Recent Advances in Pancreatic MR Imaging: A Guide on How, When, and Why to Perform

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Abstract

Imaging of the pancreas is often challenging because of its retroperitoneal location and unique set of pathologies. Conventional imaging modalities, such as transabdominal and endoscopic ultrasonography (EUS), computed tomography (CT), and magnetic resonance imaging (MRI), are well described in the literature. However, with modern demand for functional and molecular information from imaging studies, newer imaging modalities and modifications of existing modalities are developed. MRI is widely used as a problem-solving tool in pancreatic pathologies. Magnetic resonance cholangiopancreatography (MRCP) is an excellent technique for the depiction of the pancreatic ductal or biliary ductal pathologies. Newer modification of MRI including secretin MRCP, advanced diffusion-weighted imaging (DWI), perfusion imaging, and tissue composition analysis (fat and fibrosis quantification) add to the arsenal of MRI of the pancreas. In this review, we discuss the evolution of MRI of the pancreas and clinical application of advanced MR sequences.

Keywords
► diffusion magnetic resonance imaging
► magnetic resonance imaging
► pancreas
► recent advance

Introduction

Magnetic resonance imaging (MRI) has evolved over the years with significant improvement in the scan time and image quality with the improvement of MR hardware and software. Present-day MR scanners have up to 30 radiofrequency (RF) channels (receiver pathways) and more than 100 coil segments that create the magnetic field. A higher number of channels have enabled techniques, such as parallel imaging, to be implemented that have significantly shortened the scan time.

Evolution of MRI of Pancreas

One of the earliest studies of MR imaging of the pancreas was published by Tscholakoff et al in 1987. The study was done with small sample size and the scan was performed on a 0.35 Tesla (T) MR machine. Since then, there have been significant improvements seen in software and hardware in the quality with reduced scan time. The earliest literature on MR cholangiopancreatography (MRCP) was around in the early 1990s. Three decades ago only T1 and T2 sequences were available for the abdomen. Currently, the field strength of the MR machine in most centers performing abdominal imaging is 1.5 or 3 T.6 Higher field strength allows an improved signal to noise ratio, thereby allowing advanced sequences such as diffusion-weighted imaging (DWI) to be performed. Increased field strength also improves contrast resolution with better detection of focal lesions in postcontrast T1-weighted images and increasing conspicuity of lesions in T2-weighted images.

Today, the main role of conventional MRI of the pancreas is to better characterize the cystic and solid parenchymal lesions and evaluation of pancreatic ductal anatomy. However, newer sequences, such as DWI, perfusion imaging, fat, and fibrosis quantification, provide additional information about focal and diffuse pancreatic pathologies (Table 1).

Normal Morphology of Pancreas in MRI

The pancreas has an inherent high T1 signal due to the presence of water-soluble proteins, paramagnetic substances, such as manganese within the cells, and numerous endoplasmic reticula in the exocrine cells (Fig. 1). There is a progressive loss of this high T1 signal with increasing severity
in chronic pancreatitis (►Fig. 2). This change is quantified T1-relaxometry (see the section “T1 Mapping/Relaxometry”). In T2-weighted images, the pancreas appears slightly hyperintense to the skeletal muscles.

**Conventional Pancreatic MRI Protocol**

**Patient Preparation**

The patient should be on fast for at least 4 hours to reduce the stomach and duodenal contents. Fasting also improves the distension of the gallbladder and reduced peristalsis of the small bowel. The signal from fluid in the duodenal lumen can be suppressed by commercially available negative oral contrast (e.g., LumiVision, a manganese suspension). Other alternatives include blueberry or pineapple juice or dilute solution of gadolinium containing contrast medium.7–10

Conventional MRI of the pancreas routinely consists of the following sequences:

**T1-Weighted Image**

T1 signal of the pancreas is reduced in conditions such as inflammation and neoplasm. T1-weighted image is very useful to assess bleed and to see the changes in the peri-pancreatic fat. Breath-hold fast acquisition spoiled gradient sequences with short time to echo (TE) are commonly used. In-phase and out-phase sequences are useful in assessing fat within the pancreas and are routinely used in most centers.1,6,11,12 The T1-weighted image acts as a baseline for the assessment of postcontrast T1-weighted sequences.

