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ORIGINAL RESEARCH

Pancreatic neuroendocrine tumor: prediction of the tumor grade using magnetic resonance imaging findings and texture analysis with 3-T magnetic resonance

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Purpose: The purpose of this study was to evaluate the performance of magnetic resonance imaging (MRI) findings and texture parameters for prediction of the histopathologic grade of pancreatic neuroendocrine tumors (PNETs) with 3-T magnetic resonance.

Patients and methods: PNETs are classified into Grade 1 (G1), Grade 2 (G2), and Grade 3 (G3) tumors based on the Ki-67 proliferation index and the mitotic activity. A total of 77 patients with pathologically confirmed PNETs met the inclusion criteria. Texture analysis (TA) was applied to T2-weighted imaging (T2WI) and diffusion-weighted imaging (DWI) maps. Patient demographics, MRI findings, and texture parameters were compared among three different histopathologic subtypes by using Fisher's exact tests or Kruskal–Wallis test. Then, logistic regression analysis was adopted to predict tumor grades. ROC curves and AUCs were calculated to assess the diagnostic performance of MRI findings and texture parameters in prediction of tumor grades.

Results: There were 31 G1, 29 G2, and 17 G3 patients. Compared with G1, G2/G3 tumors showed higher frequencies of an ill-defined margin, a predominantly solid tumor type, local invasion or metastases, hypo-enhancement at the arterial phase, and restriction diffusion. Four T2-based (inverse difference moment, energy, correlation, and differenceEntropy) and five DWI-based (correlation, contrast, inverse difference moment, maxintensity, and entropy) TA parameters exhibited statistical significance among PNETs (*P*<0.001). The AUCs of six predicting models on T2WI and DWI ranged from 0.703–0.989.

Conclusion: Our data indicate that MRI findings, including tumor margin, texture, local invasion or metastases, tumor enhancement, and diffusion restriction, as well as texture parameters can aid the prediction of PNETs grading.

Keywords: magnetic resonance imaging, neuroendocrine tumor, pancreas, texture analysis

Introduction

Pancreatic neuroendocrine tumors (PNETs) are rare neoplasms that arise from pluripotent stem cells in ductal epithelium, accounting for less than 3% of all pancreatic neoplasms.¹ However, the incidence of PNETs has increased over the past decade, which may be due to the increased awareness, improved diagnostic imaging modalities, and the revision of its definition.²⁻⁴ In the WHO 2017 classification, PNETs are categorized based on the Ki-67 proliferation index and mitotic activity. They are divided into three tumor grades: Grade 1 (G1), Grade 2 (G2), and Grade 3 (G3) tumors.⁵ Notably, the choice of treatment of PNETs is related to tumor grades. Surgical resection should

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Construction of the full terms of this license are available at https://www.dovepress.com/terms. by and incorporate the (creative Commons Attribution — Non Commercial (unported, v3.0) License (http://creative.commons.org/licenses/by-nc/3.0). By accessing the work you hereby accept the Terms. Non-commercial uses of the work are permitted without any further permission from Dove Medical Press Limited, provided the work is properly attributed. For permission for commercial use of this work, please see paragraphs 4.2 and 5 of our Terms (https://www.dovepress.com/terms.php). always be proposed for PNETs if technically feasible, even in some cases with limited metastases. In addition, therapies with somatostatin analogs or ablation are reserved for those with unresectable or residual disease.^{1,6}

The typical imaging findings of PNETs describe them as well-defined solid masses with relatively intense enhancement at the arterial or portal venous phase, absence of ductal dilatation, and lack of vascular encasement.^{7,8} However, some recent studies demonstrated that up to 41.5% of PNETs may show arterial hypo-enhancement, especially for G2/G3 tumors.^{9,10} There are differences in tumor margin, enhancement pattern, portal enhancement ratio, and duct dilatation among PNETs.¹¹⁻¹³ Diffusion-weighted imaging (DWI) or the apparent diffusion-coefficient (ADC) map obtained from DWI is an effective imaging method in assessment of tumors or pathological treatment response to chemoradiation.14,15 Lotfalizadeh et al¹⁶ found that mean ADC and diffusion coefficient (D) became significantly lower as tumor grade of PNETs increased. Although several studies showed that computed tomography (CT) or magnetic resonance imaging (MRI) characteristics can predict tumor grade of PNET, accurate prediction of the PNET grade by CT or MRI is still limited.

Texture analysis (TA) that can extract, analyze, and interpret quantitative imaging features has been widely used in imaging fields. It allows objective quantitative assessment of lesion or tissue heterogeneity. Early evidence suggests that TA has potential for diagnosis, characterization, tumor staging, and therapy response assessment in oncologic practice.^{17–22} Several reports have investigated the value of CT TA in predicting the histopathologic grade of PNETs.^{23,24} However, to the best of our knowledge, few studies have shown the potential value of MRI TA in predicting the histopathologic grade of PNETs. TA of T2-weighted imaging (T2WI) from MRI has been used in many cancers.^{25,26} Despite the potential variability in T2 signal intensity, a previous study reported highly reproducible results of texture features from three different MRI scanners.^{27,28}

The purpose of our study was to investigate the utility of MRI findings and texture features in predicting the histopathologic grade of PNETs.

