The management of the patient with acute pancreatitis: from evidence to clinical practice

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ABSTRACT

Acute pancreatitis is an inflammatory disorder of the pancreas characterized by abdominal pain and elevation of pancreatic enzymes in the blood. The pathogenesis is complex and partly unknown and the evolution is often unpredictable. Many efforts have been made to define this disease and its complications and to classify different grades of severity in order to formulate prognostic scores that could guide the physician in choosing the optimal therapeutic setting and procedures. The management of the patient with pancreatitis is not always optimal and differs among internist, gastroenterologist or surgeon. We think that a patient with clinical suspicion of acute pancreatitis is admitted to medical or surgical department depending on the availability of beds and not according to evidence-based medicine. The aim of this monograph is to identify the optimal management of patients with acute pancreatitis admitted to hospital.

Introduction

Acute pancreatitis (AP) is an inflammatory disorder of the pancreas characterized by abdominal pain and elevation of pancreatic enzymes in the blood. The management of the patient with pancreatitis is not always optimal and differs among internist, gastroenterologist or surgeon.1 We think that a patient with clinical suspicion of acute pancreatitis is admitted to medical or surgical department depending on the availability of beds and not according to evidence-based medicine.2 The aim of this monograph is to identify the optimal management of patients with acute pancreatitis admitted to hospital.

Epidemiology

AP is one of the most common gastroenterological diseases requiring hospital admission. Its incidence varies between 4.9 and 73.4 cases per 100,000 worldwide with an increasing trend.3–7 In the US, it has been reported from 13 to 45 cases per 100,000 persons,8,9 causing 270,000 hospital admissions annually with an estimated 2.6 billion dollars per year inpatient costs. Incidence and etiology may vary in different countries in relation to different data recording methods, diagnostic criteria and facilities and local risk factors.10 About 20–30% of patients with AP have recurrence and about 10% develop chronic pancreatitis. Mortality is approximately 1%.9

AP equally affects men and women and risk increases with aging. Risk factors include alcohol, smoking, obesity and diabetes. A recent meta-analysis showed that obesity (expressed as body mass index superior to 30) is associated with increased risk of AP and severity of disease.11 Diabetes increases AP risk of 1.5–3 folds.9

Pathogenesis

Pathogenesis of this disease is not fully understood. Various experimental models have been used during decades. The most common are: the use of secretagogues (the most frequently used is cerulein, a cholecystokinin analogue. In this model pancreatitis is usually mild); the ligation of common bile duct (re-
Pathophysiology

The revised classification of Atlanta criteria identifies two phases of the disease: early and late. The first (within 1 week) is characterized by the systemic inflammatory response and/or organ failure and it is classified as mild, moderate or severe. The second is characterized by local and systemic complications.

In mild AP, the most common, there is no organ failure, local or systemic complications and resolution usually occurs in the first week.

Moderately severe AP is defined by the presence of transient organ failure, local complications or exacerbation of co-morbidities.

Severe AP is defined by persistent organ failure. The modified score of Marshall, based on three organ systems dysfunctions, considers organ failure in case of a score greater than 1. This score has the advantage of being simple, universally applicable and allows a stratification of patients easy and objective but has not been validated.

Local complications: acute peripancreatic fluid collections, pseudo cyst, acute necrotic collections and walled-off necrosis, thrombosis splenic and portal, and colonic necrosis syndrome and gastric outlet dysfunction.

Systemic complications are considered exacerbations of pre-existing conditions such as chronic obstructive pulmonary disease or coronary artery disease.

Etiology

Causes of acute pancreatitis are summarized in Figure 1.

Gallstones are the most common cause of AP in many studies. A review of 18 studies from European countries confirmed these data with the exception of Denmark and Sweden, where alcohol was the most common cause. In Italy a study on 1005 patients confirmed biliary etiology in 60% of patients. Risk of gallstones pancreatitis increases with age and is higher in women. Other causes of ampullary obstruction associated with pancreatitis include biliary ascariasis, periampullary diverticula, pancreatic and periampullary tumors, intraductal papillary mucinous neoplasms.

Considering different countries, the role of alcohol varies because of different life style. In the Unites States it accounts for the 30% of cases. In Europe it is the second cause of pancreatitis in most countries, with the exception of Sweden and Denmark. Differences are also seen in different regions of the same country. In the Italian multicenter survey on AP alcohol was responsible for 8.5% of AP (cases). Studies on different quantity of alcohol intake gave different results. Binge drinking does not appear to increase the risk of AP in general population. The risk seems indeed to be higher in subjects with a previous history of heavy drinking.

Alcohol is undoubtedly associated with AP but only 5% of drinkers develop pancreatitis, suggesting that it is not sufficient to determine the disease. Additional insult could be considered: smoking, high fat diet, obesity, genetics and infectious agents. The mechanism by which alcohol induces AP is not well understood. It has been proposed that it can cause biochemical and molecular changes in acinar cells that sensitize the pancreas to injury.

Metabolic abnormalities predisposing to AP include hypercalcemia and hypertriglyceridemia (usually more than 1000 mg/dL) that account for 1 to 4% of cases of acute pancreatitis. Type V hyperlipidemia, as well as types I and IV are prominent causes of acute pancreatitis. Secondary hyperlipidemia is caused by alcohol in-
take, pregnancy, estrogen therapy, and diabetes. Regarding hypercalcemia, it is, independently from its cause, a rare cause of pancreatitis. Calcium deposition in pancreatic duct and calcium activation of fibrinogen have been proposed as possible mechanisms.

Pancreatitis is a common complication of endoscopic retrograde cholangiopancreatography (ERCP). A systematic survey of studies from 1987 to 2003 found a 3.5% of post ERCP pancreatitis with 0.4% of patients experiencing severe AP and 0.11% death. Various mechanisms have been proposed: mechanical injury following instrumentation of papilla or thermal injury or chemical insult following injection of contrast medium, intraluminal activation of proteolytic enzymes, infections.

Drugs from different classes are a recognized cause of AP. Drug induced acute pancreatitis (DIAP) represents 0.1%-2% of overall cases. In most cases DIAP seems to be due to an idiosyncratic effect, that is to say an unpredictable abnormal interaction between the drug and the organism. Drugs may act inducing sphincter of Oddi dysfunction, like opioids or causing a pseudolithiasis like ceftriaxone, or increasing the risk of gallstones, like HMG-CoA-reductase-inhibitors. They may also generate toxic metabolites (like nucleoside reverse transcriptase inhibitors) or induce hypertriglyceridemia (like estrogens), or give an immuno-mediate reaction (like sulfonamides). Drugs inducing pancreatitis are classified in four classes: Ia (drugs with at least one case report, evidence of a positive re-challenge, and exclusion of other causes of AP, such as codeine, cytarabine, dapsone, enalapril, furosemide, isoniazid, mesalamine, metronidazole, pentamidine, pravastatin, simvastatin, sulfamethoxazole, sulindac, tetracycline, valproic acid), Ib (the same but without exclusion of other causes of AP such as amiodarone, azathioprine, dexamethasone, lamivudine, losartan, 6-MP, premarin, trimethoprim-sulfamethoxazole), II (at least four case reports with a consistent latency period for at least 75% of the cases, for example: acetaminophen, clozapine, erythromycin, estrogen, propofol, tamoxifen), III (at least two case reports but without re-challenge data or a consistent latency period, like alendronate, carbamazepine, ceftriaxone, clarithromycin, cyclosporin, hydrochlorothiazide, ribavirin, metformin, minocycline, naproxen, prednisone, prednisolone), and IV (one case report without re-challenge data, for example ampicillin, cisplatin, colchicine, cyclophosphamide, dicrofenac, doxorubicin, interleukin-2, octreotide, propoxyphene, rifampin, risperidone, sertraline, tacrolimus, vincristine).

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**Biliary gallstones, biliary sludge**

**Alcohol**

- Mechanical obstructions to flow of pancreatic juice
- ✔️ Ampullary: tumors, stricture or dysfunction of SOD
- ✔️ Ductal: stones, strictures, masses and tumors, mucus, parasites

**Metabolic: hypertriglyceridemia hypercalcemia**

**Drugs**

**Ischemia:** Hypotension, arteritis, embolic

**Hypothermia**

**Infections:** Virus, bacteria and mycobacteria and parasites

**Toxins and venoms (spider)**

**Autoimmune:** may be associate with autoimmune diseases (sicca syndrome, PBC, autoimmune hepatitis, celiac disease)

**Genetic (familial, sporadic)**

**Anatomic variants:** Pancreas divisum, choledochal cyst, duodenal diverticula

**Idiopathic**

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Figure 1. Causes of acute pancreatitis. SOD, sphincter of Oddi dysfunction; ERCP, endoscopic retrograde cholangiopancreatography; PBC, primary biliary cirrhosis.
Acute pancreatitis has also been reported during therapies with interferon (both in the standard and pegylated form) in patients with chronic hepatitis B and C (in these cases in association with ribavirin), may be due to immune modulation effects of the drug.18-20

Also therapies with tyrosine kinase receptor inhibitors (sorafenib and axitinib) could be complicated by acute pancreatitis (with a very low incidence, despite the common detection of hyperamylasemia and hyperlipasemia).21,22

Autoimmune pancreatitis is a rare cause of the disease and may present as acute or chronic. Two types with different histopathologic patterns have been identified. Type 1 is characterized by high levels of IgG4 and is associated to other extrapancreatic manifestations of IgG4 related disease; in type 2, that may be associated to inflammatory bowel disease, levels of IgG4 re normal.23

Other causes of AP may be: trauma (both blunt and penetrating, and also instrumentation, e.g. ERCP), ischemia, arteritis (for example in lupus). Infectious agents also can cause AP: Mumps, Coxsackie, cytomegalovirus, Salmonella typhi, Leptospira, Legionella, Aspergillus, Toxoplasma, Mycoplasma, Cryptosporidium, Mycobacterium tuberculosis and other mycobacteria. Among viral infections we have to remember HIV (in which pancreas can be affected by the virus itself or be damaged by drugs)14,14 and hepatitis viruses. In fact, AP has been reported during hepatitis due to hepatitis A, B, C, E virus.24 Both fulminant and, rarely, non-fulminant hepatitis may cause AP.25 Pancreatic damage may be due to direct inflammation and destruction of pancreatic acinar cell by the hepatitis virus,26 or related to immune response against virus.27

We also have to remember toxins, such as organophosphate pesticides, venoms of same arachnids and reptiles.

