Comparison of clinical course and outcome of acute pancreatitis according to the two main etiologies: alcohol and gallstone

Joon Hyun Cho, Tae Nyeun Kim* and Sung Bum Kim

Abstract

Background: Studies concerning clinical course and outcome of acute pancreatitis (AP) according to etiologies were rare, especially after year 2000. This study was designed to investigate the difference between the clinical course of alcoholic and biliary AP.

Methods: Of the 153 patients diagnosed as AP with a first attack between January 2011 and January 2013, extensive clinical data of 50 patients with AP caused by alcohol and 76 patients with AP caused by gallstone were analyzed retrospectively. We compared the severity of AP defined by revised Atlanta classification in 2012, local complications, severity scores, and computed tomography severity index (CTSI) between alcoholic and biliary AP. We also evaluated the length of hospital stay, duration of NPO, and in-hospital mortality in each group.

Results: Hemoglobin, hematocrit, and serum C-reactive protein level measured after admission for 24 h were significantly higher in the alcohol group than in the biliary group. Incidence of pseudocyst formation was significantly higher in the alcohol group than in the biliary group (20.0 % vs. 6.6 %, \( P = 0.023 \)). Among prognostic scoring systems, only CTSI showed significant difference (\( P < 0.001 \)) with a mean score of 3.0 ± 0.9 in the alcohol group and 1.7 ± 1.2 in the biliary group. Severe AP with organ failure persisting beyond 48 h was observed in 12 patients (24.0 %) in the alcohol group and one patient (1.3 %) in the biliary group (\( P < 0.001 \)). There were 4 mortalities in the alcohol group only (\( P = 0.012 \)).

Conclusion: More severe forms of AP and local complication, such as pseudocyst formation, are associated with alcoholic AP compared with biliary AP.

Keywords: Acute pancreatitis, Etiology, Alcohol, Gallstone, Clinical course

Background

Acute pancreatitis (AP) is a relatively common disease with incidence of 5 – 80 per 100,000 members of the population, although its prevalence varies in different countries and even in different areas of a given country, and the number of new cases has shown a steady increase in recent years [1–3]. Most patients with AP show spontaneous resolution without complications, however, 10 %-20 % of patients experience a severe attack with increased risk of mortality up to 25 % [4].

Gallstones and alcohol account for 70 %-80 % of all AP etiologies, although AP may be the consequence of numerous etiologic factors [5]. Identification and differentiation of etiologies of AP is important, because it can affect the specific therapeutic strategies, and eliminating causes of AP prevents further aggravation and recurrence of AP. Although many theories concerning the pathogenesis of AP caused by alcohol and gallstone have been proposed, no theory regarding the mechanisms for inducing inflammation of the pancreas in these two conditions has been widely accepted. However, the mechanisms of induction of AP by gallstones and alcohol appear to be different [6], which could lead to differences in clinical severity and rate of complications. Nevertheless, the question of whether the etiology is a
risk factor for prognosis of AP remains controversial. There have been differing reports with regard to whether etiologic factors affect clinical outcome and mortality of AP. Various studies have reported that biliary AP is more severe and is associated with higher mortality than alcoholic AP [7–9]. A few investigators have reported higher complication and mortality in patients with alcoholic AP [10, 11], while some studies found no significant difference [12–14]. Most of these studies were reported before year 2000. Atlanta Classification [15] was revised in 2012, with an emphasis on persistent organ failure. In recent years, many advances, such as endoscopic sphincterotomy and radiologic intervention, have been made in treatment of AP. These might have changed clinical course and outcome of AP from the past.

This study was designed to investigate the role of the two main etiological factors, alcohol and gallstone, on the clinical course and outcome of AP.

Methods

Patients

Between January 2011 and January 2013, a total of 153 patients were diagnosed as AP with a first attack, and, among these patients, 126 patients with AP induced by two major causes, alcohol or gallstone, were included in this study. Extensive demographic, radiographic, and laboratory data of these consecutive patients were prospectively collected and analyzed retrospectively. Patients with AP on pre-existing chronic pancreatitis and previous history of AP were excluded from this study. This study was approved by the Institutional Review Board of Yeungnam University Hospital. Written informed consent was obtained from all patients.

