Pancreatic Fibrosis and Chronic Pancreatitis: Mini-Review of Non-Histologic Diagnosis for Clinical Applications

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Abstract: Pancreatic fibrosis is the dominant reversible pathological change and diagnostic factor in early chronic pancreatitis, defined by a mechanistic approach proposed in 2016. Main guidelines for chronic pancreatitis were published by the American Pancreas Association in 2014, the Japanese Society of Gastroenterology in 2015, and United European Gastroenterology in 2017. All three sets of guidelines mentioned that the staging of chronic pancreatitis is important but challenging. There are various image modalities for the non-histologic diagnosis of pancreatic fibrosis: (1) shear wave elastography, such as acoustic radiation force impulse with a cut-off value of 1.4 m/s; (2) strain elastography using grades of strain; (3) endoscopic ultrasonography using the Rosemont criteria or endoscopic ultrasound criteria for early chronic pancreatitis proposed by the Japan Pancreas Society; (4) computed tomography using the Hounsfield scale or number of micro-calcifications; and (5) magnetic resonance imaging using the apparent diffusion coefficient and the T1w flash and T2w HASTE sequences. The clinical applications are to (1) evaluate pancreatic tumors and inflammatory disease; (2) monitor dyspepsia with early chronic pancreatitis; (3) monitor individuals with a high risk of pancreatic cancer; (4) analyze a fatty pancreas with fibrosis; (5) predict a fistula after pancreatic surgery; and (6) predict outcomes for chronic pancreatitis or pancreatic cancer. The selection of tools will be dependent on the clinical scenario. Conclusion: There are various modalities for the non-histologic diagnosis of pancreatic fibrosis. The selection of the optimal device will be dependent on the clinical scenario.

Keywords: pancreatic fibrosis; early chronic pancreatitis; non-histologic diagnosis

Chronic pancreatitis is a major disease in the digestive system with severe complications in the end stage, including exocrine and endocrine insufficiency and pancreatic duct adenocarcinoma. An epidemiological study showed that a prevalence of chronic pancreatitis is between 36.9 and 41.8 per 100,000 people in America [1] and Japan [2] and up to 120 per 100,000 people in Europe [3]. Recent guidelines for chronic pancreatitis can be traced to the American Pancreas Association published in 2014 [4], the Japan Pancreas Society in 2015 [5], and United European Gastroenterology in 2017 [3]. These guidelines mentioned that the staging of chronic pancreatitis is important but challenging. Therefore, an international consensus statement focusing on early chronic pancreatitis was published in 2018 [6]. This consensus reappraised a mechanistic definition proposed in 2016 for chronic pancreatitis [7]. This mechanistic definition proposed a concept of early chronic pancreatitis, which
can exhibit the reversible structural fibrotic change of late stage chronic pancreatitis. The fibrosis of pancreatic parenchyma can be an early stage surrogate for chronic pancreatitis. With this in mind, endoscopic ultrasound (EUS) coupled with elastography would currently be the most sensitive diagnostic modality for detecting pancreatic parenchymal fibrosis.

Pancreatic fibrosis and atrophy are two major pathological changes of chronic pancreatitis [4]. Chronic pancreatitis is a risk factor of pancreatic duct adenocarcinoma, which shows a trend of increasing incidence [8,9]. The basic study of pancreatic fibrosis has advanced toward elucidating the pathogenesis and finding a possible therapeutic agent [10–12]. The stellate cell has been found to regulate the processes of pancreatic fibrosis. In addition, pancreatic fibrosis is a surveillance surrogate for individuals with a high risk of pancreatic duct adenocarcinoma [13].

There are histologic and non-histologic approaches for the diagnosis of pancreatic fibrosis. Image localization of pancreatic tissue fibrosis is valuable before biopsy, because the fibrotic change is probably focal. Pancreatic parenchyma tissue can be obtained by EUS-guided biopsy, percutaneous biopsy, or surgical biopsy. The non-histologic diagnosis of pancreatic fibrosis is an image-based method using different bio-physical mechanisms, including ultrasound elastography, computed tomography (CT), and magnetic resonance imaging (MRI) [14]. There are two types of ultrasound elastography—strain elastography through endoscopic [15] or transcutaneous routes [16] and shear wave elastography using only the transcutaneous route [17]. CT and MRI are both non-invasive transcutaneous image reconstructions.

The selection of different tools for the non-histologic diagnosis of pancreatic fibrosis is disease specific and scenario oriented. CT is used for predicting the post-operative pancreatic fistula complication by the CT value [18] or micro-calcifications. One study designed the computed tomography as an image prediction tool for severe pancreatic fibrosis and pain relief after surgical pancreatic resection in chronic pancreatitis patients. This study found that a predictor is calcification, especially when the number is > 10 [19].