Structural abnormalities may predispose to AP, such as pancreas divisum (but only few patients develop AP in spite of frequency of this condition that interest 7-8% of white people), choledochal cyst. In this category, the most harmful are malignancies since tumors may cause mechanical obstruction at ampullary or ductal level and AP episodes may precede overt appearance of the neoplasia.14 Sphincter of Oddi dysfunction is due to stenosis or spasm of the sphincter.

Finally, we have to remember idiopathic pancreatitis recently defined as the third cause of pancreatitis. Further insights in etiology and pathophysiology should allow the decrease of unexplained cases.

Diagnosis

The American College of Gastroenterology guidelines1 state that two of these three findings should be present for the diagnosis of acute pancreatitis: i) abdominal pain consistent with the disease; ii) serum amylase and/or lipase greater than three times the upper limit of normal; and/or iii) characteristic findings from abdominal imaging (strong recommendation, moderate quality of evidence).

The new classification divides AP in three degrees of severity: mild, moderately severe and severe.4

The mild form is characterized by the absence of organ failure and local or systemic complications. It is not required to perform imaging methods and mortality is almost absent. Discharge within the first week. The moderately severe form is characterized by the presence of organ failure or transient local or systemic complications in the absence of renal organ persistent. It may resolve without intervention or may require prolonged specialized care. Its mortality rate is far lower than that of the severe form. The severe form is characterized by failure organ persistent (>48 h). When systemic inflammatory response syndrome (SIRS) is present, there is an increased risk of severe pancreatitis, and in the presence of persistent organ failure since the early days mortality can be very high and is reported up to 36-50%.

Clinical presentation

Most patients with acute pancreatitis have acute onset of persistent, severe epigastric abdominal pain.28 In some patients, the pain may be in the right upper quadrant or, rarely, confined to the left side. In patients with gallstone pancreatitis, the pain is well localized and the onset of pain is rapid, reaching maximum intensity in 10 to 20 min. In contrast, in patients with pancreatitis due to hereditary or metabolic causes or alcohol, the onset of pain may be less abrupt and the pain may be poorly localized. In approximately 50 percent of patients, the pain radiates to the back.29 The pain persists for several hours to days and may be partially relieved by sitting up or bending forward. Approximately 90 percent of patients have associated nausea and vomiting which may persist for several hours.30

 Patients with severe acute pancreatitis may also have dyspnea due to diaphragmatic inflammation secondary to pancreatitis, pleural effusions, or adult respiratory distress syndrome. Approximately 5 to 10 percent of patients with acute severe pancreatitis may have painless disease and have unexplained hypotension (e.g., postoperative and critically ill patients, patients on dialysis, organophosphate poisoning, and Legionnaire’s disease).31-35

Finally, to evaluate the evolution of pancreatitis it is important to determine the exact interval between symptom onset and hospitalization and in case of transfer center specializing in the interval between the first admission and transfer.4

Physical findings

Physical findings vary depending on the severity of acute pancreatitis. In patients with mild acute pan-
creatitis, the epigastrium may be minimally tender to palpation. In contrast, in patients with severe pancreatitis, there may be significant tenderness to palpation in the epigastrium or more diffusely over the abdomen. Patients may have scleral icterus due to obstructive jaundice caused by choledocholithiasis or edema of the head of the pancreas.

Patients with severe pancreatitis may have fever, tachypnea, hypoxemia, and hypotension. In 3% of patients with acute pancreatitis, ecchymosis discoloration may be observed in the periumbilical region (Cullen’s sign) or along the flank (Grey Turner sign).35

These findings, although nonspecific, suggest the presence of retroperitoneal bleeding in the setting of pancreatic necrosis.36 Patient may also have findings suggestive of the underlying etiology. As examples, hepatomegaly may be present in patients with alcoholic pancreatitis, xanthomas in hyperlipidemic pancreatitis, and parotid swelling in patients with mumps.

**Laboratory findings**

The two main enzymes that aid diagnosis of acute pancreatitis are amylase and lipase. Serum amylase in AP patients generally rises within a few hours after the onset of symptoms and normalizes within 3-7 days. On admission, it can remain in the normal range in a fifth of patients.37,38 Compared with lipase, serum amylase returns more quickly to values below the upper limit of normal. Due to the low sensitivity and specificity serum amylase alone cannot be used reliably for the diagnosis of AP.

Lipase rises within 4-8 h and stays elevated for 8-14 days. Lipase levels of greater than five times the upper limit of normal have 100% specificity for the diagnosis of AP. Lipase is now the laboratory measurement of choice for the diagnosis of AP. Serum lipase or amylase levels at admission do not predict severity of disease. Other enzymes can be elevated in AP. Trypsinogen activation peptide (TAP), a five amino-acid peptide that is cleaved from trypsinogen to produce active trypsin, is elevated in acute pancreatitis. Since activation of trypsin is likely an early event in the pathogenesis of acute pancreatitis, TAP may be useful in the detection of early acute pancreatitis and as a predictor of the severity of acute pancreatitis.39-42 Urinary and serum trypsinogen-2 levels are newer tests and are elevated in early acute pancreatitis. They are not readily available and additional studies are needed to determine their role in the diagnosis of acute pancreatitis.43-46

Acute pancreatitis is also associated with elevations in C-reactive protein (CRP), IL-6, IL-8, IL-10, TNF, and PMN elastase.46 A CRP level above 150 mg/dL within 48 h is associated with severe pancreatitis.

Additional laboratory tests that are useful for the diagnosis of AP are liver function tests and hematocrit.

In patients with no history of alcohol consumption, the presence of alanine aminotransferase (ALT) elevation three times the upper limit of normal has a 95% positive predictive value for acute gallstone pancreatitis.47 Normal liver function tests do not exclude the diagnosis of biliary pancreatitis, as this can occur in up to 20% of patients.48 Hemocrit concentration (defined as hemocrit greater than 44%) and failure of hemocrit to decrease at 24 h are important predictors of severe pancreatitis. Metabolic abnormalities including elevated blood urea nitrogen (BUN), hypocalcemia, hyperglycemia, and hypoglycemia may also occur. Clinical conditions that may be associated with increased levels of amylase in the absence of AP are: macroamylasemia (a syndrome characterized by the formation of large molecular complexes between amylase and abnormal immunoglobulins); decreased glomerular filtration rate; diseases of the salivary glands; extrapancreatic abdominal diseases associated with inflammation (acute appendicitis, cholecystitis, intestinal obstruction or ischemia, peptic ulcer, and gynecological diseases).

**Role of ultrasound in acute pancreatitis**

Ultrasound (US) represents the first-line imaging technique in the assessment of pancreatic disease. It has several advantages as compared to contrast-enhanced computed tomography (CT) and magnetic resonance imaging (MRI), such as low cost, real time evaluation and no radiation exposure.49 US can be performed easily at patient’s bedside and demonstrates good values of sensitivity and specificity in the assessment of AP (67 and 100%, respectively).50 Furthermore, US is useful in the follow-up of the patients, revealing local complications such as abscesses, necrosis, peripancreatic fluid, pseudocysts, abscesses and vascular disease (venous thrombosis and pseudoaneurysms).

The pancreas, however, is sometimes difficult to explore. Patients are often obese, and meteorism can limit its evaluation, for example in the case of local distension of intestinal loops by pancreatic inflammation (sentinel loop). In these cases, the visualization can be improved by water filling and changing in position (from supine to upright). With these measures, the pancreas can be fully displayed in a large percentage of patients.51

**Conventional ultrasound: B-mode and color Doppler findings**

The inflammation can involve diffusely the pancreatic gland or only some portions such as the head and body. On conventional US, the gland appears enlarged but this feature can be not so evident in the case of mild, edematous pancreatitis. An anteroposterior size of the body greater than 24 mm is traditionally suggestive of increased gland.52
Some authors, however, believe that the increase in size of the pancreas in the course of acute pancreatitis should be considered relative to the gland size in normal conditions, as the latter may vary from subject to subject and should not be strictly correlated to an absolute value.

A more pronounced edema can manifest as mass effect on the adjacent organs (gastric antrum, veins regional site, bile duct), suggesting the increase in pancreas size, avoiding further measurement of the gland in the correspondence of the body. Margins can appear unchanged or irregular, blurred or polycyclic in the case of severe disease.\textsuperscript{55}

Echogenicity

The gland appearance on B-mode US varies in relation to the clinical stage and severity of pancreatitis. In mild disease, the echogenicity is similar to that observed in normal conditions. A more pronounced inflammatory damage of the gland appears instead diffusely hypoechoic in the edematous form or occur with appreciable inhomogeneity with the necrotic-hemorrhagic involvement. The necrotic areas, in particular, can appear hypoechoic as compared to the healthy parenchyma, and their recognition is important because often enters in the differential diagnosis with neoplastic disorders such as pancreatic adenocarcinoma. A correct diagnosis is based not only on the US features but should be integrated also on clinical and laboratory findings.\textsuperscript{52,53} The appearance of hemorrhagic areas varies on US depending on the state of red blood cells aggregation and the phase in which it is detected. In the early stages, acute phase the hemorrhagic areas are hyperechoic; in the late stages, the echogenicity decreases following the clot rupture of the clot.\textsuperscript{53} Furthermore, the inflammation of the retroperitoneal tissue can be seen sometimes as a hypoechoic, perivascular halo around the portal and splenic veins.\textsuperscript{54}

Color Doppler ultrasound findings

Color Doppler US (CDUS) is an important step in the assessment of pancreas and its vascular structures, in particular the portal vein, splenic, mesenteric arterial and venous vessels, aorta and inferior vena cava.

While peripancreatic vessels are well assessed on CDUS, only few intraparenchymal vessels can be fully appreciated in normal conditions.