**T1-Weighted Postcontrast Image**

Breath-hold T1-weighted sequences are used for postcontrast MRI. These are three dimensional (3D) spoiled gradient echo sequences with fat suppression. The protocol for the postcontrast T1-weighted sequence is provided in ►Table 2. Normal pancreatic parenchyma mostly enhances the early arterial phase, while most adenocarcinomas show low and delayed enhancement (►Fig. 3). Neuroendocrine tumors show early enhancement.

**T2-Weighted Image**

The T2-weighted sequence can either be single-shot fast spin-echo (SSFSE), or half Fourier acquisition single-shot turbo spine-echo (HASTE), or multishot echo sequence. These sequences are acquired with breath-hold technique. Image quality is better with SSFSE compared with FSE or TSE sequences mainly because of faster acquisition. However, the contrast resolution and signal-noise ratio are affected due to longer echo train length and T2 decay. The fluid signal within or outside the pancreas is well visualized in T2 sequences, especially after fat suppression.13–17

**Magnetic Resonance Cholangiopancreatography**

MRCP sequence is a heavily T2-weighted sequence (FSE or SSFSE sequence) that enhances the fluid signals while suppressing other signals in the background and, therefore, shows the ductal anatomy and cystic lesions in great detail (►Fig. 4). SSFSE sequences are faster and produce fewer motion artifacts. Two dimensional (2D) sequences are performed with single-breath-hold and are thick slab images. The 3D MRCP is performed by respiratory triggering and requires uniform breath-hold technique. In the 3D MRCP sequence, thin contiguous images are obtained that can be reconstructed by thin or thick maximum intensity projections (MIP). The thin MIP reconstructions help in assessing the anatomy of the ducts (►Fig. 4); the relationship is between focal mass or cyst and duct (►Fig. 5); and dilatation, strictures, or stones in the ducts (►Fig. 6).18–21

**Quantitative Imaging and Newer Applications**

Although MRI has been established as an excellent modality to depict the anatomy of the pancreas, newer sequences are being developed for quantitative assessment and improved resolution.6

**Secretin MRCP**

Secretin is a peptide hormone that stimulates bicarbonate-rich fluid secretion from the pancreas and increases the

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**Table 1** Advanced pancreatic MRI sequences in a nutshell

<table>
<thead>
<tr>
<th>Modality</th>
<th>Use</th>
<th>Comment</th>
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<tbody>
<tr>
<td>DWI</td>
<td>Identification of early mild and focal pancreatitis</td>
<td>Routinely used. Diffusion kurtosis, tensor imaging, and IVIM are in early clinical research</td>
</tr>
<tr>
<td></td>
<td>Identification of small liver and nodal metastases in pancreatic adenocarcinoma</td>
<td></td>
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<tr>
<td></td>
<td>Identification of subtle pancreatic primary neoplasm</td>
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<td></td>
<td>Identification of solid nodules in cystic neoplasm</td>
<td></td>
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<tr>
<td></td>
<td>Characterization of focal pancreatic lesion [smaller role]</td>
<td></td>
</tr>
<tr>
<td>MR perfusion</td>
<td>Differentiation of neoplasm from benign process</td>
<td>DCE MR routinely used</td>
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<tr>
<td></td>
<td>Monitoring response to antiangiogenic therapy</td>
<td></td>
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<tr>
<td>T1 relaxometry</td>
<td>Diagnosis and monitoring of chronic pancreatitis</td>
<td>Limited clinical use</td>
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<tr>
<td>T2* / R2* mapping</td>
<td>Parenchymal iron quantification</td>
<td>Early research</td>
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<td>Fat quantification</td>
<td>Surrogate marker of metabolic disease</td>
<td>Research</td>
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<tr>
<td>MR elastography</td>
<td>Differentiation of adenocarcinoma and focal autoimmune pancreatitis</td>
<td>Research</td>
</tr>
<tr>
<td>Secretin MRCP</td>
<td>Uncover ductal communication of cystic lesions (branched IPMN)</td>
<td>Limited use, not available in India</td>
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<td></td>
<td>Assessment of exocrine function of pancreas</td>
<td></td>
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<td></td>
<td>Demonstration of ductal leak</td>
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Abbreviations: DCE, dynamic contrast enhancement; DWI, diffusion-weighted imaging; IPMN, intraductal papillary mucinous neoplasm; IVIM, intravoxel incoherent motion; MRCP, magnetic resonance cholangiopancreatography; MRI, magnetic resonance imaging.