Shear wave elastography for pancreatic fibrosis can be used in the following conditions: (1) differentiation of pancreatic cystic or solid tumor [20–23]; (2) acute and chronic pancreatitis [20,24–28]; (3) cystic fibrosis in the pancreas [14,29]; (4) prediction of post-operation pancreatic fistula [30]; and (5) surveillance of pancreatic fibrosis in a fatty pancreas [31]. The cut-off value of pancreatic fibrosis using acoustic radiation force impulse is 1.4 m/s [32]. It is well known that EUS can detect subtle parenchymal morphologic changes of pancreatic fibrosis in the early stage of chronic pancreatitis. It is evident that the EUS criteria of chronic pancreatitis have a histologic association with pancreatic fibrosis [4]. The commonly used EUS criteria for pancreatic fibrosis in chronic pancreatitis are the Rosemont criteria [33,34] and the early chronic pancreatitis EUS criteria defined by the Japan Pancreas Society [35,36]. EUS can be used for the surveillance of dyspeptic patients with early chronic pancreatitis and provides therapeutic guidance [37–39]. In these clinical studies, the diagnostic criteria of early chronic pancreatitis proposed by the Japan Pancreas Society have two parts. The first part details the clinical or functional criteria: (1) recurrent upper abdominal pain (two or more attacks); (2) abnormal serum or urine enzyme levels; (3) abnormal exocrine function; and (4) continuous heavy drinking (>80 g/d). The second part details the image features of EUS or endoscopic retrograde cholangiopancreatography (ERCP). The EUS features are (1) lobularity with honeycombing; (2) lobularity without honeycombing; (3) hyperechoic foci without shadowing; (4) Stranding; (5) cysts; (6) dilated side branches; and (7) hyperechoic main pancreatic duct margin. ERCP shows irregular dilatation of more than three duct branches [35]. In these references, the diagnosis required two or more clinical features and two or more EUS features.

Secretin-enhanced magnetic resonance cholangiopancreatography can depict the ductal system of the pancreas but lacks further analysis of parenchyma. A recent study, specified as magnetic resonance imaging as a non-invasive method for the assessment of pancreatic fibrosis (MINIMAP), was designed to incorporate both the parenchymal and ductal features of chronic pancreatitis. The aim of this study was to make a MRI-based quantitative scoring system for pancreatic fibrosis [40]. One recent study showed that body composition and pancreatic fibrosis could both be factors associated with quality of life and treatment outcomes in patients with chronic pancreatitis or
Pancreatic duct adenocarcinoma. Pancreatic fibrosis can be calculated by MRI images using T1 signals, apparent diffusion coefficient maps, and T1w flash and T2w HASTE sequences [14,41].

Here, we explain the demerits of each of the modalities. Endoscopic strain elastography is an operator-dependent invasive procedure without quantitative results. Shear wave elastography is a point-targeted evaluation and is occasionally difficult to perform in an extremely obese patient. The CT examination exposes the patient to radiation and is unable to detect the early structural change of pancreatic fibrosis. MRI is a high-cost image device and is superior in the pancreatic ductal system but not in detecting the subtle morphologic change of parenchyma in the current technique.

In addition to the image diagnostic methods, serologic research for the diagnosis of chronic pancreatitis with parenchymal fibrosis is evolving. One specific serum circulating immune signature was proved to diagnose chronic pancreatitis, pancreatic duct adenocarcinoma, and recurrent acute pancreatitis with an area under the curve of approximately 0.77–0.86 [42]. For serologic predictors of chronic pancreatitis with fibrosis, the very low level of pancreatic serum enzymes (amylase or lipase) can be regarded as a marker [43,44]. Chromogranin-A is an acidic protein and one member of the granin family of neuroendocrine secretory proteins and is located in the cells of the endocrine and nervous systems. It can be used as a plasma marker or prognostic factor for pancreatic neuroendocrine tumor [45,46] and advanced pancreatic cancer [47,48] and, theoretically, for both diseases with concomitant focal pancreatic fibrosis. Besides, the association between pancreatic fibrosis and the level of chromogranin-A in tissue has had a couple of research findings as follows: (1) an increase in chromogranin A-positive but hormone-negative endocrine cells in the pancreatic tissue of cystic fibrosis patients [49] and chronic pancreatitis patients [50] and (2) an increased expression of chromogranin-A in the duodenal mucosa of pancreatic fibrosis patients [31].

Conclusively, there are various modalities for the non-histologic diagnosis of pancreatic fibrosis. The selection of the optimal device will be dependent on the clinical scenario. The various image modalities for the non-histologic diagnosis of pancreatic fibrosis and clinical applications are summarized in Table 1.

| Image Modalities | Parameters | Clinical Scenario Application |
|------------------|------------|-----------------------------|
| Trans-abdomen strain elastography | grade of strain | pancreatic tumors |
| Trans-abdomen shear wave elastography (ARFI) | ≥ 1.4 m/s | pancreatic tumors, dyspepsia surveillance, fatty pancreas, pancreatic tumor |
| EUS | Rosemont criteria, early CP criteria | high-risk population surveillance, dyspepsia surveillance, pancreatic tumors |
| EUS: strain elastography | grade of strain | predicting fistula after pancreatic resection operation |
| CT | CT value, micro-calciﬁcations | predicting therapeutic outcomes and quality of life for chronic pancreatitis or pancreatic cancer |
| MRI | ADC, T1, T2 different sequences | |

ARFI—acoustic radiation force impulse, EUS—endoscopic ultra-sonography, CP—chronic pancreatitis, CT—computed tomography, MRI—magnetic resonance image, ADC—apparent diffusion coefficient.

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