However, the increasing diffusion of high sensitive US devices can allow a greater visualization of small, peri and intrapancreatic vessels, that is particularly important in the evaluation of vascular complications of acute pancreatitis such as venous thrombosis and pseudoaneurysms.\textsuperscript{55}

Portal vein thrombosis can manifest on CDUS as partial or complete absence of flow within intraluminal echogenic areas. CDUS can also be reliable to evaluate suspected pseudoaneurysms arising from mesenteric or splenic arteries, secondary to tunica media disruption by pancreatic inflammation; pseudoaneurysms appear as small anechoic lesions on B-mode but their vascular origin can be revealed by CDUS that show turbulent flow or typical mosaic pattern on color Doppler and arterial flow on spectral analysis.\textsuperscript{55}

Evaluation of common bile and pancreatic ducts

US evaluation of acute pancreatitis is not complete without the study of biliary and pancreatic ducts. Cholelithiasis is most frequently associated with acute pancreatitis in developed countries. Gallstones appear as hyperechoic, intraluminal formations showing posterior shadow cone, fully movable on changing the patient’s position, thus leading to the so-called rolling stone sign in case of movement along the rear wall. Duct stones in the common bile duct (CBD) have similar US characteristics; however, the mobility with the change of the position of the patient can be less easily appreciated. The biliary sludge can manifest as multiple echoes slowly floating with changes of decubitus and usually distributes along the rear wall of the gallbladder or within the CBD lumen. It usually does not generate posterior shadow cone, differently from traditional stones.\textsuperscript{56}

CBD stones can result often in the dilatation of intra or extrahepatic bile ducts, the former evident with the classic pruned-tree appearance, due to the presence of enlarged and convergent intrahepatic common bile ducts. CBD dilatation can be easily recognized as exceeding in general 8 mm of diameter. Rarely, CBD stones can be associated with dilatation of duct of Wirsung; in these cases, the dilatation derives usually from mass effect secondary to the focal edema in the head of the pancreas. Under normal conditions the size does not exceed 2-3 mm; a diameter exceeding 3 mm must raise the suspicion of obstructive and/or inflammatory etiology.\textsuperscript{53,55}

Other US signs that can be found in association with AP are cholecystitis and cholangitis.\textsuperscript{53}

Locoregional and extra-glandular complications

Complications of acute pancreatitis are classified into locoregional, if they grow in close proximity of the pancreatic gland affected by inflammation, and extra-glandular, if are localized outside of the pancreatic parenchyma. The presence of inflammatory exudate may be associated with the formation of fluid collections that can localize in or around the pancreatic gland, sometimes under the capsular layer or outside the gland, inside adjacent organs (liver, spleen and kidneys). The sonographic features of such collections consist of areas with not well-defined borders with hypo-anechoic echostructure, which sometimes include reinforcement of the wall due mostly to the
presence of fat edematous and infiltrated. The feature of different echostructural homogeneity is usually due to the presence of echogenic debris, septa and/or hemorrhagic fluid.  

The fluid collections located under diaphragm muscle can propagate in the pericardial, pleural or mediastinal cavities; in about the half of cases they regress spontaneously; Otherwise, if not reabsorbed, they may be organized in structures better defined as pseudocysts. They appear, when studied through B-mode examination, as not homogeneous areas, with variously echogenic mobile iso-anechoic content, covered with thickened wall with reinforced rear and mostly regular margins. The pancreatic bed on which pseudocysts are located is irregular with echogenic fragments; the latter are the result of necrotic and autolytic processes. Usually there is no vascular signal at CPD examination.

In this context, ultrasound contrast-enhanced ultrasound (CEUS) can add useful information for the differential diagnosis between pseudocyst and cystadenoma. The pseudocyst usually appears as non-enhanced and also the intraluminal echogenic component, if present, does not tend to take ultrasound contrast agent. Pseudocysts newly formed can have a wall hyperenhanced, unlike the old ones that have, instead, hypoechoenhanced walls.

Rarely pancreatic pseudocysts may be confused with bilomas, rare collections of abnormal bile in the intra- or extra-hepatic due to a spontaneous or iatrogenic interruption of biliary system. The sonographic appearance may be almost comparable to that of pseudocyst. The suspected diagnosis will be placed according to an history of previous trauma or surgery of the biliary tree, and confirmed by the imaging of second level (MR cholangiopancreatography), which will highlight the close relationship of continuity of the lesion with the biliary tree.

Another locoregional complication is represented by abscess or by superinfection of a fluid collection or by liquefaction of a necrotic area. It may constitute a serious complication that can lead to abrupt clinical deterioration and that needs, therefore, timely assessment of advanced cares. An abscess should be suspected in case of signs of infection (e.g., neutrophilic leukocytosis and fever) and the occurrence of abdominal pain; the sonographic appearance consists of an increase in the volume of the collection with onset within echogenic material sometimes sloping or fluctuating.

Another serious complication is represented by infected pancreatic necrosis. It has no sonographic features, which make it distinguishable from the other fluid collections, and usually it occurs with the appearance of fine small echoes within the liquid, variously mobile and floating.

### Role of contrast-enhanced ultrasound

In the last decade, conventional B-mode US has been improved by CEUS that has improved the diagnostic abilities of ultrasound thanks to a great definition of vessels and microvessels of the examined districts. US contrast agents (UCAs) are exogenous substances that can be administered intravenously to enhance the ultrasound signal. All agents contain gas-filled microbubbles with a diameter of 2-6 μm, surrounded by a shell composed of varying lipids or polymers. The gas and the shell influence the half-lives of the microbubbles and their response to insonation.

When hit by US, the microbubble sends back to the transducer a wave that has a frequency equal to that of insonation and a series of harmonics of the fundamental frequency. Among these, the second harmonic has a frequency double than the fundamental. The harmonic response is high as the acoustic pressure of the incident wave, but causes a greater destruction of microbubbles. If the microbubbles have elastic membrane, it is possible to apply a lower acoustic pressure to generate a good harmonic response. Sonovue has a better harmonic response with a frequency of 3 to 3.5 MHz; therefore, the second harmonic is 6-7 MHz. The structures produce both the fundamental frequency and harmonics of fundamental frequency (tissue harmonic imaging) generated by the distortion of the wave by crossing tissue; these harmonics are weaker than those produced by the microbubbles and employing dedicated platforms is possible to distinguish them from harmonics generated by the microbubbles. Harmonic tissue can cause artifacts when operators use higher-pressure acoustic signal.

Second generation contrast agents, such as BR-1 and Sonovue are those mainly used in Europe. Sonovue is composed of sulfur hexafluoride with a phospholipid shell, which provides stability and resistance. Sonovue generates non-linear harmonic frequencies, since at low acoustic power of insonation the degree of microbubbles expansion is greater than its destruction.

These new UCAs oscillate without destruction at low mechanical index, producing harmonic frequencies that are multiples of the transmitted frequency (non-linear fundamental echoes), allowing real-time imaging of microbubble signals. The introduction of a dedicated contrast agent software in the ultrasound equipment enables the visualization of the microbubble signals without the fundamental grey-scale echoes.

Since its infusion, contrast agent takes about 15-30 s to reach the structures of interest.

Then, gas is eliminated from the lung while the components of the membrane from liver and kidney.

Second-generation contrast agents are not indicated in case of: recent acute coronary syndrome, unstable angina, recent acute heart attack, recent coronary artery...
intervention, acute or class III or IV chronic heart failure or severe arrhythmias. No interaction with other drugs has been reported and only mild and transient adverse reactions have been reported.

CEUS examination should be performed after a conventional B-mode study. The probe used is a multifrequency curved array transducer (3-4 MHz) and an intravenous bolus of 2.4 mL of contrast agent is administered.

In the study of pancreatic disease, CEUS is not recommended for the detection of pancreatic lesions but it is useful to define more precisely lesions already found through traditional ultrasound examination.59

Different patterns of enhancement are related to different pathological conditions, depending on characteristics of the lesions (focal or widespread, several or single, solid or liquid, neoplastic or pseudo-neoplastic). The enhancement pattern of focal pancreatic lesions should be compared with the adjacent pancreatic texture. Therefore, the examination should include the mass under investigation and a portion of surrounding pancreatic parenchyma. In order to understand the meaning of different ways of contrast distribution, it is important to specify that pancreas enhancement is far different from liver enhancement: blood contribution to the pancreas is completely arterial, and CEUS shows an early and brief enhancement of pancreatic gland. Arterial phase is very early (10 to 30 s with a peak of enhancement to 15-20 s), and is followed by a transient venous phase (30 to approximately 120 s) during which spleno-mesenteric-portal venous axis is enhanced. The late phase (about 120 s after injection) is defined by enhancement of the hepatic veins. These characteristics make the study of the pancreas particularly difficult.

As contrast agents have a pure intravascular distribution, CEUS of the pancreas is suitable to discern between solid and cystic lesions, to describe focal masses, and to provide a clear differentiation between surrounding tissue, fibrosis and necrosis.

CEUS improves the diagnosis of the inflammatory pathology of the pancreas and is particularly useful in the staging of the severity of the acute pancreatitis, in the detection of area of necrosis and in the recognition of its complications.60

Edematous acute pancreatitis appears at a conventional B-mode examination, as widely described before, with a dimensional increase of the gland and a diffusely hypoechoic texture. The injection of contrast agent will show a diffused and homogeneous wash-in of the gland, with different degrees of enhancement, resulting in an increased echogenicity during the dynamic phases.

Autoimmune pancreatitis is a pathological condition characterized by inflammation around the ducts sustained by lymphocytes migration, which leads to fibrosis. The features showed at B-mode study are similar to focal acute pancreatitis: the gland appears enlarged, hypoechoic and pancreatic duct results expanded. CEUS shows different aspects: some authors have reported enhancements similar to that of the normal pancreatic parenchyma, others have described a mild or high wash-in followed by a slow wash-out of contrast agent.61

Focal acute pancreatitis appears at conventional B-mode examination as an expanded area of the pancreas, homogeneously hypoechoic. Sometimes, such images set serious problems of differential diagnosis with pancreatic tumor masses. The principal indication of the CEUS consists in the characterization of pancreatic lesions found at traditional ultrasound performed in a context of an acute pancreatitis. Focal acute pancreatitis is underlined by CEUS as a zone of increased impregnation of contrast agent. Necrotic area appears as an anechoic image on B-mode study without enhancement after contrast injection.62

Although the gold standard in the diagnosis of the necrotic areas in severe acute pancreatitis is represented by CT, CEUS constitutes currently the best technique in the follow-up of these patients after CT staging, because it reduces the exposure to radiations.63

Moreover, CEUS is useful in the recognition and in the characterization of pancreatic pseudocysts. Such lesions appear at B-mode examination as inhomogeneous areas, circumscribed by a not well-defined wall, containing sometimes echoic mobile material inside, and have a hypo-anechoic texture. CEUS will show no enhancement of the wall of pseudocyst in all phases, even if inhomogeneous on US. The sensitivity and specificity of CEUS in characterizing pseudocysts is up to 100%. Therefore, CEUS has a great importance in the differential diagnosis of pseudocysts and cystic tumors of the pancreas, whose walls result instead hyper-enhanced. Contrast agent injection makes easier the diagnosis of these formations, thanks to a more detailed characterization of the vascular component of cystic inner elements.

