Magnetic resonance imaging in the assessment of pancreatic cancer with quantitative parameter extraction by means of dynamic contrast-enhanced magnetic resonance imaging, diffusion kurtosis imaging and intravoxel incoherent motion diffusion-weighted imaging

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Abstract

Background: Despite great technical advances in imaging, such as multidetector computed tomography and magnetic resonance imaging (MRI), diagnosing pancreatic solid lesions correctly remains challenging, due to overlapping imaging features with benign lesions. We wanted to evaluate functional MRI to differentiate pancreatic tumors, peritumoral inflammatory tissue, and normal pancreatic parenchyma by means of dynamic contrast-enhanced MRI (DCE-MRI), diffusion kurtosis imaging (DKI), and intravoxel incoherent motion model (IVIM) diffusion-weighted imaging (DWI)-derived parameters.

Methods: We retrospectively analyzed 24 patients, each with histopathological diagnosis of pancreatic tumor, and 24 patients without pancreatic lesions. Functional MRI was acquired using a 1.5 MR scanner. Peritumoral inflammatory tissue was assessed by drawing regions of interest on the tumor contours. DCE-MRI, IVIM and DKI parameters were extracted. Nonparametric tests and receiver operating characteristic (ROC) curves were calculated.

Results: There were statistically significant differences in median values among the three groups observed by Kruskal–Wallis test for the DKI mean diffusivity (MD), IVIM perfusion fraction (fp) and IVIM tissue pure diffusivity (Dt). MD had the best results to discriminate normal pancreas plus peritumoral inflammatory tissue versus pancreatic tumor, to separate normal pancreatic parenchyma versus pancreatic tumor and to differentiate peritumoral inflammatory tissue versus pancreatic tumor, respectively, with an accuracy of 84%, 78%, 83% and area under ROC curve (AUC) of 0.85, 0.82, 0.89. The findings were statistically significant compared with those of other parameters (p value <0.05 using McNemar’s test). Instead, to discriminate normal pancreas versus peritumoral inflammatory tissue or pancreatic tumor and to differentiate normal pancreatic parenchyma versus peritumoral inflammatory tissue, there were no statistically significant differences between parameters’ accuracy (p >0.05 at McNemar’s test).

Conclusions: Diffusion parameters, mainly MD by DKI, could be helpful for the differentiation of normal pancreatic parenchyma, perilesional inflammation, and pancreatic tumor.

Keywords: diffusion, magnetic resonance imaging, pancreatic cancer, perfusion

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Introduction
Diagnosis of pancreatic cancer remains challenging, due to overlapping imaging features with benign lesions notwithstanding great advances with multidetector computed tomography and magnetic resonance imaging (MRI). However, a proper detection and characterization of pancreatic lesions is mandatory because the prognosis is linked to tumor type and grade, and correct staging on accurate imaging; in fact, a pancreatic cancer that infiltrates lymphatic vessels can be manifested as infiltration of peripancreatic tissue. This local invasion can determine underestimation of real extension and grade of the disease.

Thus, an imaging modality that provides higher tumor conspicuity would be desirable to improve staging and clinical outcomes. Quantitative analysis of perfusion parameters by using dynamic contrast-enhanced magnetic resonance imaging (DCE-MRI) has been considered. Moreover, diffusion-weighted imaging (DWI) is another magnetic resonance modality that is able to objectively and quantitatively assess perfusion and diffusion to aid detection of malignancies. DWI can provide additional information to identify focal pancreatic lesions, verifying more restricted diffusion in solid malignant tumors versus benign inflammatory ones. However, the diffusion-weighted signal and the apparent diffusion coefficient (ADC) values can be influenced both by molecular diffusion and by microcirculation, or blood perfusion, and, therefore, ADC values may be polluted from perfusion effects, reducing the ADC reliability to characterize pancreatic lesions.

Previous studies with the IVIM approach have demonstrated that the reduced ADC in pancreatic adenocarcinomas (PDACs) possibly relates to a difference in perfusion fraction (fp), which is reduced in PDACs; therefore, fp is the best factor among DWI-derived parameters to differentiate pancreatic adenocarcinomas from benign lesions. Microcirculation or perfusion effects can be separated by diffusion water motion biexponential curve fit analysis with the intravoxel incoherent motion (IVIM) model.

and Jensen and coworkers suggested a non-Gaussian diffusion model, known as diffusion kurtosis imaging (DKI). This model showed better performance than conventional ADC in tumor detection and staging.

The purpose of this study was to assess MRI capability in the differentiation of pancreatic tumors, peritumoral inflammatory tissue, and normal pancreatic parenchyma by means of DCE-MRI-, DKI-, and IVIM-derived parameters.