Patients and methods Patients

This study was approved by the Ethical Committee of the Affiliated Hospital of Nanjing University of Chinese Medicine and informed patient consent requirement was waived due to its retrospective nature. Patient data confidentiality was protected in accordance with the Declaration of Helsinki principles. In our study, 101 consecutive patients with pathologically confirmed PNETs between December 2013 and May 2017 were identified. Inclusion criteria were as follows: 1) patients who underwent surgical excision with histopathologic examination; 2) patients who underwent MRI examinations including DWI performed within a period of 30 days before surgery; 3) patients did not receive local treatment or systemic chemotherapy before surgery. The exclusion criteria were as follows: 1) intraoperative biopsy-proven PNETs were excluded because they could not be correctly classified according to the WHO classification (n=6); 2) no MRI examination before surgery (n=8); 3) the quality of the MRI images was poor due to movement artifacts or susceptibility artifacts (n=4); 4) patients with multiple PNETs in which the pathology report was not clear regarding the tumor grade (n=2); 5) tumors did not restrict diffusion or were too small to be depicted on DWI images (n=4). Finally, a total of 77 patients (34 women and 43 men with a mean age of 53.32±11.75 years) were enrolled in our study (Figure 1).

MRI protocol

All preoperative MRI scanning was performed using a 3.0-T MRI (Sigma HDx; GE Medical Systems, Milwaukee, WI, USA) system with an eight-channel phased-array torso coil. All patients fasted for 8 hours prior to MRI examination. MRI sequences included T1-weighted fat-suppressed liver acquisition with volume acceleration (LAVA) sequence (repetition time 3,100 ms, echo time 15 ms, imaging duration 1-2 minutes; slice thickness 5 mm, interslice gap 2 mm; flip angle 12°, matrix 384×256, field of view 22 cm × 22 cm); fast spin-echo T2-weighted fat-suppressed sequence (repetition time 6,000 ms, echo time 80 ms, imaging duration 2-3 minutes; slice thickness 5 mm, interslice gap 2 mm; flip angle 90°, matrix 384×256, field of view 22 cm × 22 cm), and DWI using respiration-triggered single shot echo-planar sequence (repetition time 8,000 ms, echo time 60 ms; slice thickness 5 mm, interslice gap 2 mm; flip angle 90°, matrix 196×133, field of view 36 cm × 30 cm; b value of 0 and 1,000 s/mm²). T1-weighted contrast-enhanced sequences were performed at the pancreatic parenchyma (35 seconds scanning delay), portal venous (70 seconds scanning delay), and delayed (240 seconds scanning delay) phase with the contrast media of Gadopentetate dimeglumine (GE Healthcare; 0.2 mmol/kg) at a rate of 2.5 mL/s.

Image analysis

Two radiologists (with 8 and 9 years experience in abdominal imaging) with no prior knowledge of detailed clinical



Figure I Flowchart of study group inclusion process.

Abbreviations: DWI, diffusion-weighted imaging; MR, magnetic resonance; MRI, magnetic resonance imaging; PNETs, pancreatic neuroendocrine tumors.

information and histopathological results, reviewed MR images independently. Consensus was reached through discussion or by referral to a third radiologist (with 13 years experience in abdominal imaging). The following imaging findings were analyzed: tumor size, location, margin (welldefined vs ill-defined), texture, local invasion or metastases, pancreatic atrophy, pancreatic duct dilatation, signal intensity of the tumor related to the adjacent parenchyma (hypo-, iso- or hyperintense), tumor enhancement (hyper- or hypoenhancement), and presence of restricted diffusion.

A well-defined margin indicated a smooth and visible margin, while an ill-defined margin indicated spiculation or infiltration on >90° of the tumor parameter. Tumor texture was divided into solid (an enhancing solid component of more than 90%), predominantly solid (an enhancing solid component of 50%-90%), and predominantly cystic (an enhancing solid component of <50%).²⁹ Local invasion or metastases referred to extrapancreatic localizations, including vascular or lymph node invasion, spleen invasion, or liver metastases, etc. Pancreatic atrophy was reported when more than the expected loss in pancreatic volume was noted. Pancreatic duct dilatation was defined as a main pancreatic duct of 4 mm or greater. Hyper- or hypo-enhancement was defined as solid components of the tumor showing greater or lower enhancement level compared to the adjacent pancreatic parenchyma at the arterial phase. Restricted diffusion within the tumor was defined as the lesion showing hyper-intensity on DWI (b=1,000 mm²/s) with corresponding hypo-intensity on the ADC maps.25

MRI TA

Preoperative axial T2WI and DWI (b value =1,000 mm²/s) maps were exported in DICOM format from the picture archiving and communication system to Omni-Kinetics software (Omni-Kinetics Version V2.0.10, GE Healthcare) to extract texture features. Prior to TA, image quality had been visually evaluated to avoid severe artifacts and mismatches between images. Two radiologists manually drew the regions of interest (ROIs) along the margin of the tumor avoiding peripheral fat, artifacts, and blood vessels in order to get rid of partial volume effect on both T2WI and DWI (Figure 2). For iso-dense tumors on T2WI maps, contrast-enhanced sequences and DWI were referred for tumor margin identification. For each lesion, we drew the ROIs by slice-by-slice segmentation of the whole tumor on each sequence. A total of 68 features were automatically extracted from T2WI or DWI maps. They were separated into five categories: 1) first-order statistics; 2) histogram; 3) gray-level co-occurrence matrix; 4) Haralick; and 5) run-length matrix.