All patients diagnosed as AP were treated with supportive care, including NPO, fluid resuscitation, and analgesics. Endoscopic retrograde cholangiopancreatography (ERCP) was considered in patients with biliary pancreatitis, and the timing of ERCP was dependent upon severity and clinical course of the patients. ERCP was performed as soon as possible in patients with concomitant acute cholangitis. We awaited spontaneous improvement of biliary obstruction for 24 – 48 hrs in patients with no associated cholangitis. Oral feeding was permitted when abdominal pain subsided and patients felt hunger sensation. Patients who remained asymptomatic with oral intake were discharged or underwent cholecystectomy if indicated.

Definitions

AP was diagnosed if a patient had more than two of the three following findings: typical abdominal pain of AP (acute onset of a persistent and severe epigastric pain often radiating to the back), elevation of serum amylase and/or lipase levels above three times upper normal limit, and findings of imaging studies, including abdominal ultrasonography or computed tomography (CT), consistent with AP [15]. Alcohol was considered a cause of AP when patients had a history of alcohol consumption within 48 h before symptom onset and other possible causes were ruled out. Biliary AP was defined when a gallstone or biliary sludge was observed on abdominal ultrasonography or CT. The etiology was considered idiopathic when causative factors could not be identified from a detailed clinical and drug history or after initial investigations.

Severity of AP was determined according to the most recently revised Atlanta Classification. Mild AP was defined by the absence of organ failure and the absence of local or systemic complications. Moderately severe AP was defined by the presence of transient organ failure, local complications such as acute peripancreatic fluid collections (APFC) and acute necrotic collections (ANC), or exacerbation of co-morbid diseases. Severe AP was defined by persistent organ failure for more than 48 h. Organ failure was defined as a score of 2 or more for one of the three systems (respiratory, cardiovascular, and renal) using the modified Marshall scoring system [16].

Pancreatic pseudocyst was defined as an encapsulated fluid collection usually outside the pancreas with a well defined wall and homogeneous fluid density occurs more than 4 weeks after the onset of interstitial oedematous pancreatitis. Walled-off necrosis (WON) was defined as a mature and encapsulated collection of pancreatic and/or peripancreatic necrosis with a well defined wall and heterogeneous with liquid and non-liquid density, with varying degrees of loculations, occurs more than 4 weeks after the onset of necrotizing pancreatitis [15].

Data collection

The following parameters were collected for each episode of AP: length of hospital stay, duration of NPO, inhospital mortality, presence of organ failure, and local complications such as APFC, pseudocyst, ANC, and WON. Acute Physiology and Chronic Health Evaluation (APACHE)-II [17] and bedside index for severity in acute pancreatitis (BISAP) [18, 19] scores were calculated using data from the first 24 h after admission and the Ranson [20] score using data from the first 48 h. CT severity index (CTSI) [21] was calculated in patients who underwent contrast-enhanced CT (CECT) within 48 h after admission.

Statistics

Data were collected in a Microsoft Excel database. After completion of data collection, the database was imported into SPSS for Windows (20.0, SPSS, Chicago, IL, USA). Continuous variables were expressed as mean with standard deviation (SD) and compared using Student’s t-
test or Mann–Whitney test. Categorical variables were expressed as absolute numbers and proportions. Pearson’s $\chi^2$ test or Fisher’s exact test was used for comparison of categorical variables. A $P$ value of $<0.05$ was considered statistically significant.

**Results**

**Etiology of AP**

In a two-year time period, 153 patients diagnosed as AP with a first attack were admitted to our institution. The etiologies of AP were gallstone in 76 patients (49.7%), alcohol in 50 patients (32.7%), idiopathic in 24 patients (15.7%), and malignancy in two patients, and hypertriglyceridemia in one patient.