As reported in The EFSUMB Guidelines and Recommendations on the Clinical Practice of Contrast Enhanced Ultrasound (CEUS): Update 2011 on non-hepatic applications the recommended uses and indications of CEUS in patients with focal pancreatic lesions identified with US are: i) characterization of ductal adenocarcinoma (Recommendation Level: A;1b); ii) differential diagnosis between pseudocysts and cystic tumors (Recommendation Level: A;1b); iii) differentiation of vascular (solid) from avascular (liquid/necrotic) components of a lesion (Recommendation Level: A;1b); iv) defining the dimensions and margins of a lesion, including its relationship with adjacent vessels (Recommendation Level: B;2b); v) management of the lesion with a better distinction between solid and cystic lesions, thus providing information for the choice
of the next imaging modality (i.e., MRI and/or endoscopic US for cystic lesions) (Recommendation Level: C;5); vi) diagnosis of cases that are indeterminate on CT (vascularization of solid pancreatic lesions; differential diagnosis between pseudocysts and pancreatic cystic tumors, especially mucinous cystic tumor) (Recommendation Level: C;5).64

A new technique of imaging, CEUS, combines the advantages of endoscopic US with amplification by contrast agents.65 After EUS examination CEUS is used to characterize micro-vascularization of the lesions, to differentiate benign from malignant masses, to improve diagnosis, staging of the lesions and to improve therapeutic procedures.

CEUS can be performed with a high or low mechanical index. This technique can be useful in the differential diagnosis of autoimmune pancreatitis. Focal and diffuse autoimmune pancreatitis result hyper-enhanced at CEUS examination. Some Authors have described that a low MI at CEUS shows hyperenhancement in focal and diffuse autoimmune pancreatitis and allows the differential diagnosis with ductal adenocarcinoma (which does not hyper-enhance).66

Role of pancreatic endoscopic ultrasound

EUS represents a new mild invasive technique of imaging that combines the advantages of B-mode ultrasound with those of endoscopy. Thanks to the proximity of EUS transducer to the organs of interest, the images obtained are frequently more accurate and more detailed than the ones obtained by traditional ultrasound. EUS can obtain information about the layers of the intestinal wall as well as adjacent areas such as biliary ducts, pancreatic structures, lymph nodes and blood vessels. These features have encouraged its use in clinical practice.67 Numerous studies have recently testified for the safety, the accuracy and the indications of this exam, whose real limit is the hold dependence from the experience of the operator.

In the acute inflammatory pathology of pancreas, EUS allows: i) to clearly identify the lithiasis etiology of acute pancreatitis thanks to its ability in detecting stones inside biliary structures or pancreatic ducts; ii) to drain pancreatic pseudocysts; iii) to differentiate between pancreatic pseudocysts and biliary or pancreatic tumors; and iv) to obtain a fine needle aspiration of structures of interest.

Even if not performed during acute pancreatitis it can be used further to clarify the causes of inflammatory process or to treat its complications.

Biliary lithiasis

The lithiasis of main biliary duct prevails in 20% of patients affected by gallbladder lithiasis and causes the development of severe complications such as acute pancreatitis and acute cholangitis. The diagnosis of lithiasis of common biliary duct is not always easy and clinical tests and conventional imaging are not endowed with elevated sensitivity. EUS offers, on the contrary, high sensitivity and specificity and represents a valid new diagnostic technique and a reliable alternative to ERCP. The suspicious cases for lithiasis of the principal biliary tract, where traditional ultrasound has not been diagnostic, require the execution of colangio-MRI or EUS. B-mode ultrasound has a low sensitivity in the detection of stones (22-55%); it rather gives indirect information about the presence of lithiasis through the detection of expansion of the biliary system, with a sensitivity from 77% to the 87%. The sensitivity of colangio-MRI in the diagnosis of lithiasis of principal biliary duct goes from 85% to 92% and the specificity from 93% to 97%, with a meaningful decrement of this value with decreasing sizes of calculi. The reliability of EUS is not conditioned instead from the dimensions of stones: EUS is able to detect biliary deposits even if they are not seen by ERCP, with a sensitivity between 89% and 94%, and a specificity that goes from 94% to 95%. The underestimated biliary lithiasis by ERCP suggests that the echo-endoscopic study of the extra-hepatic biliary ducts could represent a valid alternative to obtain diagnostic information. EUS has instead largely reduced the cases of idiopathic pancreatitis and it has progressively limited the use of ERCP in those cases, which need therapeutic endoscopy. The study of the biliary and pancreatic ducts by EUS is technically limited in case of: pneumobilia, outcomes of interventions on stomach and masses of cephalic part of the pancreas. Despite these limits, EUS reaches a sensitivity of 97% and a specificity of 100% in the diagnosis of lithiasis of common bile duct, showing a higher diagnostic accuracy than conventional US and CT68,69 and a better safety profile than ERCP.70

Drainage of pseudocyst

Pancreatic pseudocysts are intra or extra-pancreatic fluid collections composed of pancreatic secretions and inflammatory debris. Reactive granulation tissue rather than a true epithelial lining wall surrounds the fluid collection, hence the term pseudocyst. Pseudocysts originate from leaks in the pancreatic duct. The etiology may be necrosis secondary to pancreatitis, progressive ductal obstruction, or trauma.

Most of pseudocysts are asymptomatic and do not require treatment. An enlarging pseudocyst may require drainage in order to avoid rupture or hemorrhage transformation. Pseudocysts that complicate acute pancreatitis have a high probability to spontaneously resolve within 4 to 6 weeks and should be observed in this period before considering further treatment. Earlier drainage may be indicated when clinical pancreatitis fails to improve despite an aggressive medical management.
Pseudocysts complicating chronic pancreatitis usually result from pancreatic duct outflow obstruction from a stone, stricture, or accumulation of protein materials. Such pseudocysts rarely resolve on their own. Drainage is indicated to relieve symptoms associated with a space-occupying mass and neighboring organ compression such as pain, gastric outlet obstruction, and jaundice. Drainage is also indicated when pseudocysts become infected or if there is intracystic bleeding.

The application of EUS to guide pseudocyst puncture through the stomach or duodenal wall has improved the success and safety of endoscopic pseudocyst drainage. Using endoscopic guidance alone, a prominent mucosal bulge should be present to identify the site for pseudocyst puncture. Even though, the interposed tissue may contain vessels. EUS provides a highly detailed view of the pseudocyst and surrounding topographical anatomy. Surface vessels are readily detected with color Doppler. The use of a sectorial echo-endoscopic tool with a wide channel of puncture ensures safety and easy execution. With this technique, the drain of the pseudocyst is performed easily, reducing significantly operating risks.

Biliary and pancreatic tumors

The tumors of biliary ducts and pancreatic gland are characterized by high malignancy and difficult surgical treatment. These features determine a low survival. An accurate study of such tumors is desirable to select patients who are candidates for surgical intervention.

EUS has changed radically the study of the tumors of biliary and pancreatic structures: this technique has revealed to be superior to CT in the staging and in detection of invaded lymph nodes. Moreover, EUS allows performing a fine needle aspiration of the lesion and of the satellite lymph nodes.

Unlikely traditional ultrasound, EUS allows an accurate identification and staging of ampullary tumors and of tumors of the principal biliary duct.

The diagnosis of chronic pseudocyst in the diagnosis of carcinoma of the common liver duct and of its bifurcation is about 85% with B-mode US and arises 91% if micro-probes are used. Cholangiocarcinoma of the principal biliary ducts appears as a small mass projecting inside the common bile duct or as a thickening of its wall. EUS is suitable in the diagnosis of pancreatic tumors whereas a strong clinical suspect exists in the absence of comforting imaging. EUS sensitivity and specificity to detect pancreatic tumors are 90% and 89%, respectively and accuracy is 81%.

The superiority of EUS in comparison to CT in the diagnosis of pancreatic tumors increases for lesions smaller than 3 cm of diameter. Morphologically, these tumors appear as hypoechoic lesions with poor defined margins or as a mass which compresses the principal biliary tract or pancreatic duct.

Focal pancreatic lesions represent the principal indication for EUS-driven biopsy; fine needle aspiration has a sensitivity of 94% and an accuracy of 92%. In case of cystic pancreatic lesions, this technique allows to distinguish among benign and malignant forms through a cytological analysis.

Complications

In comparison with standard endoscopy the risks of perforation are slightly greater because of a higher rigidity of the tool. In case of invasive endoscopy, the risks of bleeding and infection are higher. Such complications strongly depend on the experience of the operator.

Assessment and risk stratification

The correct clinical evaluation of the patient with AP is critical to identify the proportion of patients (15-25%) that will evolve towards a severe form of acute pancreatitis. The ability to predict the severity of the AP allows the identification of patients that benefit from an early transfer to intensive care or specific invasive interventions. In literature, concerning the risk of AP, there are several predictive models based on clinical, laboratory values, radiological risk factors, severity scores and serum markers. Unfortunately, these predictive models (which can be applied to the patient at the time of access to the emergency room or in the first 48 or 72 h) have a low specificity. Low specificity associated with low prevalence of severe AP translate into low positive predictive values.

Table 24 summarizes the main features of the score system used in clinical practice.