Pathological analysis

The tumor specimens were fixed in 10% formalin for 24 hours. Then the specimens were embedded in paraffin and sectioned for H&E staining (each slice was 4 μ m thick). Immunohistochemical analysis of CgA, NSE, and Syn was used to confirm the histological diagnosis by a pathologist (with 18 years experience in biliary and pancreatic pathology). Pathological tumor grades of PNETs were determined by counting the number of mitoses per 10 HPF and detecting



Figure 2 Example of T2WI images showing segmentation of pancreatic neuroendocrine tumors.

Notes: Manual segmentation of the tumor was delineated by stacking up regions of interest slice-by-slice on the axial T2WI images (A). Histogram of the tumor on T2WI images (B).

Abbreviation: T2WI, T2-weighted imaging.

the Ki-67 proliferation index according to the WHO 2017 classification.⁵ PNET G1: <2 mitoses per 10 HPF, Ki-67 index <2; PNET G2: 2–20 mitoses per 10 HPF, Ki-67 index 3%–20%; G3:>20 mitoses per 10 HPF, Ki-67 index >20%.

Statistical analysis

Quantitative data were shown as mean \pm SD and qualitative data were expressed as numbers (percentage). Patient demographics and MRI findings were compared among three different histopathologic subtypes by using Fisher's exact tests. Inter-observer agreement for texture features was calculated using interclass correlation coefficient (ICC). ICC values $\leq 40\%$ indicated poor agreement; 41%–60%, moderate agreement; 61%-75%, substantial agreement; and >75%, perfect agreement. T2WI or DWI texture parameters were compared a cross all three histological grades of PNETs (eg, G1 vs G2, G1 vs G3, or G2 vs G3) by using Kruskal-Wallis test with Bonferroni correction (P=0.5/3=0.017). Parameters with a *P*-value <0.017 were used as the input variables for logistic regression analysis. In logistic regression analysis, a backward stepwise selection mode was adopted, with iterative entry of variables based on test results. Multi-collinearity features were eliminated when the variance inflation factor (VIF) was larger than ten.³⁰ ROC curves were performed to investigate the diagnostic performance of logistic regression model in PNET grading. The AUC was calculated for each logistic regression model, and optimal threshold was obtained by maximizing the Youden index (add the sensitivity of a diagnostic test to the specificity of the same diagnostic test, then subtract 100 from that value). All statistical analyses were performed using SPSS software (version 20.0; IBM

Corporation, Armonk, NY, USA). Unless otherwise specified, P < 0.05 was considered to indicate a statistical significance.

Results

Seventy-seven patients with pathologically confirmed PNETs were analyzed (Table 1). The histopathologic results showed that, of the 77 PNETs, 31 tumors (40.3%) were classified as G1, 29 (37.7%) as G2, and 17 (22%) as G3. Among these 77 neoplasms, 44.2% (34/77) were found at the head or neck of the pancreas, 27.2% (21/77) at the body, and 28.6% (22/77) at the tail. The mean size of all tumors (mean \pm SD) was 2.42 \pm 0.67 cm. There was a significant difference among G1, G2, and G3 PNETs with respect to the mean size of tumors (1.88 \pm 0.39 cm vs 2.52 \pm 0.47 cm vs 3.23 \pm 0.43 cm, *P*<0.001).

MRI findings

Table 2 summarizes the MRI findings of PNETs according to the tumor grade. There were significant differences among G1, G2, and G3 PNETs with respect to ill-defined margin (29% vs 58.6% vs 70.6%, P=0.01) and local invasion or metastases (0 vs 31% vs 41.2%, P<0.05). G3 demonstrated a higher frequency of a predominantly solid tumor type compared with G1/G2 tumors (64.7% vs 37.9% vs 22.6%, P=0.016). There was a significant difference of tumor enhancement at the arterial phase among G1, G2, and G3 PNETs (87.1% vs 41.4% vs 0, P<0.05). Figures 3 and 4 are typical cases of G1 and G3 PNETs.

MRITA

ICCs regarding texture features were calculated, and those features with ICCs larger than 0.75 were kept. T2WI and

Characteristics	PNETs	P-value		
	Grade I (n=31)	Grade 2 (n=29)	Grade 3 (n=17)	
Age (years) ^a	52.5±10.0	53.5±13.4	54.7±12.2	0.611
Sex ^b				0.296
Male	11 (35.5%)	13 (44.8%)	10 (58.8%)	
Female	20 (64.5%)	16 (55.2%)	7 (41.2%)	
Tumor location ^b				0.206
Head or neck	14 (45.2%)	16 (55.2%)	4 (23.5%)	
Body	7 (22.6%)	6 (20.7%)	8 (47.1%)	
Tail	10 (32.2%)	7 (24.1%)	5 (29.4%)	
Size (cm) ^a	1.88±0.39 cm	2.52±0.47 cm	3.23±0.43 cm	<0.05
Clinical symptom				
Abdominal pain	13 (41.9%)	15 (51.7%)	12 (70.6%)	0.177
Abdominal bloating or diarrhea	5 (16.1%)	5 (17.2%)	2 (11.8%)	0.925
Yellow urine or icterus	3 (9.7%)	4 (13.8%)	I (5.9%)	0.801
Marasmus	0 (0)	I (3.4%)	2 (11.8%)	0.175
Others	8 (25.8%)	4 (12.9%)	I (5.9%)	0.202
Asymptomatic	6 (19.4%)	2 (6.9%)	2 (11.8%)	0.422