**Patient characteristics of alcoholic and biliary AP**

Mean age of 126 patients with AP caused by two major causes, alcohol or gallstone, was $63.6 \pm 15.4$ years (range, 27–94) and 81 patients (64.3%) were male (Table 1). Age and BMI did not differ significantly between the alcohol group and the biliary group ($P=0.104$, and $P=0.798$, respectively). Male was more predominant in the alcohol group than in the biliary group, with statistical significance ($P<0.001$). Medical comorbidity did not differ significantly between the two groups. Initial laboratory findings of hemoglobin and hematocrit were significantly higher in patients with alcoholic AP compared to those with biliary AP ($P=0.002$, and $P=0.011$, respectively). Serum alanine aminotransferase level was significantly higher in the biliary group than in the alcohol group ($P<0.001$). Serum C-reactive protein (CRP) level measured at admission (CRPi) did not differ significantly between the two groups, and serum CRP level measured after admission for 24 h (CRP$_{24}$) was significantly higher in the alcohol group compared with the biliary group ($P<0.001$). Serum amylase and lipase level were significantly higher in patients with biliary AP compared to those with alcoholic AP ($P<0.001$, and $P=0.004$, respectively). We compared serum pancreatic enzymes between two groups except two patients with renal failure in the biliary group.

**Comparison of clinical course, outcome, and severity of alcoholic and biliary AP**

Of the 50 patients with alcoholic AP, 12 patients (24.0%) developed persistent organ failure for more

| Table 1 Baseline demographics and clinical characteristics of the patients |
|---------------------------|--------------------------|-----------------|
|                           | Alcohol group (N = 50)   | Biliary group (N = 76) | $P$ value |
| Age, years                | 52.3 ± 11.2              | 67.3 ± 14.9        | 0.104     |
| Mean ± SD                 |                          |                  |
| Median (range)            | 51 (31–80)               | 71 (27–94)        |           |
| Male                      | 41 (82.0)                | 40 (52.6)         | <0.001    |
| Male (%)                  | 82.0                     | 52.6              |           |
| BMI (kg/m$^2$)            | 22.7 ± 3.5               | 23.1 ± 3.5        | 0.798     |
| Medical comorbidity       |                          |                  |
| Hypertension              | 8 (16.0)                 | 21 (27.6)         | 0.129     |
| Liver cirrhosis           | 2 (4.0)                  | 1 (1.3)           | 0.334     |
| Diabetes mellitus         | 6 (12.0)                 | 9 (11.8)          | 0.978     |
| Chronic kidney disease    | 0 (0)                    | 2 (2.6)           | 0.413     |
| Ischemic heart disease    | 2 (4.0)                  | 4 (5.3)           | 0.744     |
| Laboratory findings       |                          |                  |
| Hemoglobin (g/dL)         | 14.7 ± 3.1               | 13.2 ± 1.8        | 0.002     |
| Hematocrit (%)            | 42.2 ± 7.6               | 38.5 ± 5.6        | 0.011     |
| AST (IU/L)                | 229.4 ± 438.9            | 284.2 ± 317.1     | 0.499     |
| ALT (IU/L)                | 98.5 ± 119.4             | 234.4 ± 226.3     | <0.001    |
| Creatinine (mg/dL)        | 1.2 ± 0.8                | 1.2 ± 0.8         | 0.758     |
| LDH (IU/L)                | 973.6 ± 860.6            | 737.5 ± 425.6     | 0.078     |
| CRPi (mg/dL)              | 8.6 ± 11.8               | 4.5 ± 6.3         | 0.106     |
| CRP$_{24}$ (mg/dL)        | 20.9 ± 10.1              | 7.7 ± 6.8         | <0.001    |
| Amylase (IU/L)            | 738.6 ± 648.4            | 2061.1 ± 1479.0   | <0.001    |
| Lipase (IU/L)             | 1090.7 ± 1036.8          | 2456.9 ± 2220.7   | 0.004     |

Values are expressed as the mean ± SD or n (%)

BMI, body mass index; AST, Aspartate aminotransferase; ALT, Alanine aminotransferase; LDH, Lactate dehydrogenase; CRPi, C-reactive protein measured at admission; CRP$_{24}$, C-reactive protein measured after admission for 24 h
than 48 h and were classified as severe AP according to the Atlanta Classification, and of the 76 patients with biliary AP, only one patient was classified as severe AP (Table 2). The number of patients with severe AP was significantly higher in the alcohol group than in the biliary group ($P < 0.001$). Thirty patients (60.0 %) in the alcohol group and 16 patients (21.1 %) in the biliary group developed moderately severe AP, respectively. Patients with moderately severe AP were more predominant in the alcohol group than in the biliary group, with statistical significance ($P < 0.001$). Eight patients (16.0 %) in the alcohol group and 59 patients (77.6 %) in the biliary group developed mild AP, respectively. The number of patients with mild AP was significantly higher in the biliary group than in the alcohol group ($P < 0.001$).

