



Added-value of 3D amide proton transfer MRI in assessing prognostic factors of cervical cancer: a comparative study with multiple model diffusion-weighted imaging

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Background: Amide proton transfer (APT) imaging has been gradually applied to cervical cancer, yet the relationships between APT and multiple model diffusion-weighted imaging (DWI) have yet to be investigated. This study attempted to evaluate the added value of 3-dimensional (3D) APT imaging to multiple model DWI for assessing prognostic factors of cervical cancer.

Methods: This prospective diagnostic study was conducted in The First Affiliated Hospital of Zhengzhou University. A total of 88 consecutive patients with cervical cancer underwent APT imaging and DWI with 11 b-values (0–2,000 s/mm²). The apparent diffusion coefficient (ADC), pure molecular diffusion (D), perfusion fraction (f), pseudo-diffusion (D*), mean kurtosis (MK), and mean diffusivity (MD) were calculated based on mono-exponential, bi-exponential, and kurtosis models. The mean, minimum, and maximum values of APT signal intensity (APT SI) and DWI-derived metrics were compared based on tumor stages, subtypes, grades, and lymphovascular space invasion status by Student's *t*-test or Mann-Whitney *U* test. Receiver operating characteristic (ROC) curve analysis was used to evaluate the diagnostic performance of the parameters.

Results: APT SI_{max}, APT SI_{min}, MK_{mean}, and MK_{max} showed significant differences between adenocarcinoma (AC) and squamous cell carcinoma (SCC) (all *P*<0.05). APT SI_{mean}, APT SI_{max}, and MK_{max} were higher and ADC_{min}, D_{mean}, D_{min}, and MD_{min} were lower in the high-grade tumor than in low-grade tumor (all *P*<0.05). For distinguishing lymphovascular space invasion, only MK_{mean} showed significant difference (*P*=0.010). APT SI_{max} [odds ratio (OR) =2.347, *P*=0.029], APT SI_{min} (OR =0.352; *P*=0.024), and MK_{mean} (OR =6.523; *P*=0.001) were the independent predictors for tumor subtype, and APT SI_{max} (OR =2.885; *P*=0.044), MD_{min} (OR =0.155, *P*=0.012) were the independent predictors for histological grade of cervical cancer. When APT SI_{min} and APT SI_{max} was combined with MK_{mean} and MK_{max}, the diagnostic performance was significantly improved for differentiating AC and AC [area under the curve (AUC): 0.908, sensitivity: 87.5%; specificity: 83.3%; *P*<0.001]. The combination of APT SI_{mean}, APT SI_{max}, ADC_{min}, MK_{max}, and MD_{min} demonstrated the highest diagnostic performance for predicting tumor grade (AUC: 0.903, sensitivity: 78.6%; specificity: 88.9%; *P*<0.001).

Conclusions: Addition of APT to DWI may improve the ability to noninvasively predict poor prognostic factors of cervical cancer.

Keywords: Cervical cancer; magnetic resonance imaging (MRI); amide proton transfer imaging (APT imaging); diffusion-weighted imaging (DWI)

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Introduction

Despite improvements in prevention, screening, and treatment, cervical cancer continues to significantly impact women's health worldwide (1). The prognosis of cervical cancer depends on tumor size, clinical stage, and some histopathological features, including tumor subtype and grade, and lymph node status (2). The presence of lymphovascular space invasion (LVSI) in early-stage tumors has a higher rate of lymph node metastases and increases the risk of recurrence from 2% to 31% at 3 years (3,4). Both the updated International Federation of Gynecology and Obstetrics (FIGO) and European Society of Urogenital Radiology (ESUR) system give added importance to magnetic resonance imaging (MRI) as a method of accurately measuring tumor size and depicting the presence of deep stromal infiltration and parametrial involvement (5,6). Although conventional MRI provides morphological evaluation for cervical cancer, it is challenging to reflect tumor microstructure without specific metrics. Different advanced MRI modalities, including molecular imaging, may provide additional complementary information for clinical diagnosis and prognostic assessment, which may help optimize the therapeutic benefit (7-9).

Diffusion-weighted imaging (DWI) plays an essential role in cervical cancer assessment by providing both excellent visual definition of tumors due to high signal intensity (SI) and quantitative information that is obtained from various mathematical models. Mono-exponential DWI-derived apparent diffusion coefficient (ADC) has been most commonly used and inversely correlated with tumor cellularity. However, the water diffusion within tissues can be complicated, for example, the capillary blood microcirculation can contribute to the water diffusion, and affect the ADC value (10). Thus, the extended intravoxel incoherent motion (IVIM) model and kurtosis model, named diffusion kurtosis imaging (DKI), have been used to reflect the biological characterization of cervical cancer (11,12). Different from DWI, amide proton transfer

(APT) imaging has recently emerged as a novel contrast agent-free MRI technique in the field of cellular and molecular imaging, which generates image contrast based on endogenous cytosolic proteins and peptides with amide protons in the peptide bond (13). This technique has been successfully applied to brain tumor, prostate cancer, endometrial carcinoma, breast cancer, and bladder cancer (14-18). Some studies have reported the potential of APT SI in differentiating the histological subtype and grade of cervical cancer (9,19,20). However, in most studies, the APT imaging has been limited to a 2-dimensional (2D) acquisition, which could not provide coverage of the whole tumor (19,20). Comparatively 3-dimensional (3D) turbo spin-echo (TSE) APT imaging, with improved B_0 field correction and image reconstruction, enables more and comprehensive analysis of the whole tumor volume (21).