Table I Demographic characteristics of patients with pancreatic neuroendocrine tumors (PNETs)

Notes: ^aData are mean ± SD. P-value was calculated with one-way analysis of variance. ^bData are number of patients, with the percentage in parentheses. P-value was calculated with chi-squared or Fisher's exact test.

Features	Grade				
	Total	Grade I (n=31)	Grade 2 (n=29)	Grade 3 (n=17)	
Tumor margin ^a					0.01
Well-defined	39	22 (71%)	12 (41.4%)	5 (29.4%)	
III-defined	38	9 (29%)	17 (58.6%)	12 (70.6%)	
Texture ^a					0.016
Solid	48	24 (77.4%)	18 (62.1%)	6 (35.3%)	
Predominantly solid	29	7 (22.6%)	11 (37.9%)	11 (64.7%)	
Predominantly cystic	0	0	0	0	
Local invasion or metastases ^a					<0.05
Present	16	0 (0%)	9 (31%)	7 (41.2%)	
Absent	61	31 (100%)	20 (69%)	10 (58.8%)	
Pancreatic atrophy ^a					0.162
Present	20	5 (16.1%)	8 (27.6%)	7 (41.2%)	
Absent	57	26 (83.9%)	21 (72.4%)	10 (58.8%)	
Pancreatic duct dilatation ^a					0.102
Present	14	3 (9.7%)	5 (17.2%)	6 (35.3%)	
Absent	63	28 (90.3%)	24 (82.8%)	11 (64.7%)	
TI-weighted imageª					-
lsointense/hypointense	77	31 (100%)	29 (100%)	17 (100%)	
Hyperintense	0	0	0	0	
T2-weighted imageª					0.950
Hypointense	8	3 (9.7%)	3 (10.3%)	2 (11.8%)	
lsointense	10	3 (9.7%)	5 (17.2%)	2 (11.8%)	
Hyperintense	59	25 (80.6%)	21 (72.5%)	13 (76.4%)	
Tumor enhancement					<0.05
Hyper	39	27 (87.1%)	12 (41.4%)	0 (0)	
Нуро	38	4 (12.9%)	17 (58.6%)	17 (100%)	
Diffusion restriction ^a	27	3 (11.1%)	14 (51.9%)	10 (37%)	<0.05

	Table 2 Magnetic resonance	e imaging finding	s and signal intensity	of pancreatic neuro	endocrine tumors b	based on tumor grade
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Notes: "Data are number of patients, with the percentage in parentheses. P-value was calculated with chi-squared or Fisher's exact test.



Figure 3 Magnetic resonance images of a 58-year-old woman with pancreatic neuroendocrine tumor Grade 1.

Notes: Fat-suppressed liver acquisition with volume acceleration sequence. TI-(A) and T2-(B) weighted imaging, a well-defined mass (arrows) located in the uncinate process of pancreas shows hypo-intensity and hyper-intensity, respectively. Diffusion-weighted images shows the tumor (arrow) was isointense (**C**). The tumor (arrows) shows persistent hyper-enhancement during the arterial (**D**), portal venous (**E**), and delayed (**F**) phase.



Figure 4 Magnetic resonance images of a 64-year-old woman with pancreatic neuroendocrine tumor Grade 3. Notes: Fat-suppressed liver acquisition with volume acceleration sequence. T1- (A) and T2- (B) weighted imaging, a well-defined mass (arrows) located in the body of pancreas shows hypo-intensity and slight hyper-intensity, respectively. Diffusion-weighted images shows the tumor (arrow) was hyper-intense (C). The tumor (arrows) shows persistent hypo-enhancement during the arterial (D), portal venous (E), and delayed (F) phase.

DWI texture features among different grades of PNETs were compared, and 22 T2WI and 37 DWI texture features showed significant differences among PNETs after Kruskal–Wallis test with Bonferroni correction (Tables S1 and S2). After assessment of VIF, there remained six (G1 vs G2), four (G1 vs G3), and four (G2 vs G3) T2WI texture features, and ten (G1 vs G2), six (G1 vs G3), and two (G2 vs G3) DWI texture features for multivariate logistic regression.