APFCs were observed on CECT in 15 of 50 patients (30.0 %) in the alcohol group and in 14 of 76 patients (18.4 %) in the biliary group (Table 2). In the early days of hospitalization, pancreatic and peripancreatic necrosis with ANC was noted in 3 patients (6 %) in the alcohol group and 3 patients (3.9 %) in the biliary group. Among these patients, 1 patient in alcohol group developed WON after 4 weeks from the onset of AP during the follow-up period. No statistical differences in occurrence of APFC and ANC were observed between patients with alcoholic AP and those with biliary AP ($P = 0.131$ and, $P = 0.597$, respectively). Pseudocyst occurred significantly more often in patients with alcoholic AP (10/50, 20 %) than in those with biliary AP (5/76, 6.6 %) ($P = 0.023$).

No significant differences in Ranson, BISAP, and APACHE-II scores were observed between the alcohol group and the biliary group (9.9 ± 5.2 vs. 7.9 ± 3.5 ($P = 0.086$), 5.4 ± 2.7 vs. 4.4 ± 2.0 ($P = 0.068$), and 3.1 ± 2.3 vs. 2.8 ± 1.7 ($P = 0.539$), respectively) (Table 3). Four mortalities (8.0 %) occurred during hospitalization in the alcohol group; No patient died in the biliary group.

**Table 2** Severity and local complications of alcoholic and biliary AP

| Severity of AP | Alcohol group (n = 50) | Biliary group (n = 76) | $P$ value |
|----------------|------------------------|-----------------------|-----------|
| Mild           | 8 (16)                 | 59 (77.6)             | <0.001    |
| Moderately severe | 30 (60)             | 16 (21.1)             | <0.001    |
| Severe         | 12 (24)                | 1 (1.3)               | <0.001    |

| Local complications | Alcohol group (n = 50) | Biliary group (n = 76) | $P$ value |
|---------------------|------------------------|-----------------------|-----------|
| APFC                | 15 (30)                | 14 (18.4)             | 0.131     |
| Pseudocyst          | 10 (20)                | 5 (6.6)               | 0.023     |
| ANC                 | 3 (6)                  | 3 (3.9)               | 0.597     |
| WON                 | 1 (2)                  | 0 (0)                 | 0.216     |

Values are expressed as the n (%)  
AP, acute pancreatitis; APFC, acute peripancreatic fluid collection; ANC, acute necrotic collection; WON, walled-off necrosis

between the two groups was only noted in CTSI, with statistical significance ($P < 0.001$).

No significant differences in hospital stay, duration of NPO, and the time to alleviation of symptoms were observed between the alcohol group and the biliary group (9.9 ± 5.2 vs. 7.9 ± 3.5 ($P = 0.086$), 5.4 ± 2.7 vs. 4.4 ± 2.0 ($P = 0.068$), and 3.1 ± 2.3 vs. 2.8 ± 1.7 ($P = 0.539$), respectively) (Table 4). Four mortalities (8.0 %) occurred during hospitalization in the alcohol group; No patient died in the biliary group.

**Table 3** Severity scores and CTSI in alcoholic and biliary AP

|               | Alcohol group (n = 50) | Biliary group (n = 76) | $P$ value |
|---------------|------------------------|-----------------------|-----------|
| Ranson score  | 3.1 ± 1.8              | 3.1 ± 1.3             | 0.861     |
| BISAP score   | 1.2 ± 1.0              | 1.2 ± 0.9             | 0.888     |
| APACHE-II score | 8.0 ± 5.7             | 6.8 ± 3.4             | 0.322     |
| CTSI          | 3.0 ± 0.9              | 1.7 ± 1.2             | <0.001    |

Values are expressed as the mean ± SD  
AP, acute pancreatitis; BISAP, bedside index of severity in acute pancreatitis; APACHE-II, Acute Physiology and Chronic Health Evaluation-II; CTSI, computed tomography severity index

