Improving the Outcome of Acute Pancreatitis

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Key Words
Acute pancreatitis · Acute necrotizing pancreatitis · Biliary pancreatitis

Abstract
Acute pancreatitis (AP) is the most common indication for hospital admission and its incidence is rising. It has a variable prognosis, which is mainly dependent upon the development of persistent organ failure and infected necrotizing pancreatitis. In the past few years, based on large-scale multicenter randomized trials, some novel insights regarding clinical management have emerged. In patients with infected pancreatic necrosis, a step-up approach of percutaneous catheter drainage followed by necrosectomy only when the patient does not improve, reduces new-onset organ failure and prevents the need for necrosectomy in about a third of patients. A randomized pilot study comparing surgical to endoscopic necrosectomy in patients with infected necrotizing pancreatitis showed a striking reduction of the pro-inflammatory response following endoscopic necrosectomy. These promising results have recently been tested in a large multicenter randomized trial whose results are eagerly awaited. Contrary to earlier data from uncontrolled studies, a large multicenter randomized trial comparing early (within 24 h) nasoenteric tube feeding compared with an oral diet after 72 h, did not show that early nasoenteric tube feeding was superior in reducing the rate of infection or death in patients with AP at high risk for complications. Although early ERCP does not have a role in the treatment of predicted mild pancreatitis, except in the case of concomitant cholangitis, it may ameliorate the disease course in patients with predicted severe pancreatitis. Currently, a large-scale randomized study is underway and results are expected in 2017.

Introduction
Acute pancreatitis (AP) is the most common gastrointestinal condition requiring acute hospital admission [1]. In 2009, more than 270,000 patients were diagnosed with AP in the United States and costs are estimated to exceed 2.5 billion dollars every year. Eighty percent of patients developed edematous pancreatitis with severe abdominal pain, which usually subsides within days without development of further complications. Around 20% of patients, however, develop necrotizing pancreatitis of which about a third get infected with mortality rates up to 30% [2, 3]. Unfortunately, no drug therapy is available to ameliorate the disease course, in particular for those who develop a systemic inflammatory response syndrome (SIRS). Prophylactic administration of intravenous antibiotics does not prevent infected pancreatic necrosis. Antibiotics are only indicated in case of a proven infection or in case of a very strong clinical suspicion of either infected necrosis or cholangitis. In the latter case, biliary drainage is mandatory. Optimal feeding has always been regarded as an im-
important means to improve the outcome in this potentially lethal disease. Starting early enteral feeding as soon as possible was thought to be important, but until recently no scientific study was available to support this. Although it is generally accepted that early ERCP has no role in patients with predicted mild biliary pancreatitis, unless there is concomitant cholangitis, there is an ongoing discussion whether in patients with a predicted severe disease course early de-obstruction of the ampulla of Vater and biliary sphincterotomy, with removal of bile duct stones and sludge, is beneficial to prevent complications.

**Treatment of Infected Pancreatic Necrosis**

Approximately, 20% of pancreatitis patients develop necrosis of the pancreatic parenchyma and/or extra-pancreatic fat tissue. Necrosis can be visualized on CT-scan as noncontrast-enhancing regions in and around the pancreas that consist of solid tissue and fluid. As long as these collections remain sterile and asymptomatic, that is, with no pain, gastric outlet obstruction or biliary obstruction, treatment is generally conservative. Between 4 and 6 weeks after disease onset, (peri)pancreatic necrosis may either disappear by resorption or perforation to the digestive tract or become liquefied and a large encapsulated fluid collection develops. In about a third of patients, pancreatic necrosis becomes infected. This is associated with a mortality rate of approximately 30% and is virtually always an indication for invasive treatment. Without intervention, mortality approaches 100% [3–6]. In the earlier days, surgical necrosectomy was performed through a laparotomy, but this carried a very high morbidity rate, as it constitutes a major ‘hit’ to patients with infected necrotizing pancreatitis. There is a ongoing discussion whether in patients with mild pancreatic necrosis, unless there is concomitant cholangitis, there is an ongoing discussion whether in patients with a predicted severe disease course early de-obstruction of the ampulla of Vater and biliary sphincterotomy, with removal of bile duct stones and sludge, is beneficial to prevent complications.

With this concept in mind of minimizing surgical trauma as much as possible, the PANTER trial was initiated within the framework of the Dutch Pancreatitis Study Group [9]. In this multicenter randomized study, 88 patients with necrotizing pancreatitis and with suspected or confirmed infected necrotic tissue were randomized to undergo primary open necrosectomy or a step-up approach treatment. The step-up approach consisted of percutaneous drainage followed, if necessary, by minimally invasive retroperitoneal necrosectomy. The primary end point, a composite of major complications (new-onset multiple-organ failure or multiple systemic complications, perforation of a visceral organ or enterocutaneous fistula, or bleeding) or death, occurred in 69% of patients assigned to open necrosectomy and in 40% of patients assigned to the step-up approach (risk ratio with the step-up approach, 0.57; 95% CI 0.38–0.87; p = 0.006). Importantly, 35% of the patients assigned to the step-up approach did not require surgical necrosectomy and were treated with percutaneous drainage only. New-onset multiple-organ failure occurred significantly less often in patients assigned to the step-up approach (12%) than in those assigned to open necrosectomy (40%). The rate of death did not differ significantly between groups.