Materials and methods

Study population
The ethical local review board of the National Cancer Institute of Naples Pascale Foundation approved this retrospective study and written informed consent for each patient was obtained. We searched the surgical database at our institution from January 2014 to October 2017 and selected 42 patients with pancreatic cancer who underwent surgical resection. The inclusion criteria for the study population were as follows: (a) patients who had pathologically proven pancreatic ductal adenocarcinomas; (b) patients who had undergone both DCE-MRI and DWI; (c) patients who had less than a 1-month interval between imaging and pathologic diagnosis; and (d) availability of diagnostic quality pictures of the resected specimens. The exclusion criteria were as follows: (a) conflict between the imaging-based diagnosis and the pathologically confirmed diagnosis; (b) limitation of pathologic imaging correlation owing to poor image quality; and (c) no available DCE-MRI and DWI.

Thirty-seven patients with pancreatic adenocarcinomas during the study period were selected. Among them, 13 patients were excluded for the following reasons: (a) 8 patients had no available DCE-MRI and DWI study; and (b) 5 patients had more than a 1-month interval between imaging and pathologic diagnosis. Thus, the study group consists of 24 patients [14 men and 10 women, median age 71 years (age range, 53–85 years)]. Characteristics of the study group are summarized in Table 1.

We also searched the radiological database of our institute during the study period and selected a
control group of patients without pancreatic lesions, confirmed by imaging and without history of increased amylases and carbohydrate antigen 19–9 (CA19–9), to reduce spectrum bias. A total of 24 patients [13 men, 11 women; median age, 56 years (age range, 33–78 years)] that underwent DCE-MRI and DWI upper abdomen studies were enrolled. Characteristics of the study control group are summarized in Table 1.

**Lesion confirmation: reference standard**
A pathologist specialized in pancreatic diseases performed histopathologic analysis of resected specimens. Twenty-four patients with pathologically proven pancreatic adenocarcinomas who underwent surgical resection (mean tumor size, 28.0 mm; range 12–52 mm) constituted the study group. Lesion confirmation was based on the pathologic diagnosis of surgically resected pancreatic specimens. Ductal adenocarcinoma composed of epithelial neoplastic cells embedded in a fibrous stroma. Neoplastic cells expressed a specific pattern of immunohistochemically detectable markers: cytokeratins (cytokeratin 7, 8, 13, 18, and 19) and CA19–9.

**MR protocol**
The MR protocol consisted of morphological and functional imaging, including DCE-MRI and DWI sequences. Imaging was performed with a 1.5 T scanner (MAGNETOM Symphony, Siemens Healthcare, Erlangen, Germany) equipped with a phased-array body coil. Patients were placed in a supine, head-first position. A free-breathing axial single-shot echo-planar DWI pulse sequence was performed with tridirectional diffusion gradients with \( b \) values of 0, 50, 100, 150, 400, 800, and 1000 s/mm\(^2\). With regards the DCE-MR

| Description | Numbers (%)/range |
|-------------|-------------------|
| **Pancreatic cancer patients (n = 24)** | |
| Sex | Men 14 (58.3%) |
| | Women 10 (41.7%) |
| Age | 71 years (range, 53–85 years) |
| Histotype | Adenocarcinoma 100% (24/24) |
| Location | |
| Head | 14 (58.3%) |
| Body/tail | 10 (41.6%) |
| Largest diameter | 28.0 mm; range 12–52 mm |
| **Control group patients (n = 24)** | |
| Sex | Men 13 (54.2 %) |
| | Women 11 (45.8%) |
| Age | 56 years; range, 33–78 years |
| Previous neoplastic history | |
| Yes | 9 (37.5%); colorectal cancer (100%) |
| No | 15 (62.5%); hepatic benign lesions |
| Previous Chemotherapy | No one |
| Previous history of pancreatitis | No one |

Table 1. Characteristics of the patients [24 pancreatic cancer and 24 control group patients].
imaging, we obtained 1 sequence before and 120 sequences (without any delay) after intravenous injection of 2 ml/kg of a positive, gadolinium-based paramagnetic contrast medium (Gadobutrol Gd-DTPA, Bayer Pharma AG, Berlin, Germany). The contrast medium was injected using a Spectris Solaris® EP MR pump (MEDRAD Inc., Indianola, PA), with a flow rate of 2 ml/s, followed by a 10 ml saline flush at the same rate. DCE-MRI T1-weighted time-resolved angiography with stochastic trajectories (TWIST) three-dimensional (3D) axial images were acquired to improve temporal resolution (3 s). MRI sequence parameters were reported in Table 2.