Although several studies have applied APT imaging or multiple diffusion models in the evaluation of cervical cancer, they were all primarily focused on the application of mean values (9,11,19,20). Due to tumor heterogeneity, the maximum or minimum values of diffusion and APT metrics within the whole lesion may also reflect the underlying aggressiveness of the tumor. Ghosh *et al.* found that ADC_{min} was more effective than ADC_{mean} in differentiating neuroblastoma with and without MYCN amplification (22). Zhuo *et al.* reported that higher APT SI_{max} were observed in H3K27M-mutant glioma than those with wildtype, whereas no significant differences of APT SI_{mean} were observed between groups (23). Therefore, the purpose of our study was to (I) compare the diagnostic performances of mean, minimum, and maximum values of APT SI for preoperative identifying the prognostic factors (tumor stage, subtype, grade, and LVSI status) of cervical cancer with those of multi-model DWI, and (II) examine the additive value of 3D-APT imaging combined with multi-model DWI for the preoperative characterization of cervical cancer. We present this article in accordance with the STARD reporting checklist (available at <https://qims.amegroups.com/article/view/10.21037/qims-23-324/rc>).

Methods

Participants

The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013). This prospective single-center study was approved by the institutional ethics board of The First Affiliated Hospital of Zhengzhou University, and all patients provided written informed consent prior to enrollment in the study. From January 2020 to April 2022, 106 patients in The First Affiliated Hospital of Zhengzhou University were selected consecutively. The inclusion criteria were as follows: (I) pathological diagnosis of cervical cancer; (II) no treatment before MR examination; (III) the availability of data from preoperative MRI, including DWI and APT imaging. The exclusion criteria were as follows: (I) rare tumors such as adenosquamous carcinoma and neuroendocrine carcinoma (n=7); (II) poor image quality (n=5); (III) tumor visibility on only one MRI slice (n=6). Finally, a total of 88 patients were enrolled in the study. Among the patients, 53 underwent surgery and 35 underwent biopsy for final diagnoses. According to the 2018 FIGO staging system (5) and histopathology results, the patients were divided into an early-stage group (< IIB) and an advanced-stage group (\geq IIB), an adenocarcinoma (AC) group and a squamous cell carcinoma (SCC) group, a high-grade group (poorly differentiated tumor) and a low-grade group (well/moderately differentiated tumor), and an LVSI-negative group and an LVSI-positive group.

Image acquisition

Cervical MR scanning was performed using a 3T system (Ingenia CX, Philips Healthcare, Best, The Netherlands) with an external pelvic phased array coil. Patients were asked to fast for at least 6 hours prior to the examination and were placed in the head-first supine position throughout the examination. The following sequences were acquired: axial T2-weighted TSE imaging, DWI, and APT imaging. APT imaging was conducted using an optimized 3D TSE pulse sequence combined with chemical shift-selective fat suppression for better robustness to field inhomogeneity and 3D volume coverage. The pre-saturation pulses consisted of three consecutive radiofrequency pulses of 500 ms duration with 500 ms interpulse delays and a time-average amplitude of 2 μ T. We acquired 6 different image volumes at saturation frequencies (± 3.5 , ± 3.42 , ± 3.58 , and $-1,560$, relative to the water resonance frequency) were acquired

for the Z-spectrum normalization and interpolation, and three acquisitions were performed at +3.5 ppm with different echo time shifts to obtain a Dixon-type B0 field map to correct for B0 inhomogeneities in the Z-spectrum frequency domain. The total scan time for the APT imaging (additional time for setup, 7–10 seconds; acquisition time for APT images, 6 minutes 7 seconds to 6 minutes 10 seconds. DWI was acquired via a single-shot echo-planar imaging (EPI) sequence for b = 0, 10, 20, 50, 100, 200, 400, 800, 1,200, 1,600, and 2,000 s/mm². Diffusion gradients were applied simultaneously along with 3 orthogonal directions. The scan time for DWI was 5 minutes 43 seconds to 5 minutes 48 seconds. The total scan time of this study protocol was 16 minutes 25 seconds to 16 minutes 37 seconds. Detailed descriptions of MRI parameters are listed in *Table 1*.

Image analysis

APT images were acquired online at the MR control console. All data were transferred to a post-processing station (IntelliSpace Portal V10, Philips Healthcare) for quantitative analysis. The APT SI was calculated as the asymmetry of the magnetization transfer rate (MTR_{asym}) at the frequency offset of +3.5 ppm using the following equation:

$$\text{APT SI} = MTR_{\text{asym}} [\Delta\omega = +3.5 \text{ ppm}] (\%) \quad [1]$$

DWI data were loaded into the application of advanced diffusion analysis (ADA) to generate parameter maps of the three diffusion models. The mono-exponential model was applied to estimate the ADC with 2 b-values (0 and 800 s/mm²). The following linear fitting equation was implemented:

$$S_b/S_0 = \exp(-b \cdot D) \quad [2]$$

where S_0 and S_b represent the MR SI under different b values (0 s/mm² and other values, respectively).

The calculation of the IVIM model used 9 b-values (0, 10, 20, 50, 100, 200, 400, 800, and 1,200 s/mm²). The following fitting equation was implemented:

$$S_b/S_0 = (1-f) \cdot \exp(-b \cdot D) + f \cdot \exp(-b \cdot (D^* + D)) \quad [3]$$

where f is the perfusion fraction, D is the diffusion coefficient representing water diffusion in the extravascular space, and D^* is the pseudo diffusion coefficient representing incoherent microcirculation within the vascular space.