The recommended dose of secretin is around 0.2 µg/kg. The peak action of the injected secretin is around 3 to 5 minutes after administration and the duodenal lumen gets filled with the bicarbonate rich fluid after 5 minutes and the maximum tone of the sphincter of Oddi. Injection of secretin before MRCP imaging distends pancreatic ducts and uncovers small ductal communications. Secretin induced MRCP is also an excellent noninvasive imaging technique to evaluate the exocrine function of the pancreas. However, this study has limited applicability due to limited availability and high cost of secretin.4,22,23

Table 2  Protocol for postcontrast T1-weighted sequence

<table>
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<tr>
<th>Contrast volume</th>
<th>0.1 mL/kg</th>
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<tr>
<td>Rate</td>
<td>2 mL/s followed by 20 mL saline flush</td>
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<tr>
<td>Arterial phase</td>
<td>20–25 seconds delay</td>
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<tr>
<td>Portal venous phase</td>
<td>55 s</td>
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<tr>
<td>Delayed</td>
<td>90–180 s</td>
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<tr>
<td>Sequence</td>
<td>VIBE/LAVA (3D spoiled gradient T1-weighted sequence)</td>
</tr>
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</table>

Abbreviations: 3D, three-dimensional; LAVA, liver acquisition with volume acceleration; VIBE, volumetric interpolated breath-hold examination.

Fig. 1 Axial T1 weighted fat-suppressed image showing relatively hyperintense pancreas (arrow).

Fig. 2 Loss of T1 signal in chronic pancreatitis. Axial T1-weighted fat-suppressed image at the level of the pancreatic tail (A) shows isointense parenchyma with loss of T1 signal (between arrows). Compare with retained normal high T1 signal in the pancreatic head (arrow in B).

Fig. 3 Dynamic/multiphasic postcontrast T1-weighted images of the pancreas. Early arterial (A), late arterial (B), and equilibrium (C) phase show early enhancement of pancreatic parenchyma (asterisk in A) and delayed enhancement of the wall of the cystic lesion at the pancreatic tail (arrow in C). Case of mucinous cystic neoplasm of the pancreas.
output is around 6 and 8 minutes. The patient is usually fasted for at least 4 hours and duodenal fluid is suppressed by a negative oral contrast agent such as very dilute gadolinium. After injection of secretin, SSTSE thick slab MRCP image (30–70-mm thickness) is obtained in the coronal plane with a breath-hold or respiratory trigger for 10 minutes postinjection, at 30-second intervals. This study is ended with a respiratory-triggered MRCP sequence. Secretin MRCP is used and volume of blood plasma (Vp) within of the pancreas semiquantitatively. Filling of the duodenum is graded as follows: grade 1, fluid filling is confined to the duodenal bulb; grade 2, fluid reached up to the second portion of the

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**Fig. 4** Demonstration of ductal anatomy in ansa pancreatica. Thick slab 2D MRCP sequence shows a characteristic loop in the main pancreatic duct (arrow). 2D, two-dimensional; MRCP, magnetic resonance cholangiopancreatography.

**Fig. 5** Demonstration of ductal communication between side branch intraductal papillary mucinous neoplasm (IPMN) of the pancreas and main pancreatic duct. A-B, Axial T2-weighted (A) and postcontrast T1-weighted (B) images show a multiloculated cystic lesion in the pancreatic head (arrow). (C) Maximum intensity projection of thin slab 3D MRCP shows thin ductal communication (thin arrow) between the cystic lesion (asterisk) and main pancreatic duct (thick arrow), establishing the diagnosis of side branch IPMN. 3D, three-dimensional; MRCP, magnetic resonance cholangiopancreatography.

**Fig. 6** Demonstration of altered ductal morphology in chronic pancreatitis. Maximum intensity projection of thin slab 3D MRCP shows dilated main pancreatic duct with prominent side branches. 3D, three-dimensional; MRCP, magnetic resonance cholangiopancreatography.
duodenum; grade 3, filling up to the third portion of the duo
denum; and grade 4, complete filling of duodenum and por-
tion of jejunum (►Fig. 7).  
Secretin MRCP can uncover small ductal channels and
communication aiding in the differentiation of branched
IPMN and other cystic lesions of the pancreas. Secretin
MRCP is also useful in the identification of pancreatic duct
leak in the postoperative pancreas and ductal anomalies
such as pancreas divisum, annular pancreas, and anomalous
pancreaticobiliary junction.