**MR image analysis**
Two expert radiologists, in consensus, simultaneously avoiding encircling any distortion artifacts, manually drew regions of interest (ROIs). One radiologist with over 20 years of clinical experience, and one with 8 years of clinical experience in interpreting abdominal MR imaging studies drew ROIs on DCE images with virtual ‘fat suppression’ obtained, subtracting the precontrast from the postcontrast image and then verifying these on the DWI image at the highest b value. For patients with pancreatic cancer, the tumor was contoured slice by slice to obtain the neoplastic volume of interest and we also selected four regions of interest in the contours of the tumor, according to the National Comprehensive Cancer Network guidelines version 3.2017, for pathologic analysis of margins, to obtain the median value of peritumoral inflammatory tissue. For the pancreatic head cancer we drew ROIs inside the superior mesenteric margin (SMA margin) corresponding to the soft tissue directly adjacent to the proximal 3–4 cm of the superior mesenteric artery, and posterior margin corresponding to the tissue between the posterior caudal aspect of the pancreatic head that merges with the SMA margin. For distal lesions, we drew ROIs inside the proximal pancreatic margin corresponding to the pancreatic body along the plane of the section, and the anterior and posterior peri-pancreatic margin corresponding to the tissue between the tumor and adjacent soft tissue.1,28 For patients without pancreatic cancer, we selected four ROIs in the pancreas parenchyma (head, neck, body, and tail) to obtain the median value of pancreatic parenchyma tissue. Features have been computed pixel by pixel to obtain the median value of ROIs.

**DCE-MRI features.** For each voxel, eight time-intensity-curve shape descriptors were computed using an approach previously reported in: maximum signal difference (MSD), the time to peak (TTP), the wash-in slope (WIS), the wash-out slope (WOS), the wash-in intercept (WII), the wash-out intercept (WOI), the WOS/WIS ratio, and the WOI/WII ratio.

DCE-MRI parameters were obtained using in-house prototype software developed in MATLAB R2007a (MathWorks Inc., Natick, MA, US).

### Table 2. MRI sequence parameters.

| Sequence                          | Orientation | TR/TE/FA (ms/ms/deg.) | FOV (mm²)  | Acquisition matrix | Slice thickness/gap (mm) |
|-----------------------------------|-------------|------------------------|------------|--------------------|--------------------------|
| HASTE T2-w                        | Axial       | 1500/90/180            | 380 × 380  | 320 × 320          | 5/0                      |
| FLASH T1-w In-out phase           | Axial       | 160/4.87/70            | 285 × 380  | 192 × 256          | 5/0                      |
| FLASH T1-w out phase              | Axial       | 178/2.3/80             | 325 × 400  | 416 × 412          | 3/0                      |
| DWI                               | Axial       | 7500/91/90             | 340 × 340  | 192 × 192          | 3/0                      |
| VIBE T1-w                         | Axial       | 4.89/2.38/10           | 325 × 400  | 320 × 260          | 3/0                      |
| TWIST T1-w Pre- and postcontrast-agent injection | Axial | 3.01/1.09/25 | 300 × 300  | 256 × 256          | 2/0                      |

- AT, acquisition time; deg., degree; DWI, diffusion-weighted imaging; FA, flip angle; FLASH, fast low-angle shot; FOV, field of view; HASTE, half-Fourier acquisition single-shot turbo spin-echo; TE, echo time; TR, repetition time; VIBE, volumetric interpolated breath-hold examination; -w, weighted.
DWI features. Per each voxel, six features were extracted from DWI data using the monoexponential model, the DKI model and the IVIM model.\(^7,8,15,16,30–38\)

DWI signal decay is most commonly analyzed using the monoexponential model.\(^\text{15,16}\)

\[
ADC = \frac{\ln\left(\frac{S_b}{S_0}\right)}{b}
\]

where \(S_b\) is the MRI signal intensity with diffusion weighting \(b\), \(S_0\) is the nondiffusion-weighted signal intensity.

For a voxel with a large vascular fraction, the MRI data decay can deviate from a monoexponential form, in particular showing a fast decay in the range of low \(b\) values generated by the IVIM effect.\(^\text{15,16,32}\) Thus, in addition to the monoexponential model, a biexponential model was used to estimate the IVIM-related parameters of pseudodiffusivity (\(D_p\), also indicated by \(D^*\)), \(f_p\) and tissue pure diffusivity (\(D_t\)) using the VARiable PROjection approach.\(^\text{38}\)

\[
\frac{S_b}{S_0} = f_p \exp\left(-b \cdot D_p\right) + (1-f_p) \cdot \exp\left(-b \cdot D_t\right)
\]

Moreover, DKI was included in the analysis to obtain the final fitted images (MD and mean of diffusional kurtosis (MK)).