After publication of this landmark study pertaining to the management of infected pancreatic necrosis, current guidelines recommend a step-up approach of delayed catheter drainage until the (peri)pancreatic collection has reached the stage of walled-off necrosis, a process that usually takes 4–6 weeks, at which time catheter drainage is followed by necrosectomy when needed.

In order to decrease surgical trauma, video-assisted retroperitoneal debridement was developed. In the latter case, the percutaneous catheter tract is surgically explored by minimal invasive means using a video endoscope, allowing identification and removal of large chunks of necrotic materials [8]. Moreover, endoscopic ultrasonography-guided drainage and necrosectomy techniques have been developed [10]. By means of this technique, the infected collection is identified using a linear ultrasonography endoscope either from the stomach or from the duodenum, and punctured by using a needle or cautery device after which the collection is drained with pigtail plastic stents or a metallic endoprosthesis. In case of ongoing infection and clinical deterioration of the patient, the same cystogastrostomy tract can be dilated up to 20 mm and used as a port d’entree to enter the collection with a regular forward-viewing endoscope, allowing endoscopic removal of necrotic tissue.

Bakker et al. [11] performed on behalf of the Dutch Pancreatitis Study Group a randomized controlled assessor-blinded pilot study (PINGUIN study) in which patients with infected necrotizing pancreatitis were randomized to endoscopic transgastric or surgical necrosectomy. Endoscopic necrosectomy consisted of transgastric puncture, balloon dilatation, retroperitoneal drainage, and necrosectomy. Surgical necrosectomy consisted of video-assisted retroperitoneal debridement or, if not feasible, laparotomy. The premise of this study was that surgical necrosectomy induces a proinflammatory response and is associated with a high complication rate, whereas endoscopic transgastric necrosectomy, a form of natural orifice transluminal endoscopic surgery, may reduce the proinflammatory response and thus reduce complications. The
primary end point in this pilot study was the postproce-
dural proinflammatory response as measured by serum
interleukin 6 (IL-6) levels. Secondary clinical end points
included a predefined composite end point of major com-
plications (new onset multiple organ failure, intra-ab-
dominal bleeding, enterocutaneous fistula, or pancreatic
fistula) or death. In total, 22 patients were randomized, 2
of whom did not undergo necrosectomy following percu-
taneous catheter drainage and therefore could not be ana-
lyzed for the primary end point. Endoscopic transgastric
necrosectomy significantly reduced the postprocedural
IL-6 levels compared with surgical necrosectomy. Twen-
ty-four hours after intervention, IL-6 levels in patient un-
dergoing surgical necrosectomy rose, while in patients un-
dergoing endoscopic necrosectomy, IL-6 levels decreased.
This difference in IL-6 levels between the 2 intervention
groups persisted 1 week after treatment. Although the
study was powered for IL-6 levels as a proxy for disease se-
verity, and was not adequately powered to definitely an-
swer the secondary endpoints, the differences were strik-
ing. The composite endpoint of major complications and
death occurred in 20% of patients treated by means of en-
doscopy compared to 80% in patients treated with sur-
gery. This difference was statistically significant and obvi-
ously carries an important clinical relevance. The same
pattern emerged for new onset organ failure; 0% after en-
doscopic treatment vs. 50% after surgical treatment.

The remarkable outcome of this pilot study formed the
basis of the design of a large-scale, adequately powered,
randomized, controlled, parallel-group superiority mul-
ticenter trial (TENSION trial) to answer the question
whether in patients with (suspected) infected necrotizing
pancreatitis with an indication for intervention and in
whom both treatment modalities are deemed possible,
endoscopic transluminal step-up treatment is superior to
a surgical step-up approach. Patient inclusion of this
study has been completed and the data are currently be-
ing analyzed. Over a 4-year time period, 98 patients have
been enrolled from 24 hospitals of the Dutch Pancreatitis
Study Group. The primary endpoint is a composite of
death and major complications within 6 months follow-
ing randomization. Secondary endpoints include complica-
tions such as pancreaticocutaneous fistula, exocrine or
endocrine pancreatic insufficiency, need for additional
radiological, endoscopic or surgical intervention, the
need for necrosectomy after drainage, the number of (re-)in-
terventions, quality of life, and total direct and indirect
costs. Soon, the TENSION trial will provide a conclusive
answer whether an endoscopic step-up approach reduces
the combined primary endpoint of death and major com-
plications, as well as hospital stay and related costs com-
pared with a surgical step-up approach in patients with
infected necrotizing pancreatitis.