Multivariate logistic regression analyses for prediction of PNET grade based on T2WI and DWI texture features are shown in Tables 3 and 4, respectively. ROC curves and AUCs of predicting models in PNETs' grading are demonstrated in Figure S1. On T2WI, inverse difference moment (OR =1.768; 95% CI, 1.322–2.364; P<0.001] was proven to be an independent predictor of pathologic grade between G1 and G2 tumors. The AUC was 0.833 (95% CI: 0.725–0.941) and the cut-off value was –0.2 with 87.1% sensitivity and 75.9% specificity. Energy (OR =3.546; 95% CI, 0.554–22.675; P=0.052), correlation (OR =2.563; 95% CI, 1.352–4.859; P=0.004), and differenceEntropy (OR =0.013; 95% CI,

Table 3 Mu	Iltivariate logistic r	egression models for	prediction of	pancreatic neuroendocrin	e tumor grade with	T2WI texture features
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	Model	Features	OR (95% CI)	P-value	AUC (95% CI)	Cut-off value
						(SEN, SPE)
Grade I vs Grade 2	Model I	Inverse difference moment ^a	1.768 (1.322–2.364)	0.000	0.833 (0.725–0.941)	-0.2 (0.871, 0.759)
Grade I vs Grade 3	Model 2	Energy ^a	3.546 (0.554–22.675)	0.052	0.989 (0.970–1.000)	–1.358 (0.935, 1.0)
		Correlation ^a	2.563 (1.352–4.859)	0.004		
		DifferenceEntropy ^a	0.013 (0.001–0.184)	0.001		
Grade 2 vs Grade 3	Model 3	DifferenceEntropy ^a	17.166 (2.42–121.764)	0.004	0.813 (0.690–0.937)	-0.711 (0.69, 0.824)

Note: ^aData are adjusted ORs per one SD change.

Abbreviations: SEN, sensitivity; SPE, specificity; T2WI, T2-weighted imaging.

Table 4 Multivariate logistic regression models for	prediction of pancreatic neuroendocrir	ne tumor grade with DWI texture features
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	Model	Features	OR (95% CI)	P-value	AUC (95% CI)	Cut-off value
						(SEN, SPE)
Grade I vs Grade 2	Model I	Correlation ^a	2.473 (1.5-4.078)	0.003	0.841 (0.766–0.949)	-1.027 (0.645, 0.931)
		Contrast ^a	0.999 (0.996–1.002)	0.031		
		Inverse difference moment ^a	1.989 (1.29–3.066)	0.073		
Grade I vs Grade 3	Model 2	Maxintensity ^a	1.012 (1.002–1.021)	0.015	0.962 (0.908–1.0)	–0.792 (0.935, 0.941)
		Entropy ^a	4.342 (0.683–27.578)	0.120		
		Inverse difference moment ^a	2.678 (1.510–4.751)	0.020		
Grade 2 vs Grade 3	Model 3	Maxintensity ^a	0.997 (0.994–1.0)	0.023	0.703 (0.544–0.862)	-0.914 (0.517, 0.824)

Note: ^aData are adjusted ORs per one SD change.

Abbreviations: DWI, diffusion-weighted imaging; SEN, sensitivity; SPE, specificity.

0.001-0.184; *P*<0.001) were independent predictors of pathologic grade between G1 and G3 tumors. The AUC was 0.989 (95% CI: 0.970-1.0) and the cut-off value was -1.358 with 93.5% sensitivity and 100% specificity. DifferenceEntropy (OR =17.166; 95% CI, 2.42-121.764; *P*=0.004) was proven to be an independent predictor of pathologic grade between G2 and G3 tumors. The AUC was 0.813 (95% CI: 0.69-0.937) and the cut-off value was -0.711 with 69% sensitivity and 82.4% specificity.

On DWI, correlation (OR =2.473; 95% CI, 1.5-4.078; P=0.003), contrast (OR =0.999; 95% CI, 0.996-1.002; P=0.031), and inverse difference moment (OR =1.989; 95% CI, 1.29-3.066; P=0.073) were independent predictors of pathologic grade between G1 and G2 tumors. The AUC was 0.841 (95% CI: 0.766-0.949) and the cut-off value was -1.027 with 64.5% sensitivity and 93.1% specificity. Maxintensity (OR =1.012; 95% CI, 1.002-1.021; P=0.015), entropy (OR =4.324; 95% CI, 0.683-27.578; P=0.12), and inverse difference moment (OR = 2.678; 95%) CI, 1.51-4.751; P=0.02) were independent predictors of pathologic grade between G1 and G3 tumors. The AUC was 0.962 (95% CI: 0.908-1.0) and the cut-off value was -0.792 with 93.5% sensitivity and 94.1% specificity. Maxintensity (OR =0.997; 95% CI, 0.994-1.0; P=0.023) was proven to be an independent predictor of pathologic grade between G2

and G3 tumors. The AUC was 0.703 (95% CI: 0.544–0.862) and the cut-off value was –0.914 with 51.7% sensitivity and 82.4% specificity.

Discussion

Tumor grade has been described as a useful predictor of clinical outcome after surgical resection for PNETs,³¹ which is associated with treatment strategy.13 Accurate pretreatment grading of the tumor is vital in determining an efficient treatment strategy for PNETs. Invasive methods, such as fine-needle aspiration, have a limited accuracy in accurately diagnosing G3 tumors, because a small tissue sample may not represent the tumor hotspot (regions which had the largest mitotic count or Ki-67 index).³² Radiologic imaging may have great potential not only in PNET grading, but also in estimating the tumor aggressiveness and deciding the treatment plan before surgery. In the present study, we showed that MRI findings, such as tumor margin or tumor enhancement, can aid in predicting the histopathologic grade of PNETs. In addition, our data indicate that four T2-based (inverse difference moment, energy, correlation, and differenceEntropy) and five DWI-based (correlation, contrast, inverse difference moment, maxintensity, and entropy) TA parameters can also assist in PNET grading. The AUCs of six predicting models on T2WI and DWI ranged from 0.703-0.989.