Multi-\(b\) diffusion-weighted images were obtained fitting voxel by voxel, using the diffusion kurtosis signal decay Equation (3) by a two-variable linear least-squares algorithm as used in previous study.\(^\text{20}\)

\[
S(b) = S_0 \exp\left(-b \cdot MD\right) + \frac{1}{6} b^2 \cdot D^2 \cdot MK
\]

In this equation, \(D\) is a corrected \(Dt\); and \(K\) is the excess diffusion kurtosis coefficient. \(K\) describes the degree that molecular motion deviates from the perfect Gaussian distribution.

The difference between \(D\) and ADC is that \(D\) is a corrected form of ADC for use in non-Gaussian circumstances.

The parameters of conventional DWI (ADC), IVIM [\(f_p\), \(D_t\), pseudodiffusivity (\(D_p\))] and DKI (MK and MD) were obtained from the multi-\(b\) DWI data with all measured \(b\) values using the prototype postprocessing software Body Diffusion Toolbox (Siemens Healthcare, Erlangen, Germany).

Statistical analysis

Continuous variables were presented as the median ± standard deviation (SD). All parameters subdivided into the three groups (normal pancreatic parenchyma, peritumoral inflammatory tissue, pancreatic tumor) were compared with each other using the nonparametric Kruskal–Wallis test. The Kruskal–Wallis test was also performed to assess differences statistically significant of the extracted parameters between head and body/tail region of the pancreas.

Receiver operating characteristic (ROC) curves were calculated to characterize each parameter value for evaluating the capability to differentiate pancreatic tumors versus peritumoral inflammatory tissue or pancreatic parenchyma tissue. The optimal cut-off values (obtained according to the maximal Youden index = sensitivity + specificity – 1), the corresponding sensitivity, specificity, positive predictive value, negative predictive value, and accuracy were calculated. McNemar’s test was used to verify statistically significant difference accuracy among parameters. Bonferroni correction was applied for multiple comparisons. A \(p\) value < 0.05 was considered statistically significant. The Statistics Toolbox of MATLAB R2007a (MathWorks Inc., Natick, MA, US) was used to perform statistical analysis.

Results

Table 3 reports the median value and SD value for pancreatic tumor, peritumoral inflammatory tissue and pancreatic parenchyma tissue.

There were statistically significant differences in median values among the three groups observing the Kruskal–Wallis test for MD, \(f_p\), and \(D_p\), while there were no significant differences among these groups for dynamic parameters (see also Figure 1). WIS showed no statically significant difference for median values in three groups (\(p\) value = 0.06): 3.75 ± 17.84 in pancreatic parenchyma tissue; 13.14 ± 20.17 in peritumoral
inflammatory tissue; and $20.91 \pm 25.49$ in pancreatic tumor. MD had a median value of $2843.20 \pm 728.35 \times 10^{-3} \text{mm}^2/\text{s}$ in normal pancreatic parenchyma while had a median value of $3211.10 \pm 796.28 \times 10^{-3} \text{mm}^2/\text{s}$ for peritumoral inflammatory tissue and of $1849.50 \pm 603.95 \times 10^{-3} \text{mm}^2/\text{s}$ in pancreatic tumor. fp had a median value of $22.50 \pm 9.04\%$ in normal pancreatic parenchyma, while having a median value of $27.71 \pm 15.46\%$ for peritumoral inflammatory tissue, and $14.42 \pm 8.15\%$ in pancreatic tumor. Dp had a median value of $135.60 \pm 57.30 \times 10^{-5} \text{mm}^2/\text{s}$ in normal pancreatic parenchyma while had a median value of $172.30 \pm 87.07 \times 10^{-5} \text{mm}^2/\text{s}$ for peritumoral inflammatory tissue and of $112.80 \pm 56.6 \times 10^{-5} \text{mm}^2/\text{s}$ in pancreatic tumor.

No statistically significant differences were observed in median values of extracted parameters between head and body/tail region of the pancreas using the Kruskal–Wallis test ($p$ value $> 0.05$).

Table 4 reports the diagnostic accuracy of MRI-extracted parameters in discriminating normal pancreatic parenchyma plus peritumoral inflammatory tissue versus pancreatic tumor. The bolded parameters having high accuracy and area under ROC curve (AUC) are WOI, WII, ADC, MD, fp, and Dp, showing an accuracy $\geq 65\%$ and AUC $> 0.6$. MD had the best results with an accuracy of $84\%$ ($p$ value $< 0.05$ using McNemar’s test) and AUC $= 0.85$.