Perfusion Imaging

Regional tissue perfusion can be assessed based on the uptake
and washout of contrast agents. Dynamic contrast enhance-
ment (DCE) MRI uses repeated acquisition of T1-weighted
gradient weighted images at short intervals and evaluated
the change in signal intensity of tissue due to contrast enhance-
ment. Dynamic susceptibility contrast (DSC) MRI uses a signal
drop in T2*-weighted images in a similar manner. Arterial spin
labeling is a novel technique that uses “intrinsic contrast” by
saturating water protons in arterial blood.

DCE is the most commonly performed technique of MR
perfusion. It can be assessed qualitatively by observing dif-
ferent phases and by generating a time-intensity curve. Semi-
quantiative parameters can be calculated from the curve
include maximal initial slope, peak signal intensity, and
washout gradient. Quantitative parameters include transfer
constant (Ktrans), volume of extravascular extracellular space
(Ve), and volume of blood plasma (Vp) within the region of
interest and rate constant (Kep). Ktrans is a marker of capillary
permeability when the overall tissue permeability is low and
it represents the tissue blood flow when the overall tissue
permeability is very high.

MR perfusion is a potentially useful tool in the differenti-
ation of neoplasms and tissue with different perfusion and
assessment of response to antiangiogenic therapy.

Diffusion-Weighted Imaging

The principle of DWI is in certain types of pathologies that
the random Brownian motion of water molecules in the
tissue is restricted. Diffusion restriction typically occurs in
tissues due to increased cell count per unit area (neoplasia)
and increased cell size (cytotoxic edema and inflammation)
resulting in a decrease in interstitial space, as well as in pre-
se of thick and dense interstitial tissue (fibrosis) or material
(pus or proteinaceous material). Pathologies, such as inflam-
mation, tumor, and fibrosis, cause restriction of this motion
and appear hyperintense on DWI with a low-apparent diffu-
sion coefficient (ADC) values.

DWI is a modification of spin-echo T2-weighted imaging
that utilizes the “ultrafast” echo-planar imaging technique.
Two opposing gradients are applied before and after the
180-degree refocusing pulse. The first gradient induces a
phase shift in the water protons that is canceled by the sec-
ond gradient, provided the protons are stationary or have
restricted motion. On the other hand, free protons are com-
pletely dephased by the second gradient. Therefore, the tis-
ue with restricted motion returns higher signals from water
protons. The amount of diffusion “weighting” is indicated
by the b-value of the sequences. In clinical practice, DWI is
usually obtained with two or more b-values. The low b-value
images (b < 200 second mm²) are similar to T2-weighted
images and higher b-values (b > 800 second mm²) are more
diffusion weighted. The DWI images can be used to gener-
ate the ADC maps. Lesions with restricted diffusion appear
hyperintense in high b-value DWI images and hypointense in

Fig. 7 Secretin MRCP in a 47-year-old woman with recurrent abdominal pain and vomiting. (A) Preinjection, (B) 1 minute, (C) 2 minutes,
(D) 5 minutes, and (E) 10 minutes of postinjection images. A mild increase in diameter of pancreatic duct 1 or 2 minutes following administra-
tion of intravenous secretin with the return to normal caliber on subsequent images. Mild exocrine insufficiency is demonstrated by filling up
of up to the third portion of duodenum in the 10-minute postinjection image (grade 3 response). (Image courtesy: Dr. Avinash Kambadakone
Ramesh, Massachusetts General Hospital, Boston, Massachusetts, United States]. MRCP, magnetic resonance cholangiopancreatography.
ADC maps. ADC of a lesion can be quantified by drawing an ROI (region of interest) on the ADC map in most commercial MR scanners. ADC is represented by the unit mm$^2$/s.

Normal pancreas should be homogeneous and slightly hyperintense DWI with slightly low signal in ADC. The average ADC of the pancreas is reported to be $1.6 \times 10^{-3}$ mm$^2$/s. Heterogeneous ADC distribution and increased ADC values are reported in type-1 diabetes mellitus along with the reduction in pancreatic volume. Common clinical utility of DWI in pancreatic imaging is given in Table 1.