Table 1 Imaging protocol parameters

Parameters	T2WI	DWI	APT imaging
Sequence/orientation	TSE/axial	EPI/axial	TSE/axial
TR (ms)	4,093	5,614	3,000
TE (ms)	100	103	1,000
Field of view (mm ²)	200×200	240×240	180×180
Slice thickness (mm)	5	5	5
No. of slices	24	24	9
Matrix	344×306	72×67	116×90
Voxel size (mm ³)	0.58×0.65×5.00	3.33×3.58×5.0	2×2×5.0
b-values (s/mm ²)	NA	0, 10, 20, 50, 100, 200, 400, 800, 1,200, 1,600, 2,000	NA
Flip angle (degrees)	90	90	90
Bandwidth (Hz/pixel)	250.6	1,784.8	647.2
Fat suppression	No	SPAIR	SPAIR
Acquisition time	4 min 30 s	5 min 37 s	6 min 0 s

T2WI, T2-weighted imaging; DWI, diffusion-weighted imaging; APT, amide proton transfer; TSE, turbo spin-echo; EPI, echo-planar imaging; TR, repetition time; TE, echo time; NA, not applicable; SPAIR, spectral attenuation with inversion recovery.

The calculation of the DKI model used 8 b-values (0, 100, 200, 400, 800, 1,200, 1,600 and 2,000 s/mm²). The following fitting equation was implemented:

$$S_b/S_0 = \exp(-b \cdot D + 1/6 \cdot b^2 \cdot D \cdot K) \quad [4]$$

where D is the corrected ADC derived from the non-Gaussian model, and K is a unitless metric of the apparent kurtosis coefficient. MD and MK are the averages of D and K among three distributed directions.

The image data were independently analyzed by two radiologists (Li S and Liu J, with over 7 years of experience in uterine MR diagnosis) who were unaware of the patient's clinical and pathologic information. The regions of interest (ROIs) of tumor area were defined on every section of DWI original map ($b = 1,200 \text{ s/mm}^2$) where mass was visible with reference to corresponding T2-weighted images. Areas of necrosis and hemorrhage were avoided. Subsequently, ROIs were copied to APT, ADC, D, D*, f, MK, and MD images, using a built-in automatic coregistration tool. Registrations were visually inspected and manually corrected if necessary. The average values of each measurement were recorded as APT SI_{mean} , ADC_{mean}, D_{mean}, D*_{mean}, f_{mean}, MK_{mean}, and MD_{mean} of the tumor, respectively. The lowest APT SI, ADC, D, and MD values of the whole tumor in all selected section were selected as the APT SI_{min} , ADC_{min}, D_{min}, and

MD_{min}. Similarly, the largest APT SI and MK value of the whole tumor was selected as the APT SI_{max} and MK_{max}.

Statistical analysis

The values of MR quantitative metrics were expressed as means ± standard deviations (SDs). The measured data were evaluated for normality using Shapiro-Wilk test. The unpaired Student's *t*-test was used to compare the continuous variables with normal distribution, whereas Mann-Whitney *U* test was used to compare variables with nonnormal distribution. Interclass correlation coefficient (ICC) was calculated to evaluate interobserver agreement on the measured metrics. ICC values greater than 0.8 were considered excellent agreement, 0.6–0.8, good; 0.4–0.6, moderate; and less than 0.4, poor agreement. Receiver operating characteristic (ROC) curves were generated to examine the ability of each metric to distinguish different prognostic factors when appropriate. The discriminating power of each metric was quantified using the area under the curve (AUC). The differences of AUCs were compared using DeLong test. Logistic regression analyses were used to identify independent factors and combination diagnosis.

Statistical analyses were performed using SPSS 22.0 (IBM Corp., Armonk, NY, USA) and MedCalc V19.0 (MedCalc

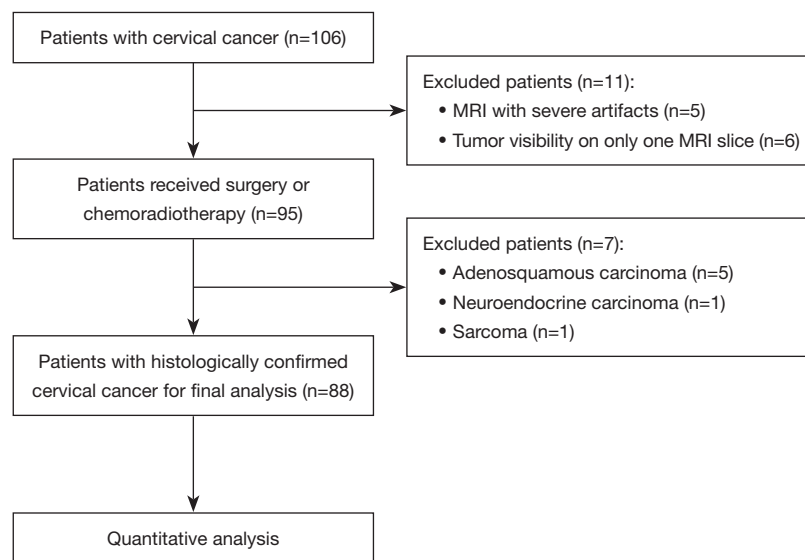


Figure 1 Flowchart of patient selection. MR, magnetic resonance; MRI, magnetic resonance imaging.