Table 4 also reports the diagnostic accuracy of MR-extracted parameters in discriminating normal pancreatic parenchyma versus peritumoral inflammatory tissue or pancreatic tumor. The parameters having high accuracy and AUC are again emphasized in bold. WII and WOI/WII showed an accuracy $\geq 63\%$ and AUC $\geq 0.6$. There were no statistically significant differences ($p$ value $> 0.05$ using McNemar’s test) between parameter accuracy, however DCE-MRI WII had the highest accuracy (68\%) and AUC (0.60).

Table 5 reports the diagnostic accuracy of MR-extracted parameters in discriminating normal pancreatic parenchyma versus pancreatic tumor and those with high accuracy and AUC are bolded. WII, MD, fp, and Dp showed an accuracy $> 70\%$ and AUC $> 0.6$. MD had the best accuracy of 78\% ($p$ value $< 0.05$ using McNemar’s test) and AUC of 0.82.
Moreover, Table 5 reports the diagnostic accuracy of MR-extracted parameters in discrimination of normal pancreatic parenchyma versus peritumoral inflammatory tissue. There were no statistically significant differences between parameter accuracy (p value > 0.05 using McNemar’s test); however, DCE-MRI WOI/WII had the best accuracy (67%) and AUC (0.67).

Finally, Table 5 reports the diagnostic accuracy of MR-extracted parameters in discriminating peritumoral inflammatory tissue versus pancreatic tumor. WII, MD, fp, and Dp showed an accuracy ≥ 72% and AUC > 0.6. MD had the best accuracy at 83% (p value < 0.05 at McNemar test) and AUC of 0.89.

Figures 2 and 3 show representative cases of pancreatic tumor with hyperintense signal on T2-weighted sequence, isohypointense signal during the portal phase of the contrast study, restricted diffusion on DWI at $b = 1000 \text{s/mm}^2$ and hypointense signal on the ADC map.

**Discussion**

DCE-MRI accuracy in the evaluation of pancreatic cancer remains unclear. In pancreatic adenocarcinoma, poorly represented microvascular components could be clarified by vessel functional impairment often observed in tumors, and by the presence of a prominent stromal matrix that embeds vessels. In addition, activated pancreatic stellate cells yield increasing fibrous stroma in tumor central areas, compressing blood vessels, leading to changes in vascularity and perfusion.\(^{39,40}\) Several studies evaluated the feasibility of DCE-MRI for the characterization of solid pancreatic diseases.\(^{39,40-9}\)

Kim and colleagues\(^{39}\) evaluated 24 patients with pancreatic cancers; 8 with pancreatic neuroendocrine tumors (PNETs), 3 with chronic pancreatitis, and 10 with a normal pancreas. They showed that $K_{\text{trans}}$ [transfer constant by extravascular extracellular space (EES) to plasma], kep (transfer constant by plasma versus EES), and iAUC (initial AUC) values in patients with pancreatic cancer were significantly lower than in patients with a normal pancreas. In addition, kep values of PNETs and normal pancreas and $K_{\text{trans}},$ kep, and iAUC values of pancreatic cancers and PNETs differed significantly. Bali and colleagues\(^{40}\) evaluated 28 patients with surgically resectable pancreatic lesions. They showed that $K_{\text{trans}}$ values were significantly lower in primary malignant tumors compared with benign lesions and nontumoral pancreatic tissue; plasma volume fraction was significantly higher in primary malignant tumors compared with nontumoral pancreatic tissue. Sensitivity and specificity for fibrosis detection were 65% and 83%, and 76% and 83% for the $K_{\text{trans}}$ one-compartment two-compartment models, respectively.

We evaluated semiquantitative descriptors of the contrast-agent time course such as MSD, TTP, WIS, WOS, WII, WOI, the WOS/WIS ratio, and the WOI/WII ratio. Our findings showed that there were no differences among three groups for dynamic parameters except a statistically nonsignificant difference for WIS comparable with $K_{\text{trans}}.\(^{30}\)

Diffusion parameters can be assessed by DWI.\(^{38}\) The IVIM approach allows separating blood volume fraction (perfusion) by diffusion and microstructural information.\(^{35,36}\) Several studies reported that IVIM is a promising tool in
Table 4. Diagnostic accuracy of MRI-extracted parameters.

