and (ii) a higher success rate with FCSEMSs vs. plastic stents (OR 5.35, 95%CI 1.35–21.19) (morbidity analysis not reported) (two studies, 250 patients) [113]. Biliary FCSEMSs were used in most patients while lumen-apposing metal stents (LAMSs) were used in 5% [138] and 6% of the patients [113]; in the studies that used a standard biliary FCSEMS, a double-pigtail plastic stent was inserted through the FCSEMS to prevent its migration. The older meta-analysis included non-comparative studies only and it found no differences between stents in terms of success rates or morbidity [137].

A meta-analysis (6 retrospective studies, 504 patients) compared LAMSs with multiple plastic stents for the treatment of PFCs but only 11% of patients had a PPC; LAMSs were associated with a higher clinical success rate (RR 2.70, 95%CI 1.49–5.00) and a lower morbidity rate (RR 0.39, 95%CI, 0.18–0.84) [139]. A decision model analysis concluded that LAMSs were less cost-effective than plastic stents [140].

**RECOMMENDATION**

ESGE recommends retrieval of transmural plastic stents at least 6 weeks after pancreatic pseudocyst regression if MPD disruption has been excluded, and long-term indwelling of transmural double-pigtail plastic stents in patients with disconnected pancreatic duct syndrome.

Strong recommendation, low quality evidence.

Transmural plastic stents are generally removed at least 6 weeks after insertion as a retrospective study showed that earlier plastic stent removal was associated with treatment failure [135]. In an RCT (28 patients, 15 of whom had a CP-related PPC), PFCs recurred more frequently in patients randomized to stent removal 2 months after drainage vs. no stent removal (38% vs 0); PFC recurrence tended to be associated with MPD rupture as identified at S-MRCP (4/5 vs 2/9, P=0.063) [126].

Disconnected pancreatic duct syndrome generally results from severe necrotizing pancreatitis and has been discussed in a dedicated ESGE Guideline [141]. Retrospective studies have shown that long-term indwelling of double-pigtail transmural plastic stents is effective, with PFC recurrence being uncommon and associated with stent migration <6 months after PFC resolution and MPD disruption at the pancreatic head level [142,143].

With respect to LAMSs, an RCT reported LAMS-related severe adverse events in 6 of 12 patients (50%), including bleeding, buried stent syndrome, and biliary stricture, all occurring >3 weeks after LAMS placement [144,145]. Stent-related morbidity dropped to levels similar to those observed with plastic stents after the study protocol was changed to removal of LAMSs within 4 weeks. The placement of a coaxial double-pigtail stent through the LAMS has also been proposed to prevent delayed adverse events [146].

Extrahepatic portal hypertension develops during the course of CP in ≥ 15% of patients [147]. The only two series that reported the results of endoscopic drainage for PFCs in patients with portal hypertension used EUS guidance; bleeding was reported in 1 of 26 patients (4%) [148, 149].

Pseudoaneurysms complicate the course of CP in 1%–10% of patients, mostly those with a PPC, and their rupture is associated with a high mortality [150]. Therefore, some authors recommend embolization of arterial pseudoaneurysms before attempting endoscopic therapy of PPC close to pseudoaneurysms [151]. This strategy has not been tested but, in patients with bleeding pseudoaneurysms, two retrospective series have reported a 94%–100% mid-term success rate with arterial embolization followed by endoscopic therapy of the PPC in a total of 40 patients [152, 153].

7 Biliary strictures

**RECOMMENDATION**
ESGE suggests performance of an ERCP when a CP patient presents with a ≥ 4-week biliary obstruction (jaundice, asymptomatic elevation of serum alkaline phosphatase (> 2 or 3 times the upper limit of normal values) and/or bilirubin) to achieve biliary decompression by means of stent placement. If follow-up shows that the obstruction is caused by a genuine fibrosis rather than transient inflammatory compression, endoscopic stent treatment should be continued in order to dilate the stricture. After 1 year of unsuccessful endotherapy, surgery should be considered.

Weak recommendation, low quality evidence.

Biliary strictures complicate the course of CP in 3%–23% of patients, with studies reporting a prevalence as high as 46% [154]. Symptoms may be absent or include jaundice, cholangitis or choleodocholithiasis. Jaundice resolves spontaneously in 20%–50% of patients within 1 month, because of resolution of edema or of a PPC in the head of the pancreas but secondary biliary cirrhosis is relatively frequent (7.3% of 288 patients in a review of 11 studies) [154]. Therefore, an asymptomatic elevation of serum alkaline phosphatase (>2 or 3 times the upper limit of normal values) and/or bilirubin for longer than 1 month are usually accepted as an indication for bile duct drainage [155].