**Table 4** Clinical course and outcome in alcoholic and biliary AP

|               | Alcohol group (n = 50) | Biliary group (n = 76) | $P$ value |
|---------------|------------------------|-----------------------|-----------|
| Hospital stay, days | 9.9 ± 5.2              | 7.9 ± 3.5             | 0.086     |
| NPO, days     | 5.4 ± 2.7              | 4.4 ± 2.0             | 0.068     |
| Time to alleviation of symptoms, days | 3.1 ± 2.3 | 2.8 ± 1.7 | 0.539     |
| Mortality     | 4 (8.0)                | 0 (0)                 | 0.012     |

Values are expressed as the mean ± SD or n (%)  
AP, acute pancreatitis; NPO, nil per os

**Discussion**

AP has many causes and the proportion of each etiologic factor varies greatly in different countries. In general, the most frequent cause of AP is cholelithiasis, followed by alcohol. Together these etiologies are responsible for approximately 80 % of all episodes of AP [5]. In a previous study including 1,068 European patients with AP, the percentage of biliary AP varied between 24 and 71 %, and that of alcoholic AP between 6
Some investigators have reported that patients et al. BMC Gastroenterology 2015). In this study, the CRP parameters for early prediction of necrotizing pancreatitis (LDH), and hematocrit have been reported to be reliable Serum markers such as CRP, lactate dehydrogenase (LDH), and hematocrit have been reported to be reliable parameters for early prediction of necrotizing pancreatitis. In this study, the CRP, lactate dehydrogenase (LDH), and hematocrit were significantly higher in patients with alcoholic AP compared to biliary AP. Serum LDH level was also higher in the alcohol group than in the biliary group, although this was not statistically significant. These differences of laboratory findings may indicate that patients with alcoholic AP have a more severe clinical course and poorer prognosis compared to those with biliary AP. Our study showed that moderately severe and severe AP defined according to revised Atlanta Classification and the mortality were significantly more common in the alcohol group than in the biliary group, and the number of patients with mild AP was significantly lower in the alcohol group compared with the biliary group, although no differences were observed with regard to the scoring systems predicting the severity of AP, such as Ranson, BISAP, and APACHE-II scores. These results indicated that there was no difference in the severity of AP on admission between alcoholic and biliary AP, but the patients with alcoholic AP were more likely to become severe during the clinical course compared with biliary AP. In our study, duration of hospital stay and NPO tended to be longer in the alcohol group than in the biliary group. Lower proportion of severe AP in the biliary group can be explained in part by the fact that progression of AP was likely to be prevented owing to early treatment of bile duct stone by ERCP. In general, many patients with biliary AP experience spontaneous passage of stone and ERCP is not recommended in these patients, while endoscopic removal of bile duct stone with ERCP is recommended if it does not pass spontaneously. The severity, clinical course and prognosis of biliary AP could be associated with concomitant acute cholangitis and cholestasis. In patients with acute cholangitis and ongoing biliary obstruction, urgent ERCP to remove bile duct stones may lessen the severity of biliary AP. In our studies, 45 of 76 patients with biliary AP showed spontaneous improvement without ERCP, in whom spontaneous passage of CBD stone was suspected. Among 31 patients who underwent ERCP, 12 patients underwent elective ERCP after improvement of AP. Urgent (<24 h) or early (<72 h) ERCP was performed in 19 patients with poor response to medical therapy, and rapid improvement of pancreatitis was observed after endoscopic stone removal in these patients. Early recognition and removal of CBD stone by ERCP or spontaneous passage of the stones might have a beneficial influence on the clinical course of biliary AP. These timely interventions might contribute to change in clinical course and improvement of outcome in biliary AP, whereas patients with alcoholic AP could be treated only by supportive care such as NPO, aggressive fluid resuscitation, and close monitoring. In the previous report, a tendency to have more peripancreatic infiltrations or fluid collections was observed for alcoholic AP compared with biliary AP. In our study, frequency of pseudocyst was significantly higher in the alcohol group than in the biliary group, and CTSI in the alcohol group was significantly higher compared with that in the biliary group. Alcoholic AP usually occurs in patients with heavy alcohol intake for a long period of time, although single heavy drinking could lead to AP, and substantial damage to the pancreas has already occurred when AP occurs. Underlying pancreatic ductal stricture and dilation due to longstanding pancreatic damage by alcohol are presumably present in patients with alcoholic AP; and, thus, they are more vulnerable to ductal disruption with subsequent leakage of pancreatic juice into the peripancreatic area and pseudocyst formation. On the other hand, biliary AP usually occurs from a single episode of bile duct obstruction caused by gallstone in relatively intact pancreas. This difference in duration of exposure to harmful impact may contribute to difference in disease severity between alcoholic and biliary AP. Lower incidence of walled-off necrosis in this study may be explained in part by lower incidence of pancreatic and/or peripancreatic necrosis at the time of admission. The reason of lower occurrence of pancreatic
necrosis in this study was unclear, but multiple factors such as BMI, genetic and environmental factors might be associated.