|                  | AUC  | SEN  | SPEC | PPV  | NPV  | Accuracy | Cut off |
|------------------|------|------|------|------|------|----------|---------|
| Discriminating normal pancreatic parenchyma plus peritumoral inflammatory tissue versus pancreatic tumor |      |      |      |      |      |          |         |
| MSD              | 0.45 | 0.09 | 1.00 | 1.00 | 0.36 | 0.40     | 95.51   |
| TTP              | 0.53 | 0.62 | 0.52 | 0.72 | 0.41 | 0.59     | 25.01   |
| WOS              | 0.51 | 0.78 | 0.35 | 0.70 | 0.44 | 0.63     | −5.88   |
| WOI              | 0.67 | 0.67 | 0.65 | 0.79 | 0.50 | **0.66** | 48.83   |
| WIS              | 0.35 | 1.00 | 0.04 | 0.67 | 1.00 | 0.68     | −44.80  |
| WII              | **0.65** | 0.53 | 0.91 | 0.92 | 0.50 | **0.66** | 33.47   |
| WOS/WIS          | 0.50 | 0.33 | 0.74 | 0.71 | 0.36 | 0.47     | 0.08    |
| WOI/WII          | 0.53 | 0.64 | 0.52 | 0.73 | 0.43 | 0.60     | −0.92   |
| ADC              | **0.63** | 0.58 | 0.78 | 0.84 | 0.49 | **0.65** | 1330.97 |
| MK               | 0.40 | 0.76 | 0.30 | 0.68 | 0.39 | 0.60     | 996.76  |
| MD               | **0.85** | 0.91 | 0.70 | 0.85 | 0.80 | **0.84** | 2168.31 |
| fp               | 0.83 | 0.80 | 0.78 | 0.88 | 0.67 | **0.79** | 199.85  |
| Dt               | 0.57 | 0.44 | 0.78 | 0.80 | 0.42 | 0.56     | 1253.63 |
| Dp               | **0.70** | 0.93 | 0.39 | 0.75 | 0.75 | **0.75** | 68.92   |
| Discriminating normal pancreatic parenchyma versus peritumoral inflammatory tissue or pancreatic tumor |      |      |      |      |      |          |         |
| MSD              | 0.50 | 0.09 | 1.00 | 1.00 | 0.70 | 0.71     | 104.04  |
| TTP              | 0.52 | 0.50 | 0.70 | 0.44 | 0.74 | 0.63     | 38.01   |
| WOS              | 0.50 | 0.64 | 0.50 | 0.38 | 0.74 | 0.54     | −1.57   |
| WOI              | 0.60 | 0.86 | 0.37 | 0.40 | 0.85 | 0.53     | 33.56   |
| WIS              | 0.43 | 0.82 | 0.28 | 0.35 | 0.76 | 0.46     | −5.19   |
| WII              | **0.60** | 0.50 | 0.76 | 0.50 | 0.76 | **0.68** | 36.70   |
| WOS/WIS          | 0.45 | 0.36 | 0.80 | 0.47 | 0.73 | 0.66     | 0.28    |
| WOI/WII          | **0.63** | 0.73 | 0.59 | 0.46 | 0.82 | **0.63** | 0.17    |
| ADC              | 0.52 | 0.55 | 0.61 | 0.40 | 0.74 | 0.59     | 1331.67 |
| MK               | 0.47 | 0.82 | 0.30 | 0.36 | 0.78 | 0.47     | 996.76  |
| MD               | 0.58 | 0.86 | 0.41 | 0.41 | 0.86 | 0.56     | 2214.80 |
| fp               | 0.55 | 0.82 | 0.39 | 0.39 | 0.82 | 0.53     | 167.83  |
| Dt               | 0.57 | 0.59 | 0.63 | 0.43 | 0.76 | 0.62     | 1147.04 |
| Dp               | 0.53 | 1.00 | 0.26 | 0.39 | 1.00 | 0.50     | 68.92   |

Diagnostic accuracy of MRI-extracted parameters in discriminating normal pancreatic parenchyma plus peritumoral inflammatory tissue versus pancreatic tumor, and in discrimination of normal pancreatic parenchyma versus peritumoral inflammatory tissue or pancreatic tumor. Parameters having high accuracy and AUC are in bold type.