**Pancreatic Inflammation**

Acute inflammation reduces the ADC of pancreatic tissue. The average ADC of acute pancreatitis is reported to be approximately $1.2-1.6 \times 10^{-3}$ mm$^2$/s depending on the severity of pancreatitis. ADC values are shown to correlate with the Balthazar score of severity of pancreatitis with lower values representing a higher score. DWI can be used to identify mild acute interstitial pancreatitis especially in patients in whom contrast study cannot be done. With successful treatment of pancreatitis, ADC values are shown to normalize. Caution should be exercised in presence of necrotizing pancreatitis, as the necrotic area may show higher ADC values and lower signal on DWI due to loss of cell membrane integrity (“facilitated diffusion”). Therefore, it is important to correlate with contrast-enhanced images when available. Areas of pancreatic necrosis do not show contrast enhancement. Some authors have suggested that the presence of superadded infection in acute pancreatic or peripancreatic collection may show diffusion restriction; however, this is not universally accepted, and infection is often proven by aspiration.

In chronic pancreatitis, pancreatic parenchyma is progressively replaced by fibrosis. ADC values decrease with increasing pancreatic fibrosis. DWI is shown to diagnose chronic pancreatitis with high specificity.

Autoimmune pancreatitis (AP) can show markedly low-ADC value (very high signal on DWI). ADC values in AP can be lower than that of adenocarcinoma. ADC is shown to be even lower in symptomatic patients. As autoimmune pancreatitis is often focal and shows strong restriction in diffusion, it is sometimes mistaken for pancreatic cancer. ADC values are useful in monitoring treatment response.

**Solid Pancreatic Lesions**

The most common solid neoplasm of the pancreas is adenocarcinoma, which is often isointensating on computed tomography (CT) scan and therefore difficult to delineate. DWI helps in identifying small, subtle pancreatic lesions. DWI has shown similar or better diagnostic performance to contrast-enhanced MRI in identifying pancreatic adenocarcinoma. However, lesions are sometimes difficult to differentiate from upstream atrophic pancreatic parenchyma because of decreased ADC value of upstream pancreas owing to fibrosis. DWI is also helpful in identifying subtle metastasis in the liver and lymph nodes.

**Cystic Pancreatic Lesions**

DWI helps in the identification of small mural nodules in cystic neoplasms such as intraductal papillary mucinous neoplasm (IPMN). A purely cystic lesion is more likely to be benign.

**Newer Diffusion Techniques**

Normal tissue is composed not only of interstitium and cells but also small blood vessels. Water diffusion in tissues is a combination of free (or coherent) diffusion through the interstitium and perfusion (or incoherent diffusion) through the microcapillary network. Diffusion of water protons detected by conventional DWI images is based on a mono-exponential model which does not take into account component of the microcapillary perfusion of tissue. This is one of the reasons why the diffusion coefficient calculated by conventional DWI is called the “apparent” diffusion coefficient. However, intravoxel incoherent motion (IVIM) MRI utilizes multiple $b$-values in a biexponential model that can calculate the component of tissue microcapillary perfusion. In addition to the conventional ADC value, IVIM also provides $D$-value, $D^*$-value, and $f$-value (perfusion fraction). $D$-value (or slow ADC value) is the true diffusion coefficient and reflects the tissue microstructure. The $D^*$-value (or fast ADC value), also referred to as the pseudo-diffusion coefficient, represents the perfusion effect of the incoherent microcirculation within the voxel. The proportion of the perfusion effect is defined as the perfusion fraction ($f$). Since pancreatic adenocarcinoma is hypovascular compared with normal...

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**Fig. 8** Mild acute pancreatitis detected in diffusion-weighted imaging (DWI). (A) axial T2-weighted HASTE image shows subtle loss of lobulations of the pancreatic tail (arrow) without any enlargement, increased T2-signal or edema. (B) axial DWI ($b = 800$ second mm$^2$) shows focal hyperintensity of the pancreatic tail (arrow). (C) apparent diffusion coefficient (ADC) map shows hypointensity in the corresponding with a low ADC value of $1.10^{-1}$ mm$^2$/s. HASTE, half Fourier acquisition single-shot turbo spine-echo.
Fig. 9 Necrotizing pancreatitis with facilitated diffusion. Axial T2-weighted HASTE image (A) shows a bulky edematous pancreas with a focal area of high signal in body portion (arrow) that shows no contrast enhancement in postcontrast T1-weighted image (arrow) (B). (C) Diffusion-weighted image ($b = 800$ second mm$^2$) shows no hyperintensity in the corresponding area with high signal on apparent diffusion coefficient (ADC) map (D) suggesting high ADC value. HASTE, half Fourier acquisition single-shot turbo spine-echo.