There were some limitations in this study. First, although all patients in this study were enrolled prospectively over the study period, some clinical data, especially for the transferred patients, were missing due to lack of availability because the data of these patients were analyzed retrospectively. Second, the number of including patients was relatively small compared to other large scale clinical studies; therefore, comparison of some variables between two groups and further analysis was difficult. Nevertheless, this study was one of the few studies to compare clinical course and outcomes of AP caused by alcohol and gallstone especially in a non-Western area.

Conclusion

In summary, results of our study demonstrated that more severe forms of AP and local complication, such as pseudocyst development, are associated with alcoholic AP compared with biliary AP. The timely interventions such as endoscopic sphincterotomy and pathophysiologic difference may contribute to difference in disease severity between alcoholic and biliary AP.

Abbreviations

AP: Acute pancreatitis; ERCP: Endoscopic retrograde cholangiopancreatography; CT: computed tomography; APFC: acute peripancreatic fluid collection; ANC: acute necrotic collection; WON: walled-off necrosis; APACHE: Acute Physiology and Chronic Health Evaluation; BISAP: bedside index for severity in acute pancreatitis; CTSI: Computed tomography severity index; CECT: contrast-enhanced computed tomography; SD: standard deviation; CRP: C-reactive protein; CRP: C-reactive protein measured at admission; CRPME: C-reactive protein measured after admission for 24 h; CBD: common bile duct; LDH: lactate dehydrogenase.

Competing interests

The authors declare that they have no competing interests.

Authors’ contributions

TN proposed the study. TN and JH designed the study. JH and SB performed research and wrote the first draft manuscript. SB and JH collected and analyzed the data. TN and JH reviewed the draft manuscript and provided feedback. All authors read and approved the final manuscript.

Acknowledgements

This work was supported by the 2014 Yeungnam University Research Grant. The authors would like to acknowledge the work of Hyun Hee Chung, MD (Division of Gastroenterology and Hepatology, Department of Internal Medicine, Yeungnam University College of Medicine) in acquisition of data and Nurisco’s editor for proof-reading the manuscript for English grammar and style.

Received: 10 February 2015 Accepted: 17 July 2015
Published online: 25 July 2015

References

1. Peery AF, Dellon ES, Lund J, Crockett SD, McGowan CE, Bulsiewicz WJ, et al. Burden of gastrointestinal disease in the United States: 2012 update. Gastroenterology. 2012;143:1179–87.

2. Fagenholz PJ, Fernandez-del Castillo C, Harris NS, Pelletier AJ, Camargo Jr CA. Direct medical costs of acute pancreatitis hospitalizations in the United States. Pancreas. 2007;35:302–7.

3. Corfield AP, Cooper MJ, Williamson RC. Acute pancreatitis: a lethal disease of increasing incidence. Gut. 1985;26:224–9.

4. Yeung VP, Lam BY, Yip AW. APACHE system is better than Ranson system in the prediction of severity of acute pancreatitis. Hepatobiliary Pancreat Dis Int. 2006;5:294–9.

5. Thomson SR, Hendry WS, McFarlane GA, Davidson AI. Epidemiology and outcome of acute pancreatitis. Br J Surg. 1987;74:398–401.