Since the step-up approach is now considered stan-
dard practice [2] and the results of the TENSION trial will
soon reveal which strategy is to be preferred, the question
of optimal timing of drainage has become the next focal
point of discussion both in terms of safety and cost-effec-
tiveness. Current literature shows that 35–64% of patients
with INP can be treated with delayed catheter drainage,
without the need for invasive necrosectomy [9, 13]. In a
recent international survey, although not evidence based,
45% of expert pancreatologists proclaimed that they im-
mediately proceed with catheter drainage once infected
pancreatic necrosis has been established [12]. Indeed,
various case series suggest that encapsulation is not a ‘sine
qua non’ for catheter drainage to be successful [13].

Therefore, optimal timing of drainage is the next re-
search question to be tackled by the Dutch Pancreatitis
Study Group. For this, the POINTER study has been de-
signed of which patient inclusion has already started. This
study investigates whether a strategy of early diagnosis of
infected pancreatic necrosis in combination with imme-
diate catheter drainage is more beneficial with regard to
clinical outcome and cost effectiveness than the current
standard of postponed catheter drainage until walled-off
necrosis has developed. With the current strategy of de-
layed intervention, patients with infected pancreatic ne-
crosis are clinically observed for long periods of time
awaiting resolution of necrosis. Consequently, the utiliza-
tion of health care resources is high with direct medical
costs exceeding €80,000 ($110,000) [9]. However, 35–
64% of patients with infected pancreatic necrosis that are
treated with postponed catheter drainage do not require
necrosectomy [9, 13]. If such a high percentage of patients
respond so favorably to delayed catheter drainage, it is
only logical to hypothesize that earlier catheter drainage
may be even more beneficial in terms of preventing seri-
ous complications to occur including organ failure and
sepsis and the need for necrosectomy.

With this new concept of very early drainage, the meth-
oodology of how to diagnose infected pancreatic necrosis
needs to be re-evaluated. With the strategy of postponed
drainage, the clinical features of either gas in the necrotic
collection on cross-sectional imaging or ongoing organ fail-
ure for several weeks after the onset of AP and a failure to
thrive, enabled a correct diagnosis of infection of the necro-
sis in 91% of patients [9]. However, with regard to setting
the correct indication for very early drainage, it will prove
impossible to distinguish between the SIRS and infected

DOI: 10.1159/000445257
pancreatic necrosis on clinical grounds. In order to make a correct diagnosis in these very early stages, additional diagnostic procedures like fine needle aspiration (FNA) are required to confirm or exclude infection in the first 2 weeks after disease onset. In a recent study, infected pancreatic necrosis was confirmed by FNA in 86% of patients, which is equal to the diagnostic performance of clinical symptoms (80%) or gas bubbles on CT-scan (94%) [14].

Feeding in Pancreatitis

Feeding in AP has always been an area of major interest to prevent secondary infection of necrosis and improve patient outcome. It is hypothesized that infection of pancreatic necrosis is mediated by bacterial translocation from the gut, provoked by disturbed intestinal motility, bacterial overgrowth, and increased mucosal permeability [15–19]. Nasoenteric tube feeding is believed to stimulate intestinal motility, which may reduce bacterial overgrowth and preserve gut mucosal integrity through the stimulation of splanchnic blood flow [20, 21]. A comprehensive meta-analysis involving 8 randomized trials with a total of 348 patients showed that nasoenteric tube feeding compared with total parenteral nutrition reduced the rate of infections and mortality among patients with severe pancreatitis [22]. A meta-analysis of randomized trials involving acutely ill patients, but no pancreatitis, showed a 22% reduction in the rate of major infections when nasoenteric tube feeding was started ≤ 36 h as compared with a later start [23]. In patients with AP, nonrandomized studies have shown that nasoenteric tube feeding started within 48 h after admission, as compared with a start after 48 h, significantly reduced the rate of major infection and in some studies even reduced mortality [24–27]. Importantly, the use of an enteric feeding tube not only has presumed benefits. Many patients dislike having a feeding tube because it may cause substantial discomfort including throat pain and excessive gagging. Moreover, feeding tubes may dislodge due to gagging or become obstructed due to impacted feeding fluid necessitating frequent replacement of the tube. A more restricted use of enteral nutrition and a feeding tube only in those patients who really benefit from it, would result in substantial avoidance of discomfort and costs.