Previous studies have identified several MRI features that can predict the histopathologic grades of PNETs. Besides well-established criteria such as local invasion or metastases and infiltration of vessels, other features associated with high tumor grade are ill-defined margin, larger tumor size, and hypo-enhancement pattern at the arterial phase. De Robertis et al¹³ reported that an ill-defined margin and significantly lower mean ADC value were valuable predictors for G2/G3 tumors. Canellas et al³³ found that G2/G3 tumors had larger tumor size (>2.0 cm), iso- or hypo- intensity on T2WI, presence of pancreatic ductal dilatation, and restricted diffusion within the tumor compared to G1 tumors. Similar results were observed in our study. Jang et al³⁴ showed that hyperenhancement at the arterial phase was more common in G1 compared with G2/G3 tumors. In our study, 87.1% of G1 tumors demonstrated a hyper-enhancement pattern compared with G2 (41.4%) or G3 tumors (0%), which is consistent with a previous study.34

TA provides an objective, quantitative assessment of tumor heterogeneity by analyzing the distribution and relationship of pixel or voxel gray levels in the image.³⁵ Tumors with high intratumoral heterogeneity tend to show a poorer prognosis or higher histopathologic grade.³⁶ Entropy is a measure of randomness of intensity images, which shows the degree of non-uniformity or complexity of the texture in the image. Canellas et al²³ reported that the odds of a tumor with high entropy values being an intermediate- or high-grade PNET were 3.7 times as high as those in tumors with low entropy values in CT TA. Energy is a measure of the intensity level distribution, which reflects the evenness of the gray level distribution and the grainy thickness of the image. If the value is high, then the distribution will present a small number of intensity levels. Kurtosis is a measure of whether the data are peaked or flat relative to a normal distribution. A high kurtosis means a distinct peak near the mean, a rapid decline, and heavy tails.³⁷ Pereira et al³⁸ demonstrated that G3 tumors had higher skewness and higher kurtosis compared with G1 tumors in histogram analysis of ADC. In our study, G2/G3 tumors showed higher entropy, higher kurtosis, lower energy, and higher maxintensity, which, in general, represent increased heterogeneity, which was consistent with previous reports.^{23,38} In addition, Choi et al¹¹ also investigated the value of CT TA in PNET grading, which also showed the potential of TA in differentiation of tumor grades. Liu et al³⁶ investigated and compared CT texture parameters between intermediate and high-grade lung adenocarcinoma, and they found high-grade lung adenocarcinoma has lower correlation and higher contrast. However, in our study, G2/G3 tumors

showed higher correlation and lower contrast compared with G1 tumors. Therefore, we speculate contrast and correlation may be predictors of histopathologic grade of tumors, but the relation between the two parameters and PNETs may require more validation studies.

To the best of our knowledge, other modalities, such as CT, single photon emission computed tomographysomatostatin receptor scintigraphy (SRS) (Octreoscan[®]), and 68Ga-DOTATOC positron emission tomography (PET)/CT, are also valuable in PNET grading.³⁹ Dromain et al⁴⁰ reported that Octreoscan is useful for PNET staging, detection of disease recurrence, or selecting patient candidates for peptide receptor radiometabolic treatment. In addition, Cloyd et al⁴¹ demonstrated that higher grade PNETs can be suspected preoperatively based on the presence of PET avidity and lack of Octreoscan uptake. Therefore, further studies addressing multimodal imaging or combination of MRI and PET/CT or SRS should be conducted.

Limitations

Our study has several limitations. First, there may have been selection bias due to the retrospective nature of the study. Second, only T2WI and DWI sequences were analyzed. Further studies about other sequences, such as contrast-enhanced images or ADC maps are needed. Third, we could not obtain inter-observer variability of the qualitative image analysis due to the consensus review by radiologists, despite the minor discrepancy during imaging analysis between two radiologists. Fourth, 3-T scanner from only one manufacturer was used in our study and it is unclear whether the results can be generalized to all manufacturers. Finally, although we built predicting models for PNET grading, we could not perform a validation test because we could not collect sufficient data on PNETs in a short term.

Conclusion

The current study indicated that MRI features, including tumor margin, texture, local invasion or metastases, tumor enhancement, and diffusion restriction, are useful for the prediction of histopathologic grade of PNETs. Notably, we also demonstrated that MRI texture parameters, such as inverse difference moment, differenceEntropy, maxintensity, entropy, and energy can aid in the prediction of PNET grading.

Ethics statement

This study was approved by the Ethical Committee of the Affiliated Hospital of Nanjing University of Chinese Medicine and patient informed consent requirement was waived due to its retrospective nature. Patient data confidentiality was protected in accordance with the Declaration of Helsinki principles.

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Disclosure

The authors report no conflicts of interest in this work.