Software, Mariakerke, Belgium). The standard for statistical significance was set at $P < 0.05$.

Results

Study population

The final study population consisted of 88 patients with pathologically diagnosed cervical cancer (Figure 1). Table 2 gives an overview of the clinical characteristics of the patients. The interval between MR examination and treatment was 1 to 15 days, with a median of 6 days.

Interobserver reliability

The interobserver agreements for the MR metrics were good or excellent. The ICCs were as follows: $APT SI_{mean}$, 0.820; $APT SI_{max}$, 0.874; $APT SI_{min}$, 0.792; ADC_{mean} , 0.863; ADC_{min} , 0.845; D_{mean} , 0.857; D_{min} , 0.817; D^*_{mean} , 0.666; f_{mean} , 0.769; MK_{mean} , 0.824; MK_{max} , 0.815; MD_{mean} , 0.881; and MD_{min} , 0.856. Therefore, two observers' mean results of each metric were used for subsequent analysis.

Comparison of metrics in different stages, subtypes, grades, and LVSI status

No significant differences were detected in APT- and DWI-derived metrics for differentiating early-stage and advanced-stage cervical cancer (all $P > 0.05$). $APT SI_{max}$, $APT SI_{min}$,

MK_{mean} , and MK_{max} showed significant differences between SCC and AC groups (all $P < 0.05$). $APT SI_{max}$, MK_{mean} , and MK_{max} were higher and $APT SI_{min}$ were lower in AC than in SCC groups. There were no significantly different metrics from mono-exponential and IVIM diffusion models between SCC group and AC group (all $P > 0.05$). $APT SI_{mean}$, $APT SI_{max}$, ADC_{min} , D_{mean} , D_{min} , MD_{min} , and MK_{max} showed significant differences between the high-grade group and low-grade group (all $P < 0.05$). $APT SI_{mean}$, $APT SI_{max}$, and MK_{max} were higher and ADC_{min} , D_{mean} , D_{min} , and MD_{min} were lower in the high-grade group. For distinguishing LVSI, only MK_{mean} showed significant difference between LVSI-positive group and LVSI-negative group ($P < 0.05$). The detailed results are shown in Table 3. Images from conventional T2-weighted, APT imaging, and the 3 diffusion models of representative images are shown in Figures 2,3. More corresponding case examples of patients with cervical cancer are shown in Figures S1-S6.

Univariate and multivariate logistic regression analyses

Univariate logistic analysis revealed that $APT SI_{max}$, $APT SI_{min}$, MK_{mean} , and MK_{max} were predictors for the histological subtype of cervical cancer, and $APT SI_{mean}$, $APT SI_{max}$, ADC_{min} , D_{min} , MD_{min} , and MK_{max} were predictors for the histological grade of cervical cancer. For APT metrics, multivariate logistic analysis showed that $APT SI_{max}$ was the independent predictor for both subtype and grade

Table 2 Clinicopathologic characteristics of patients

Clinicopathologic characteristics	Values
Age (years), mean \pm SD	53.4 \pm 12.0
FIGO stage, n	
IB	24
IIA	29
IIB	8
IIIA	3
IIIB	1
IIIC	22
IVA	1
Histological type, n	
SCC	72
AC	16
Histological grade, n	
Low-grade (well/moderately differentiated)	36
High-grade (poorly differentiated)	28
Unknown [†]	24
LVSI, n	
Positive	27
Negative	26
Unknown [‡]	35

[†], cases are of unknown tumor differentiation as a result of insignificant biopsy specimen size, which was excluded when analyzing tumor grade; [‡], cases are of unknown LVSI status as a result of insignificant biopsy specimen size, which was excluded when analyzing tumor LVSI. SD, standard deviation; FIGO, International Federation of Gynecology and Obstetrics; SCC, squamous cell carcinoma; AC, adenocarcinoma; LVSI, lymphovascular space invasion.

of cervical cancer, and APT SI_{min} was the independent predictor for tumor subtype. For DWI metrics, only MK_{mean} was an independent predictor for the histological subtype, and MD_{min} was an independent predictor for the histological grade of cervical cancer (Table 4).

Diagnostic performance of APT and DWI metrics

Table 5 and Figure 4 summarize the results of ROC curve analysis for significantly different metrics. The AUC for MK_{mean} to predict LVSI was 0.689. The AUCs for MK_{mean},

APT SI_{min}, APT SI_{max}, and MK_{max} to differentiate SCC and AC were 0.760, 0.708, 0.691, and 0.661, respectively. For distinguishing between high-grade and low-grade tumor, MD_{min} achieved the highest AUC (0.810), followed by ADC_{min} (0.740), APT SI_{max} (0.731), D_{min} (0.727), MK_{max} (0.712), APT SI_{mean} (0.693), and D_{mean} (0.652).

Additive value of APT imaging to DWI for differentiating tumor subtypes and grades

For distinction of SCC and AC, 15 combinations of prediction models were designed using MK_{mean}, APT SI_{min}, APT SI_{max}, and MK_{max}. The combination of all four metrics revealed the highest AUC (0.908), which was significantly higher than that of any single metric (all P<0.05). For prediction of high-grade cervical cancer, 127 combinations of prediction models were designed using APT SI_{mean}, APT SI_{max}, ADC_{min}, D_{mean}, D_{min}, MD_{min}, and MK_{max}. The combination of APT SI_{mean}, APT SI_{max}, ADC_{min}, MK_{max}, and MD_{min} revealed the highest AUC (0.903), which was significantly higher than that of any single metric (all P<0.05).