As underlined in Section 3, an underlying malignancy should be reasonably excluded.

A single retrospective study compared surgery vs. endoscopic therapy (multiple side-by-side plastic stents or FCSEMS) for the treatment of CP-related biliary strictures in 39 patients [156]. Compared with surgery, endoscopic therapy presented a lower procedural morbidity rate (21% vs. 83%) and a lower success rate at 2 years (15% vs. 66%). The success rate was noticeably lower than in other studies (Table 8s), including an RCT, maybe because incomplete stricture resolution at ERCP was considered a failure. Outcomes were similar in patients who had surgery as a primary treatment or following unsuccessful endoscopic therapy. The authors proposed to attempt endoscopic therapy first in the absence of associated lesions (e.g., inflammatory cephalic mass), and to evaluate its success after 12 months or three endoscopic procedures.

**RECOMMENDATION**
ESGE suggests the temporary insertion of multiple side-by-side plastic stents or of a FCSEMS for treating CP-related benign biliary strictures.

Weak recommendation, moderate quality evidence.

The strategy of endoscopic therapy for benign biliary strictures is detailed in a dedicated ESGE Clinical Guideline [157]; it consists of temporarily dilating the stricture using multiple side-by-side plastic stents or a FCSEMS (single plastic stents or uncovered SEMSs have long been abandoned because of poor long-term results (Table 8s) [158]. An RCT (60 CP patients) found that multiple plastic stents and covered SEMSs provided similar success rates 2 years after stent removal (88.0% vs. 90.9%, respectively), with similar treatment-related morbidity (23.3% vs. 28.6%, respectively) [159]. The stenting duration was 6 months in both groups. Various stenting durations have not been compared in the literature (scheduled stenting durations with multiple plastic stents and covered SEMSs have generally been for 1 year and for 6–12 months, respectively). Short biliary strictures may respond better than longer ones to stenting, as suggested by a small study (10 CP patients) [160].

**RECOMMENDATION**
ESGE recommends maintaining a registry of patients with biliary stents and recalling them for stent removal or exchange.

Strong recommendation, low quality evidence.
Patient compliance with stent exchange may be poor, giving rise to potentially fatal complications [161, 162]. To prevent this, various recall systems have proven useful in pilot studies [163, 164]. Removable FCSEMSs can result in better patient compliance since the number of ERCPs is reduced to two. Of course, patient compliance with repeat interventions should be ensured prior to endoscopic therapy and hepaticojejunostomy remains a valid option for noncompliant patients or if the stricture does not respond to endoscopic therapy.

Disclaimer
The legal disclaimer for ESGE guidelines [165] applies to the current Guideline.

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Competing interests

G. P. Aithal receives consultancy fees from Shire (September 2015 to present), Pfizer (July 2018 to present), and GSK and Agios (February 2018 to present). A. Anderloni has provided consultancy to Boston Scientific (2017–2018). M. J. Bruno has received lecturing and consultancy fees from Boston Scientific, Cook Medical, and Pentax Medical (ongoing) and consultancy fees from Mylan (ongoing); his department is involved in investigator- and industry-initiated studies with Boston Scientific, Cook Medical, and Pentax Medical (ongoing); he is a member (no financial benefit) of the Dutch Pancreatitis Study Group. J. Devière receives research support from Olympus for institutional review board-approved studies (ongoing); his department receives research support from Boston Scientific for institutional review board-approved studies (ongoing). J. E. Domínguez-Muñoz has received speaker’s honoraria from Boston Scientific (2018); his department has received financial support for educational activities from Pentax and Boston Scientific (2017–2018) and Medtronic (2018). J.-W. Poley receives speaker’s fees and travel expenses from Pentax, Boston Scientific, and Cook Endoscopy (ongoing), and consultancy fees from Boston Scientific and Cook Endoscopy (ongoing). A. Sanchez-Yague has provided paid consultancy to Boston Scientific (2015–2018). J. E. van Hooft has received lecture fees from Medtronic (2014–2015) and consultancy fees from Boston Scientific (2014–2016); her department has received research grants from Cook Medical (2014–2018) and Abbott (2014–2017). M. Arvanitakis, P. Cantú, M. Delhaye, J.-M. Dumonceau, S. Lekkerkerker, M. Ramchandani, N. Reddy, A. Tringali, and T. Vaysse have no competing interests.


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