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Supplementary materials



Figure S1 ROC curves of the multivariate logistic regression models for prediction of PNET grade with T2WI (**A**, Grade 1 vs Grade 2; **B**, Grade 1 vs Grade 3; **C**, Grade 2 vs Grade 3) and DWI (**D**, Grade 1 vs Grade 2; **E**, Grade 1 vs Grade 3; **F**, Grade 2 vs Grade 3) texture features. The AUCs are 0.833, 0.989, 0.813, 0.841, 0.962, and 0.703, respectively.

Abbreviations: PNET, pancreatic neuroendocrine tumors; T2WI, T2-weighted imaging; DWI, diffusion-weighted imaging.

Table SI Comparisons of T2WI texture features among pancreatic neuroendocrine tumors (*P*-values, Kruskal–Wallis test with Bonferroni correction)

Texture features on T2WI		Median (P-value)				
		Grade I vs Grade 2	Grade I vs Grade 3	Grade 2 vs Grade 3		
FI	Volume count	367 vs 2202 (<0.001)	367 vs 3770 (<0.001)	2202 vs 3770 (0.192)		
F2	Voxel value sum (×10 ⁵)	1.89 vs 14.98 (<0.001)	1.89 vs 20.72 (<0.001)	14.98 vs 20.72 (0.148)		
F3	Relative deviation	-185.5 vs -1081.4 (0.131)	-185.5 vs -2013.8 (<0.001)	-1081.4 vs -2013.8 (0.012)		
F4	Kurtosis	2.8 vs 3.3 (0.002)	2.8 vs 4.0 (<0.001)	3.3 vs 4.0 (0.09)		
F5	Frequency size	366 vs 2200 (<0.001)	366 vs 3769 (<0.001)	2200 vs 3769 (0.192)		
F6	Glcm total frequency	1540 vs 8350 (<0.001)	1540 vs 14494 (<0.001)	8350 vs 14494 (0.216)		
F7	Energy (×10 ³)	1.66 vs 0.47 (<0.001)	1.66 vs 0.57 (0.003)	0.47 vs 0.57 (0.691)		
F8	Inertia	734.5 vs 241.2 (<0.001)	734.5 vs 149.1 (<0.001)	241.2 vs 149.1 (0.044)		
F9	Correlation (×10 ³)	0.38 vs 0.65 (0.001)	0.38 vs 1.17 (<0.001)	0.65 vs 1.17 (0.021)		
FIO	Inverse difference moment (×10 ²)	5.85 vs 10.29 (<0.001)	5.85 vs 15.32 (<0.001)	10.29 vs 15.32 (0.034)		
FII	Cluster prominence (×10 ⁸)	2.00 vs 0.87 (0.001)	2.00 vs 0.51 (<0.001)	0.87 vs 0.51 (0.072)		
FI3	Difference variance (×10³)	2.26 vs 1.47 (0.221)	2.26 vs 0.92 (<0.001)	1.47 vs 0.92 (0.006)		
FI4	DifferenceEntropy	0.53 vs 0.48 (0.063)	0.53 vs 0.43 (<0.001)	0.48 vs 0.43 (0.001)		
F15	Inverse difference moment (×10 ²)	5.58 vs 7.37 (0.037)	5.58 vs 10.24 (<0.001)	7.37 vs 10.24 (0.001)		
FI6	Short run emphasis (×10³)	99.79 vs 99.96 (<0.001)	0.99.79 vs 0.99.98 (<0.001)	99.96 vs 99.98 (0.116)		
FI7	Long run emphasis	1.01 vs 1.00 (0.001)	1.01 vs 1.00 (<0.001)	1.00 vs 1.00 (0.121)		
F18	Gray-level non-uniformity	3.43 vs 14.19 (<0.001)	3.43 vs 43.51 (<0.001)	14.19 vs 43.51 (0.118)		
F19	Run length non-uniformity	361.26 vs 2125 (<0.001)	361.26 vs 3604 (<0.001)	2125 vs 3604 (0.192)		
F20	Low gray-level run emphasis (×10³)	5.95 vs 1.26 (<0.001)	5.95 vs 0.75 (<0.001)	1.26 vs 0.75 (0.165)		
F21	Short run low gray-level emphasis (×10³)	3.87 vs 0.85 (<0.001)	3.87 vs 0.54 (<0.001)	0.85 vs 0.54 (0.209)		
F22	Long run low gray-level emphasis (×10²)	1.43 vs 0.29 (<0.001)	1.43 vs 0.16 (<0.001)	0.293 vs 0.16 (0.160)		

Abbreviations: Glcm, Gray-Level Co-occurrence Matrix; T2WI, T2-weighted imaging.

