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Editor-in-Chief of Artificial Intelligence in Medical Imaging, Professor Xue-Li Chen is an expert in the field of biomedical photonics imaging as well as its application in early detection and accurate diagnosis of gastric cancer. Professor Chen has co-led the development of Cerenkov luminescence endoscope and further explored the application in early detection of clinical gastrointestinal tumors. Professor Chen has also developed the stimulated Raman projection tomography technology which can perform the volumetric imaging of single cells in a label-free manner. Professor Chen has served as the member of SPIE, OSA, IEEE, and as a committee member of the Branch of Contrast Technology in China Medicinal Biotech Association, the Nuclear Medicine Committee of Shaanxi Cancer Association, and the Shaanxi Society of Biomedical Engineering.

Aims and Scope

The primary aim of Artificial Intelligence in Medical Imaging (AIMI, Artif Intell Med Imaging) is to provide scholars and readers from various fields of artificial intelligence in medical imaging with a platform to publish high-quality basic and clinical research articles and communicate their research findings online. AIMI mainly publishes articles reporting research results obtained in the field of artificial intelligence in medical imaging and covering a wide range of topics, including artificial intelligence in radiology, pathology image analysis, endoscopy, molecular imaging, and ultrasonography.

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Acute pancreatitis: A pictorial review of early pancreatic fluid collections

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Abstract

Acute pancreatitis is a common acute inflammatory disease involving the pancreas and peripancreatic tissues or remote organs. The revised Atlanta classification 2012 of acute pancreatitis divides patients into mild, moderately severe and severe groups. Major changes of the classification include acute fluid collection terminology. However, some inappropriate terms of the radiological diagnosis reports in the daily clinical work or available literature may still be found. The aim of this review article is: to present an image-rich overview of different morphologic characteristics of the early-stage (within 4 wk after symptom onset) local complications associated with acute pancreatitis by computed tomography or magnetic resonance imaging; to clarify confusing imaging concepts for pancreatic fluid collections and underline standardised reporting nomenclature; to assist communication among treating physicians; and to facilitate the implications for clinical management decision-making.

Key words: Acute pancreatitis; Computed tomography; Magnetic resonance imaging; Acute peripancreatic fluid collection; Acute necrotic collection; Complication

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Core tip: To our best of knowledge, this is the first pictorial review that determines the spectrum of magnetic resonance imaging features in patients with acute pancreatitis of distinct early acute necrotic collection compared with acute peripancreatic fluid collection.

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INTRODUCTION

Acute pancreatitis is a common digestive disease, which is related to an acute onset of epigastric pain with/without nausea and vomiting. Cholelithiasis, alcoholism and hyperlipidaemia are the most widely recognised etiological factors in acute pancreatitis patients[1]. Clinically, physicians often make the accurate diagnosis based on clinical manifestations and biochemical parameters (sufficiently elevated serum lipase or amylase) for the majority of patients with acute pancreatitis[2]. Indeed, the routine medical imaging for this disease is unwarranted. However, the natural history and consequences of critically ill patients (particularly moderately severe or severe acute pancreatitis) can result in a variety of local complications[3]. These developments thus prompt imaging to detect clinical complications.

Imaging approaches, especially computed tomography (CT) and magnetic resonance imaging (MRI), are valuable in detecting local complications associated with acute pancreatitis in both the early-phase and the late-phase of disease. With the increasing application of the revised Atlanta classification criteria 2012[4], radiologists play a crucial role in relevant imaging diagnosis, scientific research and multidisciplinary team communication. Although this classification updates the definitions of acute pancreatitis and many pancreatitis-associated complications, some inappropriate terms of the radiological diagnosis reports in the daily clinical work or available literature may still be found.

Therefore, the purpose of this pictorial article is: To serve as an image-rich overview of different morphologic characteristics of the early-stage (within 4 wk after symptom onset) local complications associated with acute pancreatitis by CT or MR; to clarify confusing imaging concepts and enable standardised reporting nomenclature; to assist communication among treating physicians; and to facilitate the implications for treatments.