Therapy of acute pancreatitis

The treatment of acute pancreatitis is divided into three stages: i) treatment of acute phase; ii) treatment of complication; iii) treatment of predisposing factors.
Table 1. Clinical predictors, laboratory and radiological of acute pancreatitis gravity.

<table>
<thead>
<tr>
<th>Clinical</th>
<th>Laboratory</th>
<th>Radiological</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clinical judgment</td>
<td></td>
<td></td>
</tr>
<tr>
<td>It is the result of the impression of the physician evaluating the patient with AP at the emergency room. It is based on clinical and laboratory data.</td>
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<tr>
<td>Homocentrization</td>
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<td>Chest-X-ray</td>
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<tr>
<td>The studies that evaluated the hematocrit as a predictor of the severity of AP have produced variable results. It seems that a normal or low hematocrit at admission and during the first 24 hours is generally associated with a milder clinical course.</td>
<td>The presence of pleural effusion or pulmonary infiltrates during the first 24 h may be associated with pancreatic necrosis and organ failure.</td>
<td></td>
</tr>
<tr>
<td>Age</td>
<td></td>
<td>Abdominal computed tomography with contrast medium</td>
</tr>
<tr>
<td>Numerous studies have established that advanced age is an adverse prognostic factor, although the age limit is variable (55-75 age).</td>
<td>C-reactive protein Abdominal magnetic resonance and colangiopancreatic magnetic resonance</td>
<td></td>
</tr>
<tr>
<td>The value of the CRP is directly proportional to the severity of pancreatitis. It is an inexpensive and ready available test. CRP value &gt;150 mg/L after 48 h of onset of AP are a watershed between mild and severe forms.</td>
<td>Magnetic resonance imaging is of comparable diagnostic and prognostic value with computed tomography in the staging of acute pancreatitis.</td>
<td></td>
</tr>
<tr>
<td>Sex</td>
<td></td>
<td></td>
</tr>
<tr>
<td>It is not considered a predictor of outcome in many studies.</td>
<td>BUN</td>
<td></td>
</tr>
<tr>
<td>In some studies blood urea was found to be laboratory examination more useful in predicting mortality of AP. Values of BUN ≥20 mg/dL on admission, are associated with an increased risk of mortality as well as any subsequent increases in blood that occurred within 24 h.</td>
<td></td>
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<tr>
<td>Alcohol</td>
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<td></td>
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<tr>
<td>In many studies, alcohol is associated with an increased risk of pancreatic necrosis with the need for invasive treatments.</td>
<td>Serum creatinine</td>
<td></td>
</tr>
<tr>
<td>An elevated serum creatinine within the first 48 hours can predict the development of pancreatic necrosis. A normal creatinine has a high negative predictive value for the development of pancreatic necrosis. A normal creatinine in the absence of complications, may obviate the need to perform an abdominal CT.</td>
<td></td>
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<tr>
<td>Obesity</td>
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<tr>
<td>Obesity (BMI &gt;30) is a factor predictive of severe AP in many studies. A meta-analysis with 739 patients estimated: OR 2.9 (95% CI 1.8 to 4.6) for serious AP; OR 2.3 (95% CI 1.4 to 3.8) for systemic complications; OR 3.8 (95% CI 2.4 to 6.6) for local complications; OR, 2.1 (95% CI 1.0 to 4.8) for mortality</td>
<td>Other serum markers</td>
<td></td>
</tr>
<tr>
<td>Other serum markers have been variously tested in several clinical studies to predict the severity of the AP, including: urinary trypsinogen activation peptide, procalcitonin, amylase, lipase, polymorphonuclear elastase, pancreatic-associated protein, serum glucose, serum calcium, pro-carboxypeptidase B, carboxypeptidase B activation peptide, serum trypsinogen-2, phospholipaseA-2, serum amyloid protein-A, substance P, antithrombin III, platelet activating factor, interleukins 1, 6, and 8, tumor necrosis factor-alpha or soluble tumor necrosis factor receptor, many of which are not even available for daily use, and therefore they will not be considered due to their unproven clinical utility.</td>
<td></td>
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<tr>
<td>Short time interval between start of the symptoms and hospitalization</td>
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</tr>
<tr>
<td>A time interval less than 24 h is associated with increased severity of pancreatitis.</td>
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<td></td>
</tr>
<tr>
<td>Organ failure</td>
<td></td>
<td></td>
</tr>
<tr>
<td>The presence of early and persistent organ failure is widely regarded as a predictor of severe AP, increased mortality and prolonged hospital stay.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

AP, acute pancreatitis; CRP, C-reactive protein; CT, computed tomography; BUN, blood urea nitrogen; MRI, magnetic resonance imaging; BMI, body mass index; OR, odds ratio; CI, confidence interval.
The management of the patient with acute pancreatitis

Therapy of acute phase

In this phase the choice of an appropriate care setting is mandatory.\textsuperscript{1,115} There are indications for admission or transfer in a setting of intensive/sub-intensive type: hypotension unresponsive to fluid loading, multi-organ failure, persistent SIRS, increased urea or creatinine and hematocrit, the presence of severe cardiovascular-pulmonary comorbidities.

Referral to a specialist center is necessary if multi-organ failure persists and serious complications appear. It has been demonstrated that admission to a reference center is a factor that can influence the prognosis.\textsuperscript{116} The cornerstones of treatment at this stage are: i) fluid resuscitation and support of the circle; ii) pain control; iii) nutrition; iv) infection control; v) monitoring and correction of hypoxia; vi) control of blood glucose.

Fluid resuscitation

Hydration is the only therapy that has been shown to improve survival in AP.\textsuperscript{3,117,118} Therefore, the first therapeutic approach consists of an accurate assessment of the state of hydration and hemodynamic balance, which can be compromised by vomiting, pain, diaphoresis, increased perspiration, vasodilation linked to the release of cytokines and pancreatic enzymes that alter the microcirculation and increase edema and extravascular fluid loss.\textsuperscript{3}

The hydration therapy must be aggressive in the early stages (the first 6-12 h) to correct the blood volume and prevent the mesenteric hypoperfusion with the risk of ischemia and increased size of necrosis.\textsuperscript{119} In this way the hydration is also able to reduce the pain due to hypoperfusion.

Aggressive therapy consists in the infusion of 250-500 cc of isotonic crystalloid hourly. Physiological saline or Ringer’s lactate can be infused, but Ringer seems preferable unless in the presence of hypercalcemia.\textsuperscript{120} In cases of severe volume depletion a bolus of 20 mL/kg/h may be indicated for the first hour followed by a maintenance to 3 mL/kg/h (decrease to 1.5 mL/kg/h if BUN reduces) checking every 6-8 hours clinical status and vital signs.\textsuperscript{112,121}

The assessment of the state of hydration and the need for replenishing liquids should be guided by clinical and can be facilitated by some laboratory parameters such as BUN, creatinine and hematocrit.

The following targets are considered indicators of adequate hydration: heart rate <120 bpm, median arterial pressure between 65 and 85 mmHg, diuresis >0.5-1 mL/kg/h, decreased hematocrit and BUN.\textsuperscript{120,122} In a context of ICU type the variation of the stroke volume can be monitored for this purpose.

Infusion quantity and quality should consider degree of dehydration, electrolyte imbalances (including calcium and magnesium), diuresis and the presence of renal and/or cardiac complications and should be repeatedly evaluated at regular intervals and adjusted in the first 24-48 h.\textsuperscript{123}

After the first 24 h aggressive rehydration showed no benefit and may be counterproductive because it can cause abdominal compartment syndrome, volume overload and pulmonary edema.\textsuperscript{117,121,124} If you cannot note a reduction in azotemia, you could be in the presence of acute tubular necrosis.

Pain control

Pain control is essential for the well being of the patient and to prevent further hemodynamic impairment. It can be achieved in an effective and safe way with opioids (meperidine, morphine and fentanyl).\textsuperscript{125}

There are no studies that indicate the superiority of an opioid analgesic.

Nutrition

In the first 24-48 h, in cases of mild to moderate pancreatitis, fluid supplementation may be useful. In these cases, the re-feeding can take place as soon as abdominal pain, nausea and vomiting stop and inflammation markers decrease. The diet should not necessarily be liquid or semi-liquid, but should be low-fat and low residue.\textsuperscript{126,127}

In severe cases, artificial feeding is necessary with endoscopic or radiological positioning of naso-jejunal tube (preferred) and enteral nutrition formulas rich in protein and low in fat (with a caloric need of 25 kcal/kg of ideal weight). This approach allows, with respect to the total parenteral nutrition (TPN), to maintain the integrity of the intestinal barrier, to reduce the risk of intestinal atrophy and bacterial/fungal translocation and to reduce the risk of sepsis related to the use of central venous access (CVC) and TPN. This way the risk of necrosis infection also reduces.\textsuperscript{128}

In case it is impossible to position the naso-jejunal tube, data are in favor of nasogastric tube rather than the TPN: these data, however, should be confirmed.\textsuperscript{129}

Antibiotic therapy

There are only two conditions in which antibiotic therapy is recommended: infection and prophylaxis of sterile necrosis.\textsuperscript{3,115,130,131}

Infection of extrapancreatic sites (lung, urinary, biliary, CVC related) should be identified and treated with antibiotic therapy. Empirical therapy will be discontinued if culture tests are negative.

In case of pancreatic necrosis infection, antibiotic therapy can be empirical or targeted by means of CT guided fine-needle aspiration. CT-guided fine needle aspiration is not routinely recommended because combination of clinical and radiological data seems more accurate. CT guided fine-needle aspiration can be a reasonable approach in the case of no clinical improvement.
### Table 2. Scoring system of acute pancreatitis gravity.