Discussion

In this study, we explored the value of multi-model DWI and APT imaging in the prediction of clinical-pathological factors in patients with cervical cancer. Our results demonstrated that both APT imaging and multi-model DWI can be used to identify the histological subtype and grade of cervical cancer. Multiparametric analysis showed that adding APT to DWI can significantly improve the diagnostic performance with an AUC of 0.908 to predict tumor subtype, and an AUC of 0.903 to predict histological grade. In addition, higher MK_{mean} were observed in LVSI-positive tumors.

By focusing on amide protons, APT imaging has been used to visualize endogenous mobile proteins and peptides, which indirectly reflects changes of the internal environment (13). We observed a significantly higher APT SI_{max} and a significantly lower APT SI_{min} in AC than SCC. Cervical AC originates from endocervical cells that have abundant glandular structures and strong secretion function. This may result in increased concentrations of macromolecules in tumor tissue and consequently increased APT SI_{max}. The APT contrast is also affected by tissue pH. Decreased APT SI_{min} in AC may indicate the different local pH changes of the two microenvironments. However, APT

Table 3 APT- and DWI-derived metrics in differentiating tumor stages, subtypes, grades, and LVSI status of cervical cancer

Parameters	Tumor stage			Tumor subtype			Tumor grade			LVSI status		
	Early (n=53)	Advanced (n=35)	P value	SCC (n=72)	AC (n=16)	P value	High-grade (n=28)	Low-grade (n=36)	P value	Positive (n=27)	Negative (n=26)	P value
APT SI _{mean} (%)	2.82±0.65	2.66±0.87	0.095 [†]	2.74±0.79	2.84±0.53	0.312 [†]	3.15±0.95	2.67±0.53	0.008 ^{**†}	2.87±0.55	2.74±0.75	0.483 [‡]
APT SI _{max} (%)	5.28±1.41	5.68±1.77	0.308 [†]	5.24±1.45	6.38±1.78	0.017 ^{**†}	6.27±1.85	4.92±1.23	0.002 ^{**†}	5.59±1.63	5.02±1.09	0.206 [†]
APT SI _{min} (%)	0.07±1.47	0.13±1.23	0.846 [‡]	0.30±1.23	-0.83±1.62	0.002 ^{**†}	0.12±1.59	0.37±1.11	0.470 [‡]	0.26±1.32	0.10±1.44	0.673 [‡]
ADC _{mean} (×10 ⁻³ mm ² /s)	0.84±0.19	0.84±0.14	0.695 [†]	0.84±0.18	0.84±0.15	0.948 [†]	0.80±0.15	0.88±0.19	0.167 [†]	0.81±0.10	0.88±0.24	0.119 [†]
ADC _{min} (×10 ⁻³ mm ² /s)	0.52±0.14	0.52±0.14	0.852 [‡]	0.53±0.15	0.48±0.09	0.218 [‡]	0.46±0.11	0.57±0.15	0.002 ^{**†}	0.53±0.11	0.52±0.19	0.745 [‡]
D _{mean} (×10 ⁻³ mm ² /s)	0.70±0.12	0.71±0.09	0.831 [†]	0.71±0.11	0.70±0.12	0.665 [†]	0.68±0.13	0.74±0.09	0.038 ^{**†}	0.70±0.10	0.71±0.13	0.294 [†]
D _{min} (×10 ⁻³ mm ² /s)	0.41±0.13	0.43±0.10	0.739 [†]	0.43±0.12	0.38±0.11	0.176 [‡]	0.37±0.11	0.46±0.12	0.004 ^{**†}	0.43±0.11	0.40±0.14	0.374 [‡]
D* _{mean} (×10 ⁻³ mm ² /s)	44.26±18.93	43.02±17.53	0.759 [†]	44.92±17.48	38.56±21.42	0.210 [†]	46.74±19.51	45.08±18.00	0.726 [‡]	47.94±19.57	44.12±18.16	0.464 [‡]
f _{mean}	0.16±0.04	0.18±0.11	0.711 [†]	0.16±0.05	0.20±0.14	0.086 [†]	0.17±0.11	0.17±0.04	0.261 [†]	0.15±0.03	0.17±0.05	0.117 [†]
MK _{mean}	0.77±0.21	0.82±0.18	0.218 [‡]	0.75±0.17	0.96±0.23	0.001 ^{**†}	0.80±0.21	0.73±0.17	0.282 [†]	0.82±0.17	0.69±0.21	0.010 ^{**†}
MK _{max}	1.48±0.35	1.50±0.32	0.597 [†]	1.45±0.30	1.67±0.43	0.044 ^{**†}	1.59±0.35	1.38±0.26	0.004 ^{**†}	1.45±0.31	1.46±0.33	0.769 [†]
MD _{mean} (×10 ⁻³ mm ² /s)	1.03±0.18	1.05±0.15	0.848 [†]	1.04±0.17	1.05±0.20	0.996 [†]	1.00±0.18	1.08±0.17	0.161 [†]	1.03±0.12	1.06±0.22	0.650 [†]
MD _{min} (×10 ⁻³ mm ² /s)	0.55±0.14	0.54±0.13	0.718 [†]	0.56±0.14	0.52±0.11	0.303 [‡]	0.47±0.10	0.61±0.14	<0.001 ^{**†}	0.55±0.13	0.55±0.16	0.888 [†]

Data are presented as mean ± standard deviation. **, indicates the comparison with statistical significance; †, comparisons were performed by Mann-Whitney U test; ‡, comparisons were performed by independent t-test. APT, amide proton transfer; DWI, diffusion-weighted imaging; LVSI, lymphovascular space invasion; SCC, squamous cell carcinoma; AC, adenocarcinoma; SI, signal intensity; ADC, apparent diffusion coefficient; D, pure molecular diffusion; D*, pseudo-diffusion; f, perfusion fraction; MK, mean kurtosis; MD, mean diffusivity.