Texture features on DWI		Median (P-value)				
		Grade I vs Grade 2	Grade I vs Grade 3	Grade 2 vs Grade 3		
FI	Maxintensity	270 vs 411 (0.003)	270 vs 628 (<0.001)	411 vs 628 (0.034)		
F2	Median Intensity	213 vs 264 (0.029)	213 vs 370 (<0.001)	264 vs 370 (0.021)		
F3	Mean value	201 vs 261 (0.020)	201 vs 373 (<0.001)	261 vs 373 (0.019)		
F4	Standard deviation	35.5 vs 56.4 (0.018)	35.5 vs 67.2 (<0.001)	56.4 vs 67.2 (0.049)		
F5	Variance	1260 vs 3185 (0.018)	1260 vs 2553 (<0.001)	3185 vs 2553 (0.049)		
F6	Volume count	89 vs 382 (<0.001)	89 vs 1017 (<0.001)	382 vs 1017 (0.271)		
F7	Voxel value sum	19661 vs 153737 (<0.001)	19661 vs 444309 (<0.001)	153737 vs 444309 (0.155)		
F8	RMS	205 vs 266 (0.019)	205 vs 379 (<0.001)	266 vs 379 (0.019)		
F9	Range	163 vs 291 (0.002)	163 vs 422 (<0.001)	291 vs 422 (0.062)		
F10	Mean deviation	54.3 vs -5.6 (0.02)	54.3 vs -117.7 (<0.001)	-5.6 vs -117.7 (0.019)		
FII	Energy (×10)	0.19 vs 0.10 (<0.001)	0.19 vs 0.09 (<0.001)	0.10 vs 0.09 (0.119)		
FI2	Entropy	5.90 vs 6.88 (<0.001)	5.90 vs 7.06 (<0.001)	6.88 vs 7.06 (0.129)		
FI3	Frequency size	88 vs 381 (<0.001)	88 vs 1016 (<0.001)	381 vs 1016 (0.271)		
FI4	Quantile5	138 vs 168 (0.087)	138 vs 260 (<0.001)	168 vs 260 (0.007)		
F15	Quantile I 0	150 vs 182 (0.045)	150 vs 290 (<0.001)	182 vs 290 (0.011)		
FI6	Quantile25	175 vs 213 (0.036)	175 vs 324 (<0.001)	213 vs 324 (0.017)		
FI7	Quantile50	212 vs 264 (0.027)	212 vs 370 (<0.001)	264 vs 370 (0.021)		
F18	Quantile75	222 vs 295 (0.021)	222 vs 422 (<0.001)	295 vs 422 (0.019)		
F19	Quantile90	246 vs 335 (0.015)	246 vs 465 (<0.001)	335 vs 465 (0.02)		
F20	Quantile95	255 vs 360 (0.011)	255 vs 481 (<0.001)	360 vs 481 (0.025)		
F21	Glcm total frequency	272 vs 1748 (<0.001)	272 vs 3786 (<0.001)	1748 vs 3786 (0.297)		
F22	Correlation (×10 ³)	0.17 vs 0.38 (<0.001)	0.17 vs 0.42 (<0.001)	0.38 vs 0.42 (0.41)		
F23	Inverse difference moment	0.03 vs 0.05 (<0.001)	0.03 vs 0.06 (<0.001)	0.05 vs 0.06 (0.305)		
F24	Haralick correlation (×108)	3.13 vs 6.37 (0.001)	3.13 vs 7.43 (<0.001)	6.37 vs 7.43 (0.343)		
F25	Haralick entropy	0.49 vs 0.62 (<0.001)	0.49 vs 0.67 (<0.001)	0.62 vs 0.67 (0.320)		
F26	Angular second moment (×10³)	4.50 vs 1.14 (<0.001)	4.50 vs 0.71 (<0.001)	1.14 vs 0.71 (0.325)		
F27	Contrast (×10³)	5.48 vs 2.94 (0.005)	5.48 vs 2.25 (<0.001)	2.94 vs 2.25 (0.101)		
F28	Sum entropy	0.71 vs 0.81 (<0.001)	0.71 vs 0.82 (<0.001)	0.81 vs 0.82 (0.640)		
F29	Difference variance (×103)	8.36 vs 4.50 (0.004)	8.36 vs 3.61 (<0.001)	4.50 vs 3.61 (0.110)		
F30	Inverse difference moment (×10²)	2.12 vs 4.09 (<0.001)	2.12 vs 4.15 (<0.001)	4.09 vs 4.15 (0.149)		
F31	Short run emphasis	0.99 vs 1.00 (<0.001)	0.99 vs 1.00 (<0.001)	1.00 vs 1.00 (0.233)		
F32	Long run emphasis	1.04 vs 1.01 (<0.001)	1.04 vs 1.00 (<0.001)	1.01 vs 1.00 (0.231)		
F33	Gray-level non-uniformity	1.71vs 4.82 (<0.001)	1.71 vs 7.58 (<0.001)	4.82 vs 7.58 (0.313)		
F34	Run length non-uniformity	85.52 vs 372.26 (<0.001)	85.52 vs 996.25 (<0.001)	372.26 vs 996.25 (0.272)		
F35	Low gray-level run emphasis (×10²)	2.54 vs 0.62 (<0.001)	2.54 vs 0.24 (<0.001)	0.62 vs 0.24 (0.259)		
F36	Short run low gray-level emphasis (×10³)	1.64 vs 0.39 (<0.001)	1.64 vs 0.18 (<0.001)	0.39 vs 0.18 (0.327)		
F37	Long run low gray-level emphasis (×10³)	6.54 vs 1.50 (<0.001)	6.54 vs 0.52 (<0.001)	1.50 vs 0.52 (0.232)		

 Table S2 Comparisons of DWI texture features among pancreatic neuroendocrine tumors (P-values, Kruskal–Wallis test with Bonferroni correction)

Abbreviations: DWI, diffusion-weighted imaging; Glcm, Gray-Level Co-occurrence Matrix; RMS, root mean square.

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