**Ranson’s criteria**

**A:**
- Age in years; white blood cell count; blood glucose; serum AST; serum ALT
- Serum calcium; hematocrit fall; hypoxemia; BUN increased after IV fluid hydration; base deficit; sequestration of fluids

- If the score ≥3, severe pancreatitis likely
- If the score <3, severe pancreatitis is unlikely

**APACHE II score**

**C:**
- 1. AaDO2 or PaO2 (depending on FiO2);
- 2. Temperature (rectal);
- 3. Mean arterial pressure;
- 4. pH arterial;
- 5. Heart rate;
- 6. Respiratory rate;
- 7. Sodium (serum);
- 8. Potassium (serum);
- 9. Creatinine;
- 10. Hematocrit;
- 11. White blood cell count;
- 12. Glasgow Coma Scale

<table>
<thead>
<tr>
<th>Score</th>
<th>Mortality</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;8</td>
<td>&lt;4%</td>
</tr>
<tr>
<td>≥8</td>
<td>11-18%</td>
</tr>
</tbody>
</table>

**BUPAP score**

**D:**
- BUN >25 mg/dL; impaired mental status; ≥2 SIRS criteria; age >60; pleural effusion present

- BUPAP score of 0 <1% risk of mortality
- BUPAP score >5 22% risk of mortality

**Harmless acute pancreatitis score**

**E:**
- Rebound tenderness and/or guarding; normal hematocrit; normal serum creatinine

If all three are present it is possible to predict with 98% accuracy a mild course

**Organ failure based score includes:**
- Goris multiple organ failure score
- Bernard score
- SOFA score (sequential organ failure assessment)
- Marshall organ dysfunction score

**CT severity index**

The modified CT severity index is an extension of the original CT severity index that was developed by Balthazar and colleagues in 1994 for distinguishing mild, moderate and severe forms of acute pancreatitis. Scores are generated by estimating pancreatic inflammation and necrosis to give a score out of 10. The finding of necrotizing pancreatitis on CT abdomen may modify the therapeutic approach to acute pancreatitis.

AST, aspartate aminotransferase; ALT, alanine aminotransferase; BUN, blood urea nitrogen; SIRS, systemic inflammatory response syndrome; HAPS, harmless acute pancreatitis score; CT, computed tomography.
The management of the patient with acute pancreatitis after several weeks from onset in the absence of clear signs and symptoms of infected necrosis.\textsuperscript{132} The antibiotics of choice are those capable of penetrating necrosis (carbapenems, quinolones, metronidazole). It is not recommended the routine administration of antifungal drugs as prophylaxis or therapy.\textsuperscript{133}

Other therapies Hypoxia should be corrected with supplements of oxygen and, if necessary, with mechanical ventilation, invasive and not. Hypoxia may be associated with acute respiratory distress syndrome (ARDS), pleural effusion, aggressive fluid resuscitation or superimposed pulmonary infections. Hyperglycemia may be due to a pre-existing diabetes mellitus, pancreatic disease, stress or artificial nutrition and should be corrected with insulin therapy with target blood glucose of 140-180 mg/dL. The use of protease inhibitors, in the normal practice, has no clear role and there is no evidence to support it.\textsuperscript{134}

Treatment of complications The presence of local complications or organ failure sustained beyond 48-72 h requires the transfer in a reference center for the treatment of AP.

Infectious complications The probability of developing an infected necrosis is independent of the degree of necrosis and is the main predictor of mortality. Unlike previously believed, the infection may be an early complication of necrosis and Gram-negative intestinal bacteria are frequently involved. The use of antibiotics has expanded in the last decades. There are no indications of efficacy for probiotics. If necessary, the timing of surgery must be carefully assessed in each case by a multidisciplinary team (gastroenterologists and surgeons). In case of stable patients, necrosectomy should be deferred over the 4th week, regardless of the location and extent of necrosis. If the patient remains unstable despite antibiotic therapy, necrosectomy (percutaneous/endoscopic/laparoscopic or laparotomy surgery; if possible a minimally invasive approach is always preferred) is required. However, it would be better to postpone surgery as much as possible so that necrosis can be organized.\textsuperscript{132}

Abdominal compartment syndrome Abdominal compartment syndrome is defined as a persistent increase of intra-abdominal pressure >20 mmHg associated with the emergence of a new organ failure. It may be due to ascites, intestinal obstruction, peripancreatic inflammation and intensive hydration. The intra-abdominal pressure is measured in the bladder. The treatment of compartment syndrome can be surgical or medical.\textsuperscript{135} Medical therapy includes viscera detention through naso-jejunal tube and rectal probe, prokinetic, endoscopic decompression, ultrafiltration or diuretics in case of fluid accumulation, analgesia and sedation until neuromuscular blockade to decrease the muscle tone of the abdominal wall. Surgical therapy includes paracentesis, midline or subcostal laparostomy and subcutaneous fasciotomy. Surgical decompression can be lifesaving.

Pancreatic duct disruption Pancreatic duct disruption produces pleural effusion, ascites and extension of the collections. Symptoms may include shortness of breath, abdominal pain, vomiting and dyspepsia. A focal disconnection can be treated with stents placed endoscopically. If it occurs in the context of a large necrotic area, it requires a specialized surgical approach.\textsuperscript{136}

Pseudocyst Asymptomatic pseudocysts do not require therapy regardless of the location and size. The symptomatic pseudocyst is an indication for surgery.

Thrombosis of splanchnic vessels The porto-spleno-mesenteric vein thrombosis occurs in 50\% of patients with necrotizing pancreatitis and is rare in the absence of necrosis. Treatment consists in anticoagulants, which seem safe but ineffective.\textsuperscript{137}

Pseudoaneurysms Pseudoaneurysms are rare but they can lead to fatal consequences. They are responsible for severe gastrointestinal bleeding and sudden reductions in hematocrit. The treatment of choice is transcatheter arterial embolization. In case of failure the surgical approach is necessary.

Systemic complications They are represented by the following clinical conditions: decompensated diabetes mellitus, exacerbation of preexisting comorbidities, extrapancreatic infections, alcohol withdrawal, respiratory failure due to ARDS, atelectasis or pleural effusion.

Urgent surgical indication in acute pancreatitis Urgent surgical indications in acute pancreatitis are the following two: i) the biliary obstruction associated with cholangitis unresolved by ERCP; ii) infected
necrosis with clinical deterioration associated with persistent signs of infection (>48-72 h). Postponing surgery in conditions of stability (after at least four weeks) is demonstrated to be beneficial.

The conditions in which surgery may be necessary are: biliary obstruction with cholangitis; necrosectomy of infected necrosis; cholecystectomy in case of gallstone pancreatitis although already performed sphincterotomy (indicated at the admission); abdominal compartment syndrome unresponsive to conservative therapy; bleeding from pseudoaneurysms; intestinal ischemia; ductal disruption; fistula of the colon; symptomatic pseudocyst; ongoing gastric outlet, intestinal or biliary obstruction due to mass effect. It should be noted that in case of surgery for abdominal compartment syndrome, bleeding or intestinal ischemia, necrosectomy of sterile necrosis is not recommended, because it can cause necrosis infection.

**Therapy of predisposing factors**

**Gallstones**

In the case of choledocholithiasis with obstruction of the bile duct associated with pancreatitis and cholangitis, an urgent ERCP with sphincterotomy (within 24-72 h) is indicated. This indication is not valid if there are gallstones without cholangitis. ERCP should be considered, in a stable patient with clinical signs of obstruction. To reduce the risk of post-ERCP pancreatitis, pancreatic duct stents and non-steroidal anti-inflammatory drugs (NSAIDs) given rectally have been found to be useful, especially in patients at high risk of severe pancreatitis. Probably NSAIDs should be administered before the procedure (indomethacin 100 mg or diclofenac 100 mg).

ERCP is also indicated in case of residual common bile duct stones after cholecystectomy. ERCP indications for acute biliary pancreatitis are: pancreatitis with biliary obstruction signs associated with fever (urgent-within 24-72 h); pancreatitis with biliary persistent signs of biliary obstruction (election-timing undefined).

In gallstone-associated pancreatitis, cholecystectomy should be performed (even in those who underwent ERCP) with a timing based on the pancreatitis severity: during the same hospitalization in case of mild pancreatitis; later for other grades of severity. Surgery should be performed after resolution of the acute inflammatory phase or after stabilization and organization of necrosis. It can be performed during the intervention of necrosectomy if indicated.

Cholecystectomy may be performed after two episodes of acute idiopathic pancreatitis if there is an increase in transaminases and cholestasis markers (suspected microlithiasis).

Sphincterotomy, that reduces the risk of pancreatitis recurrence, may be adequate in patients with multiple comorbidities or in the frail elderly, in which the operative risk is high.

**Alcoholism**

It is advisable to prevent alcohol withdrawal syndrome and reintegration of thiamine.

**Hypertriglyceridemia**

The levels of plasma triglycerides >1000 mg/dL should be considered as the cause for acute pancreatitis. In this case, apheresis, insulin and heparin (even combined) are suggested in addition to supportive therapy for pancreatitis early treatment with fibrates.

**Hypercalcemia**

Causes of hypercalcemia should be identified and treated.

**The management of the patient with pancreatitis: rationale and objectives**

Acute pancreatitis is a dynamic clinical entity whose severity may vary during hospital stay. Its management may require the cooperation of many specialties (emergency medicine, internal medicine, gastroenterology, surgery, anesthesiologists) in order to ensure the right setting for the right patient, so that each possible complication may be treated in the best way. Therefore, a common language is of pivotal importance to clarify diagnosis, severity and complications and to exactly stratify the patients with the aim of guaranteeing a standard of care, under the therapeutic and diagnostic profile.

**The management of the patient with pancreatitis: methodology**

In order to provide evidence-based recommendations on the management of patients with acute pancreatitis, first of all we verified the existence of guidelines (GL) on this topic.

With this aim we performed a bibliographic search on this guidelines database: i) Scottish Intercollegiate Guidelines Network (SIGN); ii) Institute for Clinical Systematic Improvement (ICSI); iii) National Institute for Health and Clinical Excellence (NICE) (NHS evidence); iv) National Guideline Cleringhouse; v) Canadian Medical Association, CMA infobase; vi) New Zealand Guidelines Group; vii) National System Guidelines; viii) Clinical Practice Guidelines Portal; ix) eGuidelines.

The search has been independently performed by four authors using the following key words: pancreatitis or pancreas where the database included the pos-
sibility of search, and manually examining the guidelines regarding the digestive system disease or biliary tract disease in the other cases. Given the poor result, we conducted further research using both Google search engine and the general database Medline, PubMed using the following search strategies: i) in the first case: pancreatitis AND guidelines; ii) in the second case: pancreatitis [MeSH Terms] and Guideline as limit and pancreatitis [MeSH Terms] and Consensus Development Conference as filter.