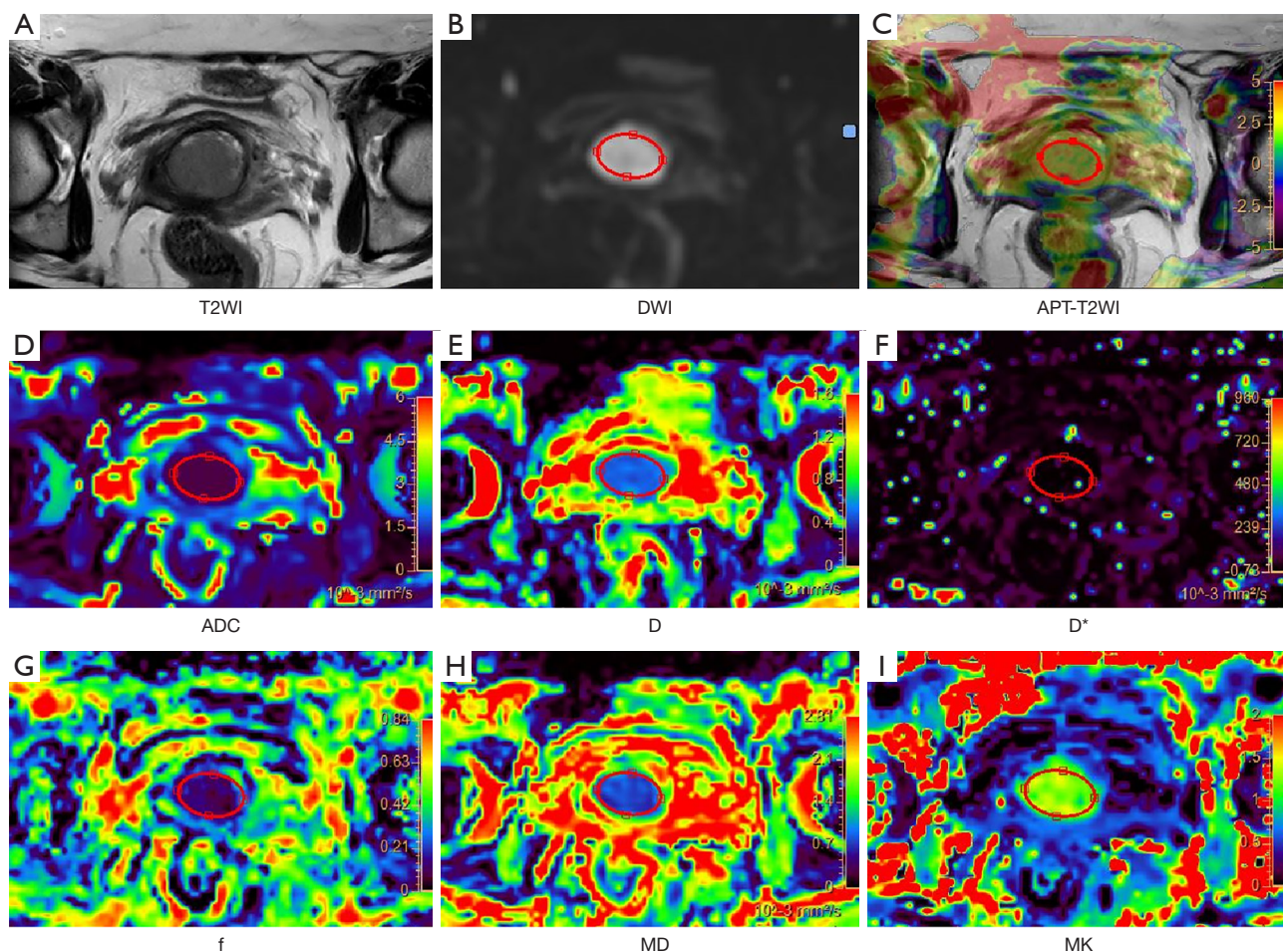


Figure 2 Images from a 45-year-old woman with histologically proven cervical squamous cell carcinoma (red circle, FIGO Ib2; G3; LVSI-negative), including T2WI (A), DWI with $b = 1,200 \text{ s/mm}^2$ (B), APT-T2 merged image (C), ADC map (D), D map (E), D^* map (F), f map (G), MD map (H), and MK map (I). T2WI, T2-weighted imaging; DWI, diffusion-weighted imaging; APT, amide proton transfer; ADC, apparent diffusion coefficient; D, pure molecular diffusion; D^* , pseudo-diffusion; f, perfusion fraction; MD, mean diffusivity; MK, mean kurtosis; FIGO, International Federation of Gynecology and Obstetrics; G3, grade 3; LVSI, lymphovascular space invasion.