DEFINITION AND DIAGNOSIS OF EARLY-STAGE COLLECTIONS ASSOCIATED WITH ACUTE PANCREATITIS

In general, acute pancreatitis is classically divided into two types: Interstitial oedematous pancreatitis and necrotising pancreatitis. Clinically, the majority of patients with acute pancreatitis are present as interstitial oedematous pancreatitis. They have diffuse or localised enlargement of the pancreas owing to inflammatory oedema. On the other hand, necrotising pancreatitis accounts for 20%-30% of acute pancreatitis patients[5], and it is subdivided into three subtypes on the basis of contrast-enhanced CT according to the new Atlanta classification[6]: (1) Combined pancreatic necrosis and peripancreatic necrosis (most common, approximately 75% of all necrotising pancreatitis); (2) Peripancreatic tissue necrosis alone (less common, with an incidence of approximately 20%)[7]; and (3) Pancreatic parenchymal necrosis alone (rare, with an incidence of only 5%)[8-9]. For radiologists, it is crucial to make a distinction between interstitial oedematous pancreatitis and necrotising pancreatitis in diagnosing acute pancreatitis. The correct diagnosis of imaging type can assist in the recognition of subsequent pancreatitis-related complications and application of the proper terminology.

Currently, there are two diagnostic terms for local complications in the early stage of acute pancreatitis: Acute peripancreatic fluid collections (APFCs) and acute necrotic collections (ANCs)[10]. On the one hand, the characteristics of acute peripancreatic fluid collections include: (1) Arising from interstitial oedematous pancreatitis; (2) Nonencapsulated collections (lack of a well-defined capsule/definable wall); (3) Collection age within the initial 4 wk after symptom onset; (4) Peripancreatic location (surrounding or adjacent to pancreas); and (5) A homogeneous or simple fluid appearance (containing pure fluid) (Figure 1)[10].

On the other hand, the characteristics of acute necrotic collections include: (1) Occurring only in the setting of necrotising pancreatitis; (2) Collections without an encapsulating capsule or over time with poorly organised wall, (3) Collection age within the first 4 wk of this disease; (4) Peripancreatic and pancreatic different
Figure 1  Schematic diagram of acute peripancreatic fluid collections and magnetic resonance imaging of a patient. A: Schematic diagram of acute peripancreatic fluid collections within 4 wk of onset of interstitial oedematous pancreatitis. The term “acute peripancreatic fluid collections” applies only to interstitial oedematous pancreatitis patients; B: A 66-year-old woman with interstitial oedematous pancreatitis. Magnetic resonance imaging axial T2WI image shows a homogeneous fluid finding (arrows) around the pancreas. APFCs: Acute peripancreatic fluid collections; P: Pancreas.

locations (surrounding the pancreas or intrapancreatic extension or both); and (5) A heterogeneous appearance due to containing variable amounts of inflammatory fluid and liquefied or nonliquefied necrotic debris (Figure 2)[7-10].

LOCATION, SHAPE, SIZE OF APFCs/ANCs

On CT/MRI, acute peripancreatic fluid collections are predominantly localised in peripancreatic areas (e.g., the lesser sac) and the retroperitoneal spaces (e.g., left anterior pararenal space) or peripancreatic fascial planes. They exhibit variable shape and size but mostly present as a uniform linear or strip-shaped liquid appearance[3-6]. In addition, the volume of APFCs is relatively smaller due to fluid generally confined by a simple retroperitoneum space and/or normal peripancreatic fascial planes (Figure 3).

On the other side, acute necrotic collections often break through the limitation of the interfascial planes and can affect multiple retroperitoneal spaces, interfascial planes, subperitoneal spaces and other abdominal spaces. In fact, they are most frequently situated in the lesser sac and the anterior pararenal spaces, followed by transverse mesocolon, mesenteric root, and thereafter, gastrohepatic, gastrosplenic and gastrocolic ligaments[11]. Furthermore, these collections may additionally involve the remote regions, such as the pelvic sidewalls and mediastinum. Thus, acute necrotic collections are generally numerous (multiple), irregular and loculated, and the volume of effusion often appears larger than that of acute peripancreatic fluid collections (Figure 4)[9-11].

DENSITY/SIGNAL INTENSITY, ENHANCEMENT CHARACTERISTICS OF APFCs/ANCs

Acute peripancreatic fluid collections have homogeneous fluid appearances. They are uniformly hypoattenuating on CT and T1 hypointense and T2 hyperintense on MRI. After intravenous contrast-material administration, acute peripancreatic fluid collections are not enhancing owing to the pure fluid nature[8-10].