pancreatic parenchyma because of the destruction of the normal capillary network, adenocarcinoma shows lower perfusion fraction ($f$).45

Free diffusion is isotropic, that is, occurs in all directions. However, in biological tissue diffusion is anisotropic, that is, more in one or more direction and restricted in others. This is due to the presence of cell membranes, myelin sheath, or tissue architecture. Diffusion tensor imaging (DTI) detects the degree of the restriction diffusion of water molecules and evaluates the different directions of diffusion. DTI can provide some details on the microstructure of tissues that are not available in conventional imaging.46 There are five main parameters in the DTI, including mean diffusivity (MD), three eigenvalues $\lambda_1, \lambda_2, \lambda_3$, and fractional anisotropy (FA). MD is theoretically more representative of the diffusivity of water protons. The FA represents the fraction of anisotropic water diffusion. Nissan et al showed that DTI parameters are lower in pancreatic adenocarcinoma than that of the distal normal pancreatic tissue.47 FA and mean ADC were found to lower in acute pancreatitis compared with the normal pancreas.48

Diffusion kurtosis imaging (DKI) utilizes a diffusion model that addresses the nonuniform (non-Gaussian) movement of water protons. In this model, the heterogeneity of tissue is taken into account and DKI is supposed to represent the complex microstructure of tissue. DKI is still in its early research phase. It is technically demanding and requires the acquisition of high and very high $b$-values. The parameters obtained from DKI are mean kurtosis (MK) and corrected diffusion coefficient ($D_K$). $D_K$ is shown to be able to differentiate between tumor and normal tissue,49 correlate with HbA1c levels, and may potentially serve as a biomarker of glycemic control.50

Limitations of Diffusion Imaging

The major technical limitations of DWI are the inherent low signal-to-noise ratio (SNR), low spatial resolution, and susceptibility to artifacts. SNR can be improved in 3 T at the cost of more artifacts. SNR is also improved by smaller matrix size, larger FOV, shorter TE, and use of free-breathing sequences.4 Newer techniques, such as zoomed EPI-DWI (ZOOMit, Siemens Healthcare, Erlangen, Germany) used 2D spatially-selective RF excitation pulses combined with a reduction in the FOV in the phase-encoding direction leading to improved spatial resolution and decreased
Fig. 10 Subtle pancreatic mass demonstrated in diffusion-weighted imaging (DWI). (A), contrast-enhanced axial CT shows atrophic pancreas with abrupt duct cutoff and altered areas of increased enhancement in the body of the pancreas. (B, C) Fat-suppressed T2-weighted axial HASTE (B) and postcontrast T1-weighted image shows subtle ill-defined hyperintense mildly enhancing lesion in the body portion. (D), coronal 3D MRCP in maximum intensity projection shows abrupt cut-off of duct in the same location. (E), axial DWI ($b = 800$ second mm$^2$) clearly shows a hyperintense lesion in the corresponding location. Compare the conspicuity of the lesion in DWI to that in other images. Biopsy revealed adenocarcinoma. 3D, three-dimensional; HASTE, half Fourier acquisition single-shot turbo spine-echo; MRCP, magnetic resonance cholangiopancreatography.

Fig. 11 Demonstration of liver metastases in diffusion-weighted imaging (DWI). (A) Axial T2-weighted HASTE image shows a well-defined hyperintense mass lesion in the head of the pancreas (solid arrow) with few ill-defined faintly hyperintense nodules in the liver (open arrow). (B), Axial DWI ($b = 800$ second mm$^2$) offers a much better demonstration of liver nodules (open arrow). HASTE, half Fourier acquisition single-shot turbo spine-echo.
susceptibility artifacts. To improve image quality with similar FOV, sequences, such as readout-segmented echo-planar diffusion methods (rs-DWI), have been developed (RESOLVE [readout segmentation of long variable echo trains], Siemens Healthcare, Erlangen, Germany).