The results have been analyzed by each author and then discussed. Thus we (so) realized that most studies analyzed acute pancreatitis guidelines with the AGREE method, considering 2008 as a time limit of bibliography. We therefore decided to analyze with the AGREE method (Appraisal of Guidelines, Research and Evaluation II) only the GL produced after 2008. The AGREE method evaluates the respect of 23 items, collected in 6 domains, that consider explanation of the purpose, clarity, involvement of all stakeholders, applicability, editorial independence and two final considerations on global evaluation. Each author evaluates the respect of each item with a score going from 1 (total disagreement) to 7 (total agreement); scores given by each author are summed within each domain and related to maximum and minimum score that is possible to obtain in relation to the number of included item and the number of authors.

**Clinical approach to patients with pancreatitis**

The management of pancreatitis consisted of four steps: i) diagnosis and evaluation etiology of pancreatitis; ii) initial evaluation and prognostic assessment; iii) initial management: fluid therapy, role of antibiotics and nutritional support; iv) management of complications.

**Diagnosis and evaluation etiology of pancreatitis (Grade 1B, strong agreement)**

The diagnosis of AP is possible when two of the following three criteria are present: i) abdominal pain consistent with the disease; ii) serum amylase and/or lipase greater than three times the upper limit of normal. Lipase is more specific. Possible modifications of the limit are admitted for diabetic patients who normally have a higher average value; iii) characteristic findings from abdominal imaging.

Concerning etiology, since the most frequent cause of AP is cholelithiasis and alcohol intake, an abdomen ultrasound examination should always be performed in the patient referred to the hospital for AP in addition to blood tests on cholestasis (ALT >150 U/L in the first 48 h represents a positive predictive value >85%).

In the absence of clinical history of alcohol or gallstones, a careful medical history should be performed in order to identify previous unknown episodes of AP, use of drugs, abdominal trauma, infections, recent invasive procedures such as ERCP and other laboratory parameters, such as serum triglyceride (considered the etiology if >1000 mg/dL) and calcium.

The tumor etiology must always be regarded as possible cause of AP especially given the decreasing age average of onset of pancreatic adenocarcinoma or other malignant or benign tumors.

In case of idiopathic pancreatitis diagnosis there is no consensus about referring patients to a center of excellence. The guidelines IAP/APA recommend EUS that can point out micro lithiasis <3 mm, neoplasia or chronic pancreatitis and in case of negative examina-
The update of 2002 IAP/APA guidelines is based on an evidence-based approach to AP management. It includes the revised Atlanta criteria of 1998, and divides the clinical condition into four scenarios identifying what are the techniques of image to be taken for each scenario. The focus of the guidelines is on the diagnosis and subsequent assessment of patients with AP. It is definitely useful because it adds information on diagnostic tools in different AP onset.

Guideline for the practice of endoscopy, developed by the American Society for Gastrointestinal Endoscopy by using an evidence based methodology.

It represents the 2013 review of the original criteria of 1998, and divides the clinical condition in four scenarios identifying what are the techniques of image to be taken for each scenario. The focus of the guidelines is on the diagnosis and subsequent assessment of patients with AP. It is definitely useful because it adds information on diagnostic tools in different AP onset.

Table 3. Guidelines found on databases of guidelines, PubMed and Google.

<table>
<thead>
<tr>
<th>Databases of guidelines</th>
<th>Guidelines found on databases of guidelines, PubMed and Google.</th>
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</thead>
<tbody>
<tr>
<td>IAP/APA evidence-based guidelines for the management of acute pancreatitis. Pancreatology 2013;2013:e1-e15</td>
<td>The update of 2002 IAP/APA guidelines is based on an evidence-based approach to AP management. It includes the modified Atlanta criteria and the 38 recommendations regarding 12 aspects of AP. This guideline is graded with the GRADE method.</td>
</tr>
<tr>
<td>Classification of acute pancreatitis-2012: revision of the Atlanta classification and definitions by international consensus. Gut 2013;62:102-11</td>
<td>It is the revision of the 1998 classification criteria of Atlanta. It is a consensus document made through an iterative process and web-based of a working group consisting of 11 national and international pancreatic society. The main goal is to provide unique classification criteria. It includes a clinical assessment of the severity of the AP proposing objective criteria (and therefore measurable and quantifiable) to describe the complications of local AP. It is not a guideline management of AP, neither gives therapeutic indications. It is included in two more recent guidelines, ACG and IAP/APA.</td>
</tr>
<tr>
<td>Clinical pathways for acute pancreatitis. Recommendations for early multidisciplinary management. Med intensiva 2012;36:351-357</td>
<td>It does not represent a true guideline. It is based on the recommendations address to 2005 guidelines SEMICYUC ICU. It was proposed by Petrov et al. in 2010 as classification criteria.</td>
</tr>
<tr>
<td>Practical guidelines for acute pancreatitis (official guidelines of the Italian association for the study of the pancreas regarding the medical, endoscopic and surgical management of acute pancreatitis). Pancreatology 2010;10:523-535</td>
<td>The document is not a real guideline but a review of the existing guidelines on the management of acute pancreatitis using ADAPTE methodology. The literature search was conducted looking all guidelines 1996-2007 in pubmed cochrane library and other databases. The guidelines selected (9 from 21) were evaluated using the AGREE instrument and the working group concluded that all 9 guidelines met criteria to answer specific clinical questions about acute pancreatitis.</td>
</tr>
<tr>
<td>AGA institute medical position statement on acute pancreatitis. Gastroenterology 2007;132:2019-2021</td>
<td>These recommendations are written in order to provide evidence based on prospective.</td>
</tr>
<tr>
<td>Practice guidelines in acute pancreatitis. American journal of gastroenterology 2006</td>
<td>It considers various aspects of acute pancreatitis such as diagnoses, assessment of risk factors and severity, supportive care therapy, fluids therapy, the indication of ICU, nutrition, antibiotics, necrosis treatment.</td>
</tr>
<tr>
<td>JPN guidelines for the management of acute pancreatitis. J Hepatobiliary Pancreat Surg 2006;13:56-60</td>
<td>It contains several elements which are inconsistent with the latest guidelines, especially regarding therapeutic aspects.</td>
</tr>
</tbody>
</table>

AP, acute pancreatitis; ACR, American College of Radiology; IAP/APA, International Association of Pancreatology/American Pancreatic Association; ACG, American College of Gastroenterology; ICU, Intensive Care Unit.
The management of the patient with acute pancreatitis

Hemodynamic status should be assessed immediately upon presentation and resuscitative measures begun as needed. Risk assessment should be performed to stratify patients into higher- and lower-risk categories to assist triage, such as admission to an intensive care setting (strong recommendation, moderate quality of evidence).

The best score to predict severe acute pancreatitis on admission and at 48 hours is the presence of SIRS. Persistent (>48 h) SIRS is associated with multi-organ failure and mortality (25%) in acute pancreatitis (Grade 2B, weak agreement).

<table>
<thead>
<tr>
<th>Domain</th>
<th>IAP/APA</th>
<th>ACG</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>94%</td>
<td>92%</td>
</tr>
<tr>
<td>2</td>
<td>58%</td>
<td>47%</td>
</tr>
<tr>
<td>3</td>
<td>91%</td>
<td>67%</td>
</tr>
<tr>
<td>4</td>
<td>97%</td>
<td>100%</td>
</tr>
<tr>
<td>5</td>
<td>24%</td>
<td>0</td>
</tr>
<tr>
<td>6</td>
<td>100%</td>
<td>100%</td>
</tr>
</tbody>
</table>

IAP/APA, International Association of Pancreatology/American Pancreatic Association; ACG, American College of Gastroenterology.

The best strategy to predict outcome of acute pancreatitis during admission is a 3-dimension approach combining: i) host risk factors (e.g., age, co-morbidity, body mass index); ii) clinical risk stratification (e.g., persistent SIRS); iii) monitoring response to initial

### Table 5. Difference between the two main guidelines.

<table>
<thead>
<tr>
<th>If diagnosis of idiopathic pancreatitis</th>
<th>Guidelines of ACG</th>
<th>Guidelines of IAP/APA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Refer patient to centers of expertise; Genetic testing may be considered in young patients (&lt;30 years old) if no cause is evident and a family history of pancreatic disease is present (conditional recommendation, low quality of evidence)</td>
<td>If EUS is negative, secretin-stimulated-MRCP can be used to identify rare morphologic abnormalities If etiology remains unidentified, genetic counseling should be considered, especially after a second attack of AP (Grade 2C, weak agreement)</td>
<td></td>
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</tbody>
</table>

| Definition of specialist center | None | High volume center with up-to-date intensive care facilities including options for organ replacement therapy, and with daily access to interventional radiology, interventional endoscopy with EUS and ERCP assistance as well as surgical expertise in managing necrotizing pancreatitis (Grade 2C, weak agreement) |

| Prevent severe post-ERCP pancreatitis in high-risk patients | Pancreatic duct stents and/or post-procedure rectal non-steroidal anti-inflammatory drug suppositories (conditional recommendation, moderate quality of evidence) | None |

| The role of antibiotics in acute pancreatitis | Extrapancreatic infection, such as cholangitis, catheter-acquired infections, bacteremia, urinary tract infections, pneumonia; (strong recommendation, high quality of evidence) Infected necrosis (strong recommendation, low quality of evidence) | None |

| Consider and treat ACS | None | ACS is defined as a sustained intra-abdominal pressure >20 mmHg (via the bladder with a maximal instillation volume of 25 mL of sterile saline) that is associated with new onset organ failure. It should be considered in mechanically ventilated patients with severe acute pancreatitis, especially in case of clinical deterioration. Medical treatment of ACS consists in decreasing intra-abdominal pressure |

ACG, American College of Gastroenterology; IAP/APA, International Association of Pancreatology/American Pancreatic Association; EUS, endoscopic ultrasonography; MRCP, magnetic resonance cholangiopancreatography; AP, acute pancreatitis; ERCP, endoscopic retrograde cholangiopancreatography; NSAID, non-steroidal anti-inflammatory drug; ACS, abdominal compartment syndrome.
therapy (e.g., persistent SIRS, blood urea nitrogen, creatinine) (Grade 2B, strong agreement).