ACC, accuracy; AUC, area under curve; Dp, pseudodiffusivity; Dt, tissue pure diffusivity; fp, perfusion fraction; MD, mean diffusivity; MK, mean of diffusional kurtosis; MRI, magnetic resonance imaging; MSD, maximum signal difference; NPV, negative predictive value; PPV, positive predictive value; SEN, sensitivity; SPEC, specificity; TTP, time to peak; WII, wash-in intercept; WIS, wash-in slope; WOS, wash-out slope; WOI, wash-out intercept.
Table 5. Diagnostic accuracy of MRI-extracted parameters.

|                  | AUC | SEN | SPEC | PPV | NPV | Accuracy | Cut off |
|------------------|-----|-----|------|-----|-----|----------|---------|
| MSD              | 0.47| 0.14| 0.96 | 0.75| 0.54| 0.56     | 92.21   |
| TTP              | 0.54| 0.59| 0.61 | 0.59| 0.61| 0.60     | 31.02   |
| WOS              | 0.51| 0.64| 0.48 | 0.54| 0.58| 0.56     | −1.54   |
| WOI              | 0.68| 0.86| 0.48 | 0.61| 0.79| 0.67     | 30.87   |
| WIS              | 0.36| 1.00| 0.04 | 0.50| 1.00| 0.51     | −44.80  |
| WII              | 0.67| 0.55| 0.91 | 0.86| 0.68| **0.73** | 33.49   |
| WOS/WIS          | 0.47| 0.36| 0.78 | 0.62| 0.56| 0.58     | 0.17    |
| WOI/WII          | 0.59| 0.77| 0.52 | 0.61| 0.71| 0.64     | −0.92   |
| ADC              | 0.61| 0.55| 0.78 | 0.71| 0.64| 0.67     | 1330.99 |
| MK               | 0.42| 0.82| 0.30 | 0.53| 0.64| 0.56     | 997.00  |
| MD               | **0.82**| 0.86| 0.70 | 0.73| 0.84| **0.78** | 2168.48 |
| fp               | **0.79**| 0.82| 0.70 | 0.72| 0.80| **0.76** | 167.81  |
| Dt               | 0.59| 0.55| 0.74 | 0.67| 0.63| 0.64     | 1197.58 |
| Dp               | 0.67| 1.00| 0.39 | 0.61| 1.00| 0.69     | 68.91   |

Discriminating normal pancreatic parenchyma versus peritumoral inflammatory tissue

|                  | AUC | SEN | SPEC | PPV | NPV | Accuracy | Cut off |
|------------------|-----|-----|------|-----|-----|----------|---------|
| MSD              | 0.53| 0.73| 0.43 | 0.55| 0.63| 0.58     | 30.24   |
| TTP              | 0.51| 0.50| 0.70 | 0.61| 0.59| 0.60     | 38.01   |
| WOS              | 0.50| 0.64| 0.52 | 0.56| 0.60| 0.58     | −1.63   |
| WOI              | 0.53| 0.27| 0.91 | 0.75| 0.57| 0.60     | 105.40  |
| WIS              | 0.50| 0.82| 0.35 | 0.55| 0.67| 0.58     | −5.20   |
| WII              | 0.52| 0.73| 0.43 | 0.55| 0.63| 0.58     | 8.50    |
| WOS/WIS          | 0.44| 0.36| 0.83 | 0.67| 0.58| 0.60     | 0.28    |
| WOI/WII          | **0.67**| 0.73| 0.61 | 0.64| 0.70| **0.67** | 0.18    |
| ADC              | 0.43| 0.73| 0.35 | 0.52| 0.57| 0.53     | 1139.20 |
| MK               | 0.53| 1.00| 0.13 | 0.52| 1.00| 0.56     | 600.80  |
| MD               | 0.35| 1.00| 0.00 | 0.49| –   | 0.49     | 1479.50 |
| fp               | 0.30| 1.00| 0.04 | 0.50| 1.00| 0.51     | 33.76   |
| Dt               | 0.54| 0.59| 0.61 | 0.59| 0.61| 0.60     | 1147.04 |
| Dp               | 0.39| 1.00| 0.13 | 0.52| 1.00| 0.56     | 67.22   |

(Continued)
Kang and colleagues evaluated the diagnostic performance of ADC- and IVIM-derived parameters to distinguish pancreatic tumors, chronic pancreatitis, and normal pancreas and to characterize intraductal papillary mucinous neoplasms (IPMNs). They reported that incoherent microcirculation (Dfast) and fp values of PDACs were significantly lower than those of normal pancreas, chronic pancreatitis, and NETs. In differentiating PDACs from NETs, fp and Dfast showed a significant difference. Malignant IPMNs had significantly lower ADC and slow component of diffusion values, while benign IPMNs had significantly higher Dfast and fp values. In ROC analysis, fp showed the highest ROC AUC in distinguishing malignant from benign IPMNs. They concluded that perfusion might be a more important factor than diffusion in discriminating PDAC from normal pancreas, chronic prostatitis and NETs. In addition, fp showed the highest AUC by ROC analysis in differentiating malignant from benign IPMNs among ADC- and IVIM-derived parameters. Klau and colleagues investigated the correlation between IVIM-derived parameters and histologically determined microvascularity in PDACs and PNETs. They showed that blood volume fraction fp was significantly lower in PDACs compared with PNETs, and that the Dt was significantly higher in PDAC.