6. Sakorafas GH, Tsitou AG. Etiology and pathogenesis of acute pancreatitis: current concepts. J Clin Gastroenterol. 2000;30:343–56.

7. Imrie CW, Whyte AS. A prospective study of acute pancreatitis. Br J Surg. 1975;62:490–4.

8. Ranson JH, Rifkind KM, Roses DF, Fink SD, Eng K, Spencer FC. Prognostic signs and the role of operative management in acute pancreatitis. Surg Gynecol Obstet. 1974;139:68–81.

9. Frey CF. Gallstone pancreatitis. Surg Clin North Am. 1981;61:923–8.

10. Bank S, Wise L, Gersten M. Risk factors in acute pancreatitis. Am J Gastroenterol. 1983;78:67–40.

11. Agarwal N, Pitchumoni CS. Assessment of severity in acute pancreatitis. Am J Gastroenterol. 1991;86:1385–91.

12. Uhl W, Issenmann R, Curti G, Vogel R, Beger HG, Büchler MW. Influence of etiology on the course and outcome of acute pancreatitis. Pancreas. 1996;13:335–43.

13. Gullo L, Migliori M, Olah A, farkas G, Levy P, Arvanitakis C, et al. Acute pancreatitis in five European countries: etiology and mortality. Pancreas. 2002;24:223–7.

14. Andersen AM, Novovic S, Enbol AK, Hansen MB. Mortality in Alcohol and Biliary Acute Pancreatitis. Pancreas. 2008;36:432–4.

15. Acute Pancreatitis Classification Working Group. Classification of acute pancreatitis–2012: revision of the Atlanta classification and definitions by international consensus. Gastroenterology. 2013;145:114–5.

16. Marshall JC, Cook DJ, Chistou NV, Bernard GR, Sprung CL, Sibbald WJ. Multiple organ dysfunction score: a reliable descriptor of a complex clinical outcome. Crit Care Med. 1995;23:1638–92.

17. Larvin M, McMahon MJ. APACHE-II score for assessment and monitoring of severe acute pancreatitis. Lancet. 1989;2:201–5.

18. Wu BJ, Johannès RS, Sun X, Tabak Y, Conway DL, Banks PA. The early prediction of mortality in acute pancreatitis: a large population-based study. Gut. 2008;57:1698–703.

19. Singh VK, Wu BJ, Bollen TL, Repas K, Maurer R, Johannes RS, et al. A prospective evaluation of the bedside index for severity in acute pancreatitis score in assessing mortality and intermediate markers of severity in acute pancreatitis. Am J Gastroenterol. 2009;104:966–71.

20. Ranson JH, Rifkind KM, Roses DF, Fink SD, Eng K, Localio SA. Objective early identification of severe acute pancreatitis. Am J Gastroenterol. 1974;61:443–51.

21. Balthazar EJ, Robinson DL, Megibow AJ, Ranson JH. Hemococoncentration is an early marker for organ failure and necrotizing pancreatitis. Pancreas. 1993;5:399–401.

22. Sperpoint E, Berka A, Blasi A, Brenchley MJ, Cloonan F, Desai N, et al. Early ductal decompression prevents the progression of biliary pancreatitis: an experimental study in the opossum. Gastroenterology. 1993;105:157–64.

23. Fan ST, Lai EC, Mok FP, Lo CM, Zheng SS, Wong J. Early treatment of acute biliary pancreatitis by endoscopic papillotomy. N Engl J Med. 1993;328:228–32.
29. Dutch Acute Pancreatitis Study Group. Early endoscopic retrograde cholangiopancreatography in predicted severe acute biliary pancreatitis: a prospective multicenter study. Ann Surg. 2009;250:68–75.

30. Tse F, Yuan Y. Early routine endoscopic retrograde cholangiopancreatography strategy versus early conservative management strategy in acute gallstone pancreatitis. Cochrane Database Syst Rev. 2012;5, CD009779.

31. Kim YS, Kim Y, Kim SK, Rhim H. Computed tomographic differentiation between alcoholic and gallstone pancreatitis: Significance of distribution of infiltration or fluid collection. World J Gastroenterol. 2006;12:4524–8.

32. Singh M, Simsek H. Ethanol and the pancreas. Current status. Gastroenterology. 1990;98:1051–62.