SI_{mean} was unable to distinguish between SCC and AC in this study. It could be that the heterogeneity of both SCC and AC affected the endogenous mobile proteins and tissue pH in a similar extent as depicted by APT SI_{mean} . Meng *et al.* found that the APT SI of the SCC was significantly lower than that of AC by using a 2D EPI sequence (19). The inconsistent results may be due to the differences in the study population, sequence acquisition method, and ROI placement. To the best of our knowledge, this is the first report on the application of maximum and minimum values of APT SI in cervical cancer. Previous studies were performed at a single slice containing the maximum cross-sectional tumor area (9,19,20). In the present study, APT

imaging was acquired using a 3D TSE sequence, which allowed the measurement of maximum and minimum values of APT SI based on volumetric coverage. 3D-TSE-APT imaging could provide geometrically undistorted images with high quality, and reduced lipid artifact (24).

This study also found that the APT SI_{mean} and APT SI_{max} of the high-grade tumor were generally higher than those of the low-grade tumor, which was consistent with previous studies (9,16,18,19,25). Choi *et al.* confirmed that high-grade gliomas had higher concentrations of mobile proteins and peptides than low-grade gliomas (25). Li *et al.* stated that APT SI was positively correlated with the SCC grades (9). The pathological features of high-grade cervical cancer

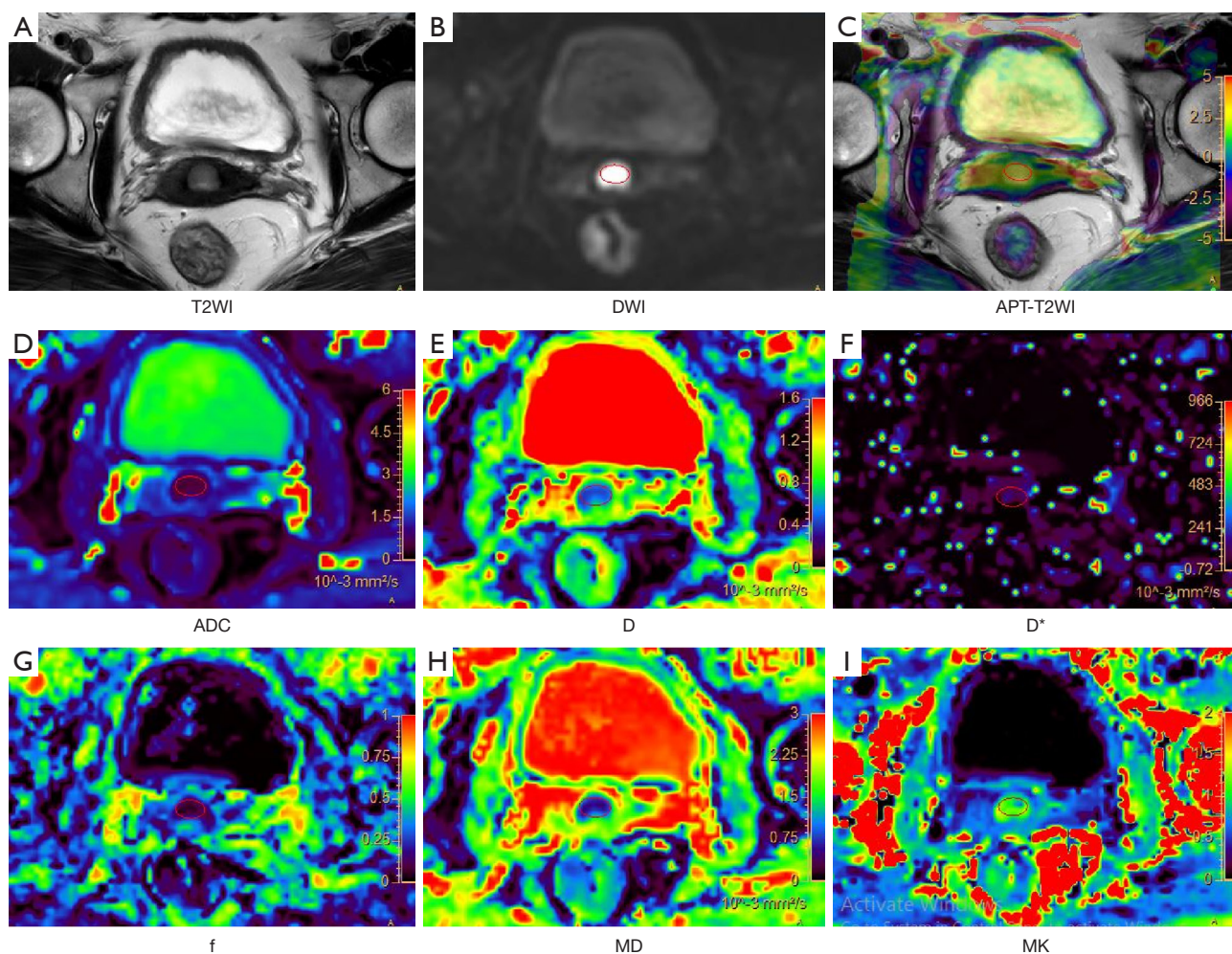


Figure 3 Images from a 54-year-old woman with histologically proven cervical adenocarcinoma (red circle, FIGO Ib1; G2; LVSI-positive), including T2WI (A), DWI with $b = 1,200 \text{ s/mm}^2$ (B), APT-T2 merged image (C), ADC map (D), D map (E), D^* map (F), f map (G), MD map (H), and MK map (I). T2WI, T2-weighted imaging; DWI, diffusion-weighted imaging; APT, amide proton transfer; ADC, apparent diffusion coefficient; D, pure molecular diffusion; D^* , pseudo-diffusion; f , perfusion fraction; MD, mean diffusivity; MK, mean kurtosis; FIGO, International Federation of Gynecology and Obstetrics; G2, grade 2; LVSI, lymphovascular space invasion.