In contrast, acute necrotic collections are heterogeneous (Figure 5). There are relatively hyperdense materials (necrotic fragments of pancreas) and/or markedly hypodense fat globules (peripancreatic fat) among hypodense fluid on CT[6-8]. Similarly, there are varying degrees of round, patchy, strip-shaped T2-hypointense components (adipose fragments or necrotic pancreatic tissue) among T2-hyperintense fluid on MR images[3-6]. During contrast-enhanced CT/MRI, the internal necrotic debris or trapped fat within acute necrotic collections often does not show enhancement (Figure 5), while the immature, fibrous granulation tissue wall of these collections may be detectable as slight to confluent enhancement (Figure 6).
Figure 2 Schematic diagram of acute necrotic collections and computed tomography of a patient. A: Schematic diagram of acute necrotic collections within 4 wk of onset of necrotizing pancreatitis. The term “acute necrotic collections” is diagnosed only in necrotizing pancreatitis patients; B: A 56-year-old man with necrotizing pancreatitis. Axial contrast-enhanced computed tomography image in the venous phase shows a large area of necrosis (asterisks) in the pancreatic body and tail; therefore, a lesser omental sac collection (arrows) should be diagnosed as acute necrotic collections. ANCs: Acute necrotic collections; P: Pancreas; N: Necrosis.

Figure 3 A 40-year-old woman with interstitial oedematous pancreatitis. Magnetic resonance imaging axial T2WI image shows uniform linear liquid hyperintense (acute peripancreatic fluid collections) in the left pararenal anterior spaces (arrowheads) and retromesenteric plane (arrows).

Figure 4 A 53-year-old man with necrotizing pancreatitis. Axial contrast-enhanced computed tomography image in the venous phase shows extensive heterogeneous collections (acute necrotic collections) in the left pararenal anterior spaces (arrows) and the subperitoneal spaces/transverse mesentery areas (asterisks) as well as greater omentum zones (arrowheads).

SECONDARY OR CONCOMITANT CT/MRI FINDINGS OF APFCs/ANCs

In general, (peri)pancreatic fluid collections may also be associated with a variety of complications, which can make the clinical condition more complex. After conservative treatments, acute peripancreatic fluid collections are often absorbed quickly with rare follow-up complications. Most of these patients are discharged within 1-2 wk after admission.1-9.
Clinically, necrotising pancreatitis with acute necrotic collection is mainly seen in patients with moderate severe acute pancreatitis and severe acute pancreatitis, with a longer disease course (lasting several weeks or months). Consequently, secondary infectious complications are more likely to occur in acute necrotic collections, compared with acute peripancreatic fluid collections. Infection should be suspected when there are secondary clinical signs of sepsis, such as a new occurrence of fever and leucocytosis. On CT images, the sign of multiple extraluminal gas or a gas-fluid level in the peripancreatic zones and retroperitoneal spaces is highly suggestive of acute necrotic collections complicated by infection. If clinical manifestations are concordant or needle-guided aspiration confirms the development of infection, then these collections should be classified as infected acute necrotic collections. In this setting, percutaneous aspiration or drain insertion can be performed for the treatment of an infected collection. Moreover, when necrotising pancreatitis affects a large area of intraparenchymal pancreas, it involves the main pancreatic duct (necrosis of the pancreatic duct). Over time, the pancreatic duct rupture or disrupted integrity of the pancreatic duct accompanied with the intrapancreatic acute necrotic collections (liquefied pancreatic tissue) can result in the formation of “disconnected duct syndrome”. As for this condition, a collection communication with main pancreatic duct is usually evident on MRI and MR cholangiopancreatography. This syndrome may alter treatment but does not affect acute necrotic collection classification, and these patients often require surgical management for a complete recovery.
**Figure 7** A 59-year-old man with necrotizing pancreatitis complicating infection. Axial contrast-enhanced computed tomography image in the venous phase shows multiple extraluminal gas bubbles (arrowheads) in the peripancreatic and the retroperitoneal spaces, consistent with a pathognomonic sign of the infected necrosis.