In our study, we evaluated ADC and the IVIM-related parameters (Dp, fp and Dt), so the kurtosis coefficient that is linked to the deviation of
Figure 2. Female, 43 years, body pancreatic adenocarcinoma. The lesion shows hyperintense signal in T2-w sequence: (a) HASTE T2-w in axial plane with isohypointense signal during portal phase of contrast study; (b) VIBE FS in axial plane. In DWI (c) $b = 1000 \text{s/mm}^2$; the lesion shows restricted diffusion with hypointense signal on the ADC map (d).

ADC, apparent diffusion coefficient; DWI, diffusion-weighted imaging; HASTE, half-Fourier acquisition single-shot turbo spin-echo; T2-w, T2 weighted; VIBE FS, volumetric interpolated breath-hold examination fat saturated.

Figure 3. Female, 45 years, tail pancreatic adenocarcinoma. The lesion shows hyperintense signal in T2-w sequence (a) HASTE T2-w in axial plane with isohypointense signal during portal phase of contrast study; (b) VIBE FS in axial plane. In DWI (c) $b = 1000 \text{s/mm}^2$; the lesion shows restricted diffusion with hypointense signal on the ADC map (d).

ADC, apparent diffusion coefficient; DWI, diffusion-weighted imaging; HASTE, half-Fourier acquisition single-shot turbo spin-echo; T2-w, T2 weighted; VIBE FS, volumetric interpolated breath-hold examination fat saturated.
tissue diffusion from a Gaussian model, and the Dt with the correction of non-Gaussian bias by DKI. Recently, DKI was used to assess therapy response in different kinds of tumors.\textsuperscript{43–45} According to our results, there was a statistically significant difference in median values among the three groups observed by Kruskal–Wallis test for MD, fp and Dp. In our study, the perfusion-related factors of PDAC, fp and Dp, and MD of DKI, differed from those seen in patients with normal pancreatic parenchyma and in peritumoral tissue, and showed better diagnostic performance than did ADC. Although the differential diagnosis of PDAC and normal pancreatic parenchyma is usually considered straightforward, overlap in imaging features can make this differentiation difficult. Therefore, the significantly different perfusion-related factors of PDAC and normal pancreatic parenchyma might be helpful for determining the most accurate diagnosis. Increased fp and MD in peritumoral inflammation seem to suggest that DWI-derived parameters fit in the anticipated physiologic phenomena. Our results support the hypothesis that the kurtosis effect could have a better performance in differentiating pancreatic tumors, peritumoral inflammatory tissue, and normal pancreatic parenchyma, although our data were acquired with a maximum \( b \) value of 1000 s/mm\(^2\). In general, in brain applications, very high \( b \) values are recommended for the assessment of a non-Gaussian kurtosis effect;\textsuperscript{1,20} while in abdominal applications, for the lower signal-to-noise ratio and lower T2-relaxation times, very high \( b \) values are not usually applied. Recently, various authors have shown that kurtosis effects could be detectable in abdominal and whole-body applications also using, as maximum, \( b \) values of 800 s/mm\(^2\) or less at 3T.\textsuperscript{1,7,24,27} We applied multiple \( b \) values with a maximum of 1000 s/mm\(^2\) that, coupled with the use of a parallel imaging factor, resulted in images with acceptable signal-to-noise ratio (SNR) at 1.5T.

Some limits in our study must be highlighted. First, the retrospective nature of this study. A larger number of patients will be needed to confirm our results. We believe further studies with a larger study population are warranted for its validation. Second, we did not assess the interobserver variability regarding the drawing of ROIs. However, we used median values both for DCE-MRI and for DWI-derived parameters. Third, we used only 4 \( b \) values <200 s/mm\(^2\) to estimate IVIM diffusion parameters, which could be seen as a weakness; however, we used a robust algorithm, the VARiable PROjection approach, superior to the conventional Levenberg–Marquardt algorithm for curve fitting and diffusion parameters estimation of intravoxel incoherent motion method.

**Conclusion**

IVIM and DKI-derived parameters could be helpful in the discrimination of normal pancreatic parenchyma tissue, perilesional inflammation, and pancreatic tumor. Overall, MD of DKI is the parameter that allows the best classification among normal pancreatic parenchyma tissue, perilesional inflammation, and pancreatic tumor.

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**Conflict of interest statement**

Robert Grimm is an employee of Siemens Healthcare.

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