Recently, a large multicenter randomized controlled study from the Dutch Pancreatitis Study Group did not show that in patients with AP at high risk for complications, an early start of nasoenteric tube feeding is superior compared to the introduction of an oral diet after 72 h (with tube feeding only on-demand) [28]. There was no statistically significant difference with regard to the composite end point of major infection or death or for the secondary endpoints including development of necrotizing pancreatitis, infected pancreatic necrosis, bacteremia, CT severity index, ICU admission, and organ failure. With the oral diet and on-demand tube feeding strategy, only approximately one third of patients required a nasojejunal feeding tube. It is difficult to speculate on factors that explain the observed differences with previous trials and observational studies, which suggested an improved outcome after early nasoenteric tube feeding as compared with total parenteral nutrition, other than study design and methodology [8, 10–13]. The timing of early nasoenteric tube feeding was similar to the timing mentioned in previous studies and similar criteria for enrolling patients at high risk for complications were used. One of the major issues with observational studies is that one cannot differentiate between cause and effect; for instance, less severely ill patients may have been fed earlier. This discussion on (early) feeding to ameliorate the disease course is not limited to patients with pancreatitis. In most current ICU guidelines, early enteral feeding is recommended, but the methodologic quality of the included trials has been criticized [29–31]. American and European nutritional societies recommend routine early nasoenteric tube feeding in all patients with severe pancreatitis [32–34]; guidelines from gastroenterologic and pancreatic societies, however, state that, regardless of disease severity, tube feeding is indicated when patients are predicted not to be able to tolerate an oral diet for up to 7 days [35, 36]. Unfortunately, it takes several days after admission to make such assessment and by that time the window of opportunity for effective prevention of infection with early tube feeding may have passed. Based on the trial by Bakker et al. [11], it is now right to state that current evidence does not support an early start of nasoenteric tube feeding in all patients with severe AP in order to reduce the risks of infection and death [14].

ERCP in Biliary Pancreatitis

In a majority of cases, AP is initiated by the obstruction of gallstones at the level of the ampulla of Vater [36, 37]. Entrapment of biliary stones or sludge is believed not only to initiate an attack of pancreatitis but also to sustain and possibly aggravate the disease. This is the reason why (early) biliary decompression by means of ERC with sphincterotomy was proposed as a means to ameliorate the disease course [38, 39]. The presumed benefits of such
intervention have to be weighed against potential complications that are associated with ERC and sphincterotomy such as pancreatitis, bleeding and perforation [40, 41].

Several studies on the indication and outcome of ERC in patients with biliary pancreatitis have been published [42]. Guidelines declare that there is an undisputed indication for ERC in patients with concomitant cholangitis [2, 36]. In patients with a predicted mild disease course, early ERC is not indicated, as in these patients the risk for complications does not outweigh the potential benefit. For patients with pancreatitis with concomitant cholestasis, guidelines suggest that ERC with sphincterotomy be performed, but the quality of evidence for this recommendation is moderate at best. In patients with a biliary pancreatitis with a predicted severe disease course but without concomitant cholangitis, data are conflicting. A meta-analysis did not show a beneficial effect of routine early ERC compared to conservative treatment [42]. Unfortunately, each of the included studies has notable shortcomings that prevent us from drawing a reliable conclusion from the aggregated data, in particular for patients with predicted severe biliary pancreatitis but without cholangitis [43]. For one, patients with a low pre-likelihood of a biliary etiology were included, as well as patients at low risk for developing complications, and patients with cholangitis at presentation. Second, patient selection criteria and study endpoints varied considerably between studies. Third, patients with and without cholestasis are not presented separately, and this precludes doing an important subgroup analysis. Fourth, routine ‘early’ ERC was not performed within 24 h after hospital admission but after 48–72 h, and this may be too late to prevent complications from severe disease or increase the risk for ERC related complications. Fifth, study protocols did not specify if and when sphincterotomy should be performed. This resulted in many patients who underwent ERC without sphincterotomy. It is our opinion that sphincterotomy should be performed routinely in order to decompress the biliary duct, even in the absence of visible gallstones or sludge during ERC [44–46]. Sixth, no criteria were set to guarantee that ERCs were performed by experienced endoscopists, although ERC is an intervention that requires considerable training and expertise [47, 48]. Finally, even if the data of all randomized trials are pooled, such analysis still has insufficient power to detect statistically significant effects of early ERC with sphincterotomy in patients with severe biliary pancreatitis without concomitant cholangitis with respect to major complications or death [42, 49]. The Dutch Pancreatitis Study Group is currently conducting a multicenter randomized study in 232 patients to investigate whether early ERC with sphincterotomy compared to conservative treatment improves outcome in patients with biliary pancreatitis that are at high risk for complications but do not have cholangitis. Patient inclusion is already more than halfway and results are awaited in 2017.

Disclosure Statement

M.J.B. was a consultant and lecturer to Boston Scientific and Cook Medical and consultant to SOCAR and Uniqure.

References