**Figure 8** Schematic diagram of disconnected pancreatic duct syndrome and medical images of patients. A: Schematic diagram of disconnected pancreatic duct syndrome. The pancreatic duct rupture and interruption (arrows) resulting from a wide range of liquefied pancreatic body tissue is seen; B: A 60-year-old man with necrotizing pancreatitis. Axial contrast-enhanced computed tomography image in the venous phase shows extensive parenchymal transmural necrosis (arrows) in the region of neck and body of the pancreas. Disconnected main pancreatic duct was proved at surgery; C: A 57-year-old woman with necrotizing pancreatitis. Magnetic resonance imaging axial T2WI image shows a majority of liquefied necroses (arrowheads) in the pancreatic body; D: The main pancreatic duct (arrow) of the tail of pancreas is interrupted (arrowhead) by the mentioned-above lesion. P: Pancreas; N: Necrosis.

**TERMINOLOGY MISUSE IN IMAGING REPORTS OF APFCs/ANCs**

Depending on the revised Atlanta classification and our clinical practice, common terminology misuse conditions in the daily imaging reports are summarised as follows: (1) On CT/MRI, no necrosis finding was observed in the pancreatic parenchyma, which was assumed to be “interstitial oedematous pancreatitis.” Then the accumulation of fluid in the peripancreatic regions may be misinterpreted as “acute peripancreatic fluid collection.” However, a pitfall is probably present because we may ignore the presence of small “adipose tissue debris” in peripancreatic collections ([Figure 9](#)). At this point, it should be diagnosed as “necrotising pancreatitis (peripancreatic necrotic type),” due to necrotic adipose fragments around the pancreas. Therefore, the nomenclature of the collection is referred to as an “acute
Figure 9  Schematic diagram of necrotizing pancreatitis and computed tomography image of a patient. A: Schematic diagram of necrotizing pancreatitis (peripancreatic necrotic type). Acute necrotic collection involving peripancreatic fat only (arrows) is seen; B: A 58-year-old woman with necrotizing pancreatitis (peripancreatic necrosis only). Axial contrast-enhanced computed tomography image in the venous phase shows normal pancreatic parenchymal enhancement without definite necrosis. A peripancreatic collection may be misdiagnosed as “acute peripancreatic fluid collection.” However, multiple heterogeneous, nonliquid adipose components (arrows) are revealed among the collection. For this reason, it should be considered as acute necrotic collection. P: Pancreas; APFC: Acute peripancreatic fluid collection; ANC: Acute necrotic collection.

EXISTING PROBLEMS AND PROSPECTS

Although there are many differences between acute peripancreatic fluid collections and acute necrotic collections, in the clinical practice it is sometimes difficult to accurately distinguish acute peripancreatic fluid collections from acute necrotic collections on CT at the early stage of acute pancreatitis (especially within 2 d of symptom onset). The reasons may be related to the following factors: (1) It may allow sufficient time for completed necrosis of the pancreas and/or peripancreatic fat (findings of solid necrotic materials to liquefy over several days). Thus, heterogeneous contents may not be found within the early-phase fluid. This also explains why an early contrast-enhanced CT may underestimate the eventual degree of (peri)pancreatic necrosis. In this setting, further CT studies after an interval of between 5 d and 7 d should be performed[8-12]; and (2) Due to the low contrast resolution of CT, it is difficult to detect a small amount of heterogeneous contents. For this purpose, MRI may be required for this distinction because it is very sensitive to the detection of internal architecture of collections (even a small area of heterogeneous debris)[6-9]. Moreover, MRI diffusion weighted imaging combined with ADC value measurement is helpful for the differential diagnosis of interstitial oedematous pancreatitis and necrotising pancreatitis[4-6]. Whether diffusion weighted imaging is also valuable for the early differential diagnosis of acute peripancreatic fluid collection and acute necrotic collection may become a direction of future research.

CONCLUSION

To sum up, the natural history and consequences of different pancreatic and peripancreatic collections are now better described and understood. The differential
Figure 10  Schematic diagram of necrotizing pancreatitis and magnetic resonance imaging of a patient. A: Schematic diagram of necrotizing pancreatitis (pancreatic parenchymal necrosis alone). The peripancreatic collections are homogeneous; B: A 47-year-old man with necrotizing pancreatitis complicated with haemorrhage. Magnetic resonance imaging axial T1WI image shows peripancreatic homogeneous fluid with greater hyperintense signal. For this reason, it may be misdiagnosed as “acute peripancreatic fluid collection.” However, the necrosis and haemorrhage (arrow) of the body and tail of the pancreas can be indicated. Therefore, the collection should be diagnosed as acute necrotic collection. P: Pancreas, N: Necrosis, H: Haemorrhage; APFC: Acute peripancreatic fluid collection; ANC: Acute necrotic collection.