Because of the absence of any available test to determine severity, assessing early fluid losses, hypovolemic shock, and organ dysfunction symptoms is crucial for any clinicians (Figure 2) to predict which patients with AP will develop severe disease.

Optimal timing for initial CT assessment is at least 72/96 h after onset of symptoms (Grade 1C, strong agreement). Early CT may be useful to rule out bowel ischemia or intra-abdominal perforations in patients presenting with both acute pancreatitis and acute abdomen.

Indications for follow-up scanning (CT/MR) in acute pancreatitis are the following (Grade 1C, strong agreement): i) lack of clinical improvement; ii) clinical deterioration; iii) invasive intervention is considered, and only a portal venous phase (monophasic) is generally sufficient.

The optimal CT and MR protocol to detect necrosis should be applied.

Indications for ERCP and sphincterotomy are: patients with biliary pancreatitis and cholangitis (Grade 1B, strong agreement) and biliary pancreatitis with common bile duct obstruction (probably indicated) (Grade 1C, strong agreement).

The optimal timing of ERCP in patients with biliary pancreatitis without cholangitis is not known.

The urgent ERCP (<24 h) is required in patients with acute cholangitis (Grade 2C, strong agreement). As the exact timing of early ERCP (24/72 h) is not known, it is reasonable to await spontaneous improvement of biliary obstruction for 24/48 h. ERCP should be performed as soon as possible in patients with cholangitis.

The role of MRCP and EUS in biliary pancreatitis is to prevent some ERCPs that would otherwise be performed for suspected common bile duct stones in patients with biliary pancreatitis who do not have cholangitis, without influencing the clinical course. EUS is superior to MRCP in excluding the presence of small (<5 mm) gallstones. MRCP is less invasive, less operator-dependent and probably more widely available than EUS. Therefore, there is no clear superiority for either MRCP or EUS in clinical practice (Grade 2C, strong agreement).

**Initial management: fluid therapy, role of antibiotics and nutritional support**

Initial management includes patients’ stabilization and their allocation according to AP severity.

**Fluid therapy**

Early fluid resuscitation is associated with decreased rates of persistent SIRS and organ failure (Grade 1C, strong agreement).

Ringer’s lactate (Grade 1B, strong agreement) at the rate of 5-10 mL/kg/h (250-500 mL/h) (Grade 1B, weak agreement) is the best fluid to use for initial fluid resuscitation in acute pancreatitis, unless cardiovascular and/or renal comorbidities exist. Early aggressive intravenous hydration is most beneficial during the first 12-24 h, and may have little benefit beyond. A more rapid repletion (bolus) may be necessary in a patient presented with hypotension and tachycardia for severe volume depletion. Fluid requirements should be reassessed at frequent intervals within 6 hours from admission and for the following 24-48 h. The goal of aggressive hydration should be to decrease the blood urea nitrogen.

To assess the response to fluid therapy, one or more of the following points should be considered (Grade 2B, weak agreement): i) non-invasive measurement: heart rate <120/min; mean arterial pressure between 65 and 85 mmHg (8.7-11.3 kPa); and urinary output >0.5/1 mL/kg/h; ii) invasive clinical targets: stroke volume variation and intrathoracic blood volume determination; iii) biochemical targets of hemocrit 35/44%.

Indications for admission to an intensive care unit in acute pancreatitis are showed in Figure 3 (Grade 1C, strong agreement).

**Antibiotics therapy**

Antibiotics should be given for an extrapancreatic infection, such as cholangitis, catheter-acquired infections, bacteremia, urinary tract infections, pneumonia. In patients with infected necrosis, antibiotics known to penetrate pancreatic necrosis, such as carbapenems, quinolones, and metronidazole, may be useful in delaying or avoiding intervention, thus decreasing morbidity and mortality.

Routine administration of antifungal agents along with prophylactic or therapeutic antibiotics is not recommended (conditional recommendation, low quality of evidence).

**Nutritional support**

In patients with predicted mild pancreatitis, oral feeding can be restarted once abdominal pain is resolved and inflammatory markers improved (Grade 2B, strong agreement). Feeding can be started with a full solid diet without waiting for normalization of lipase levels before restarting oral feeding.

In patients with predicted severe acute pancreatitis who require nutritional support enteral tube feeding should be the primary therapy and can be administered via either the nasojejunal or nasogastric route (Grade 1B, strong agreement). If nasojejunal tube feeding is not tolerated and nutritional support is required, parenteral nutrition can be administered as second-line therapy (Grade 2C,
The management of the patient with acute pancreatitis

Management of complications

The indications for referral patients with AP to a specialist center (high volume center with up-to-date intensive care facilities including options for organ replacement therapy, (and with) daily access to interventional radiology, interventional endoscopy with EUS and ERCP assistance as well as surgical expertise in managing necrotizing pancreatitis) is severe acute pancreatitis and patients with local complication who may need interventional radiologic, endoscopic, or surgical intervention.

<table>
<thead>
<tr>
<th>Patient characteristics</th>
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<tbody>
<tr>
<td>Age &gt;55 years</td>
</tr>
<tr>
<td>Obesity (BMI &gt;30 kg/m²)</td>
</tr>
<tr>
<td>Altered mental status</td>
</tr>
<tr>
<td>Comorbid disease</td>
</tr>
</tbody>
</table>

The systemic inflammatory response syndrome

Presence of >2 of the following criteria:
- pulse > 90 beats/min
- respiration > 20/min or PaCO₂ >32 mmHg
- temperature > 38°C or < 36°C
- WBC count > 12,000 or < 4000 cells/mm³ or > 10% immature
- neutrophils (bands)

Laboratory findings

| BUN >20 mg/dL |
| Rising BUN   |
| HCT > 44%    |
| Rising HCT   |
| Elevated creatinine |

Radiology findings

| Pleural effusions |
| Pulmonary infiltrates |
| Multiple or extensive extrapancreatic collections |

Figure 2. Clinical findings associated with a severe course for initial risk assessment (the presence of organ failure and/or pancreatic necrosis, defines severe acute pancreatitis). BMI, body mass index; WBC, white blood cell; BUN, blood urea nitrogen; HCT, hematocrit.

Indications for intervention in necrotizing pancreatitis (either radiological, endoscopic or surgical)

Common indications for intervention are (Grade 1C, strong agreement): i) clinical suspicion or documented infected necrotizing pancreatitis with clinical deterioration, preferably when the necrosis has become walled-off; ii) presence of organ failure for several weeks after the onset of acute pancreatitis also in the absence of documented infected necrotizing pancreatitis.

Less common indications for intervention are (Grade 1C, strong agreement): i) abdominal compartment syndrome; ii) acute bleeding; iii) bowel ischemia; iv) gastric, intestinal, or biliary obstruction due to mass effect from large walled-off necrosis (arbitrarily >4/8 weeks after onset of pancreatitis).
Routine percutaneous needle aspiration of peripancreatic collections (to document infected necrotizing) is not indicated because clinical signs (i.e., persistent fever, increasing inflammatory markers) and imaging signs (i.e., gas in peripancreatic collections) are accurate predictors of infected necrosis in the majority of patients (Grade 1C, strong agreement).

**Indications for intervention (either radiological, endoscopic or surgical) in sterile necrotizing pancreatitis**

Indications for intervention (either radiological, endoscopic or surgical) in sterile necrotizing pancreatitis are (Grade 1C, strong agreement): i) gastric outlet, intestinal, or biliary obstruction due to mass effect of organized necrosis (i.e., arbitrarily >4-8 weeks after onset of acute pancreatitis); ii) persistent symptoms (e.g., pain, persistent malaise) in patients with organized necrosis without signs of infection (i.e., arbitrarily >8 weeks after onset of acute pancreatitis); iii) disconnected duct syndrome (i.e., full transection of the pancreatic duct in the presence of pancreatic necrosis) with persisting symptomatic (e.g., pain, obstruction) collection(s) with necrosis without signs of infections (i.e., arbitrarily >8 weeks after onset of acute pancreatitis).

The presence of asymptomatic pseudocysts and pancreatic and/or extrapancreatic necrosis does not warrant intervention, regardless of size, location, and/or extension.

**Timing of cholecystectomy**

After mild biliary pancreatitis cholecystectomy appears safe and is recommended during index admission. Interval cholecystectomy after mild biliary pancreatitis is associated with a substantial risk of readmission for recurrent biliary events, especially recurrent biliary pancreatitis (Grade 1C, strong agreement).

After severe biliary pancreatitis with peripancreatic collections, cholecystectomy should be delayed until the collections resolve. In case they persist beyond 6 weeks, cholecystectomy can be performed safely (Grade 2C, strong agreement).

Cholecystectomy is also indicated in patients with biliary pancreatitis and previous history of sphincterotomy, since neither ERCP nor sphincterotomy prevent biliary colic and cholecystitis (Grade 2B, strong agreement).

**Conclusions**

The two guidelines considered above emphasize the latest clinical evidence in the context of diagnosis, etiology, and therapeutic approach to the patient with AP.

Their diffusion and application in the clinical setting can uniform clinical practice and improve patient outcomes.

In addition, the new definitions of severity and complications of AP used in these guidelines give us a universal language to properly stratify the patient for correct allocation.

For this reason, it is important that the internist knows diagnosis, management and treatment of this complex and multidisciplinary disease.

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**One or more of the following parameters (106)**

1) Pulse <40 or >150 beats/min;

2) Systolic arterial pressure <80 mmHg (<10.7 kPa) or mean arterial pressure <60 mmHg (<8.0 kPa) or diastolic arterial pressure >120 mmHg (>16 kPa).

3) Respiratory rate >35 breaths/min.

4) Serum sodium <110 mmol/L or >170 mmol/L.

5) Serum potassium <2.0 mmol/L or >7.0 mmol/L.

6) PaO₂ <50 mmHg (<6.7 kPa).

7) pH <7.1 or >7.7.

8) Serum glucose >800 mg/dL (>44.4 mmol/L).

9) Serum calcium >15 mg/dL (>3.75 mmol/L).

10) Anuria.

11) Coma.

Figure 3. Criteria for admission to intensive care unit.
References


108. Lankisch PG, Weber-Dany B, Hebel K, et al. The harm-