**Fat Quantification**

Pancreatic fat content increases with age and body mass index. Although pancreatic fat content may be increased in patients with diabetes and prediabetes, this relationship is not established. More recent studies suggest that the fat content of pancreatic tail may be related to the future development of diabetes.

Fat quantification can be done by various techniques including two-point Dixon, three-point Dixon, and chemical shift imaging. Dixon sequence generates a set of in-phase, out-phase, fat-only, and water-only images from the same acquisition. Fat containing lesions are hypointense in opposed-phase images (“signal drop”) and hyperintense in fat-only images (∼Fig. 12).

MRI proton density fat fraction (MRI-PDFF) is obtained by a quantitative DIXON sequence that utilizes multiple TE. This sequence automatically generates maps of fat fraction, T2, T2*, and R2*. The fat fraction can be calculated from placing the ROI in the pancreas (∼Figs. 13 and 14). The normal fat content of the pancreas is reported to be between 4.6 and 4.9%.

**T1 Mapping/Relaxometry**

The pancreas has a relatively short T1 relaxation and is hyperintense compared with adjacent soft tissues due to the presence of acinar proteins and rough endoplasmic reticulum. The T1 relaxation time of pancreatic tissue increases in diffuse pancreatic pathology such as fibrosis, atrophy, or edema. An increase in T1 relaxation time decreases the T1 signal intensity. A subtle decrease in T1 signal can be an early marker of chronic pancreatitis. An increase in T1 relaxation time can be semiquantitatively assessed by quantifying the loss of high T1 signal in comparison to control tissue, such as muscle. Quantitative assessment of T1 relaxation time requires the generation of parametric T1 maps from which T1 relaxation time can be assessed by placing an ROI (∼Fig. 15).

**T2*/R2* Mapping**

T2*/R2* mapping is widely used in the liver as a measure of parenchymal iron content but large validation studies in the pancreas are lacking. Correlations between logarithmic pancreatic T2* values and β-cell function is shown in some studies. Early research has shown that the pancreatic R2* values correlate well with liver iron overload and may have potential use as a surrogate marker for hereditary hemochromatosis.

**Magnetic Resonance Elastography**

Magnetic resonance elastography (MRE) is “virtual palpation” where the stiffness of tissue is calculated indirectly by calculating the velocity of shear waves propagated through tissue generated by an external source of vibration. It is currently an established technique to measure the stiffness of the liver that acts as an indirect marker for fibrosis. MRE of the pancreas is limited due to the retroperitoneal location of the pancreas and overlying stomach and transverse colon. Chronic pancreatitis and carcinoma pancreas are shown to increase pancreatic stiffness. Early studies of MRE show...
potential in the differentiation between autoimmune pancreatitis and pancreatic adenocarcinoma.\textsuperscript{50}

**Conclusion**

MRI has a problem-solving role in pancreatic imaging in routine clinical practice. Advanced diffusion techniques can localize and characterize focal and diffuse pathologies serving as a useful adjunct to conventional MRI. Secretin MRCP can improve visualization of ductal anatomy and pathologies and can potentially demonstrate pancreatic leak. Advanced imaging sequences and specialized sequences are useful in the quantitative and semiquantitative assessment of pancreatic fat, iron, fibrosis, and tissue characters that can help in better characterization of diffuse pancreatic diseases in the near future.

Fig. 13 Normal fat fraction map of the pancreas with fatty liver. In phase (A) and opposed-phase images (B) show diffuse loss of signal in the liver in opposed phase with normal signal in pancreas suggesting fatty deposition of the liver. (C) Fat fraction map shows fat deposition in liver (green, compare with visceral fat) with a pancreatic fat fraction of 0.9%, within normal limit.

Fig. 14 Steatosis of the pancreas with a normal liver. In phase (A) and opposed-phase images (B) show patchy loss of signal in the pancreas in opposed-phase with a normal signal in liver suggesting pancreatic steatosis. (C) Fat fraction map shows normal liver (blue, compare with image 13) with a pancreatic fat fraction of 5.5%.

Fig. 15 T1 mapping of normal pancreas. (A) Axial T1-weighted fat-suppressed image shows normal pancreas. (B) ROI placed on the body of the pancreas on T1 map shows T1 relaxation time of 976 milliseconds.

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