Figure 11  Schematic diagram of acute necrotizing pancreatitis, and extrapancreatic fluid and computed tomography of a patient. A: Schematic diagram of acute necrotizing pancreatitis and extrapancreatic fluid with extension within the pancreatic parenchyma (arrow); B: A 56-year-old woman with acute necrotizing pancreatitis. Axial contrast-enhanced computed tomography image in the late arterial phase shows a peripancreatic homogeneous collection (arrowheads). It may be misinterpreted as “acute peripancreatic fluid collection.” However, note the peripancreatic fluid extends into the parenchyma of the head and neck of the pancreas (arrow). Therefore, it should be diagnosed as an “acute necrotic collection.” P: Pancreas; APFC: Acute peripancreatic fluid collection; ANC: Acute necrotic collection.

diagnosis of these collections within 4 wk of symptom onset is succinctly summarised in Table 1. The accurate description of pancreatitis-associated collections, including location (pancreatic, peripancreatic, others), the presence of contents (liquid, solid, gas), the thickness of collection wall (thin, thick) and the presence or absence of infectious findings will facilitate the radiologic reports in daily practice. Finally, radiologists should be fully aware of the standardised imaging nomenclature on the basis of associated morphologic descriptions. It is necessary for accurate documentation and reporting of academic research, and it is also important to direct implications of care plans for patients with acute pancreatitis.
Xiao B. Pictorial review of early pancreatic fluid collections

Table 1 Key points of clinical and imaging differential diagnosis between acute peripancreatic fluid collection and acute necrotic collection

<table>
<thead>
<tr>
<th>Key points</th>
<th>Acute peripancreatic fluid collection</th>
<th>Acute necrotic collection</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clinical severity</td>
<td>Mostly mild acute pancreatitis</td>
<td>Moderately severe acute pancreatitis or severe acute pancreatitis</td>
</tr>
<tr>
<td>Management algorithm</td>
<td>Conservative treatment (usually resolves spontaneously without intervention)</td>
<td>Likely increased morbidity and intervention rates (drainage or surgical treatment)</td>
</tr>
<tr>
<td>Course and prognosis</td>
<td>The hospital stay is usually about one week after onset; a good prognosis</td>
<td>Hospitalization often lasts from weeks to months; increased infection and mortality rates</td>
</tr>
<tr>
<td>CT/MRI imaging pattern</td>
<td>Occurs only in the setting of interstitial oedematous pancreatitis</td>
<td>Occurs in the setting of acute necrotising pancreatitis (including peripancreatic necrosis only)</td>
</tr>
<tr>
<td>Location and number of collections on CT/MRI</td>
<td>Mostly confined to simple retroperitoneal space or interfascial plane</td>
<td>Mostly in transabdominal-pelvic cavities and multiple spaces or interfascial planes</td>
</tr>
<tr>
<td>Shape, size, edge</td>
<td>Linear/strip-shaped, a small amount of collections, clear edge</td>
<td>Large patchy-shaped, a large amount of collections, unclear or irregular edge</td>
</tr>
<tr>
<td>Density/intense, enhancement characteristics</td>
<td>Homogeneous low density/hypointense T1 hyperintense T2 signal, no enhancement</td>
<td>Mixed features, mainly low density/hypointense T1 /hyperintense T2 signal, containing low density fat/fat signal intensity and low density or hypointense pancreas fragments; fragments are not enhancing</td>
</tr>
<tr>
<td>Secondary or concomitant signs</td>
<td>Rare</td>
<td>Frequent secondary infection with “bubble sign” (caused by infection itself or intestinal fistula with adjacent intestine); when a large area of intrapancreatic collections is present, “pancreatic duct disruption syndrome” may occur (further invasive operation is often required)</td>
</tr>
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</table>

CT:Computed tomography;MRI:Magnetic resonance imaging.

REFERENCES