include a higher cell density, greater nuclear atypia, and more microscopic necrosis than low-grade cervical cancer. The increased number of tumor cells which contain more proteins and peptides may account for the increase in the APT SI in high-grade tumor (9,18,25,26). Nuclear atypia has been reported to cause an interaction between the hydrophobic cell membrane and macromolecule substance and promote the release of proteins and peptides (16,19). Necrosis may be the third possible factor of increased APT SI in high-grade tumor. Togao *et al.* found that the diffuse gliomas with microscopic necrosis showed higher APT SI than those without necrosis (27). The highly concentrated

proteins and peptides in the extracellular space, such as microscopic necrosis or fluid collection in the microcystic space, may result in higher APT SI (27,28). Different tumor stages or different LVSI status, however, may not alter the overall macromolecular concentration or structure of the whole tumor, and the corresponding APT metrics showed no significant difference in our study.

Mono-exponential DWI, IVIM, and DKI have been widely reported to provide insight into typing and grading of cervical cancer (11,29,30). However, most studies only utilized one or two diffusion models. Our study processed three diffusion models through a single scan to predict

Table 4 Univariate and multivariate analyses for identifying histological subtypes and grades of cervical cancer

Parameters	SCC and AC (univariate analyses)		SCC and AC (multivariable analyses)		High- and low-grade (univariate analyses)		High- and low-grade (multivariable analyses)	
	OR (95% CI)	P value	OR (95% CI)	P value	OR (95% CI)	P value	OR (95% CI)	P value
APT S _{mean} (%)	1.129 (0.676–1.884)*	0.644	–	–	2.085 (1.062–4.095)*	0.033**	1.949 (0.664–5.720)*	0.225
APT S _{max} (%)	1.966 (1.156–3.344)*	0.013**	2.347 (1.090–5.053)*	0.029**	2.532 (1.329–4.822)*	0.005**	2.885 (1.028–8.096)*	0.044**
APT S _{min} (%)	0.432 (0.236–0.791)*	0.006**	0.352 (0.142–0.874)*	0.024**	0.796 (0.471–1.344)*	0.393	–	–
ADC _{mean} ($\times 10^{-3}$ mm ² /s)	0.957 (0.548–1.673)*	0.878	–	–	0.572 (0.288–1.137)*	0.111	–	–
ADC _{min} ($\times 10^{-3}$ mm ² /s)	0.702 (0.400–1.232)*	0.217	–	–	0.424 (0.221–0.815)*	0.010**	2.002 (0.392–10.220)*	0.404
D _{mean} ($\times 10^{-3}$ mm ² /s)	0.937 (0.546–1.609)*	0.814	–	–	0.550 (0.293–1.033)*	0.063	–	–
D _{min} ($\times 10^{-3}$ mm ² /s)	0.693 (0.407–1.181)*	0.177	–	–	0.462 (0.254–0.840)*	0.011**	1.641 (0.403–6.678)*	0.489
D* _{mean} ($\times 10^{-3}$ mm ² /s)	0.686 (0.381–1.237)*	0.210	–	–	1.085 (0.663–1.774)*	0.746	–	–
f _{mean}	1.538 (0.910–2.598)*	0.108	–	–	1.049 (0.659–1.669)*	0.841	–	–
MK _{mean}	3.797 (1.833–7.862)*	<0.001**	6.523 (2.081–20.441)*	0.001**	1.495 (0.833–2.683)*	0.178	–	–
MK _{max}	1.815 (1.072–3.071)*	0.026**	0.921 (0.376–2.258)*	0.858	2.168 (1.144–4.109)*	0.018**	3.856 (0.991–15.008)*	0.052
MD _{mean} ($\times 10^{-3}$ mm ² /s)	1.062 (0.616–1.830)*	0.829	–	–	0.613 (0.344–1.093)*	0.097	–	–
MD _{min} ($\times 10^{-3}$ mm ² /s)	0.748 (0.432–1.296)*	0.301	–	–	0.269 (0.127–0.571)*	0.001**	0.155 (0.036–0.662)*	0.012**

All factors with $P < 0.05$ in univariate analyses were included in multivariate regression analyses. *, OR for per 1 standard deviation; **, indicates the logistic regression analyses with statistical significance. SCC, squamous cell carcinoma; AC, adenocarcinoma; OR, odds ratio; CI, confidence interval; APT SI, amide proton transfer signal intensity; ADC, apparent diffusion coefficient; D, pure molecular diffusion; D*, pseudo-diffusion; f, perfusion fraction; MK, mean kurtosis; MD, mean diffusivity.

Table 5 ROC analysis of APT- and DWI-derived metrics

Category	Threshold	AUC (95% CI)	Sensitivity	Specificity	P value
SCC vs. AC					
APT SI _{max} (%)	4.91	0.691 (0.584–0.786)	81.2%	51.4%	0.008
APT SI _{min} (%)	−0.65	0.708 (0.601–0.800)	62.5%	80.6%	0.005
MK _{mean}	0.91	0.760 (0.657–0.844)	68.7%	88.9%	<0.001
MK _{max}	1.54	0.661 (0.553–0.759)	56.2%	73.6%	0.042
Combined diagnosis [†]	NA	0.908 (0.827–0.959)	87.5%	83.3%	<0.001
Tumor grade: high vs. low					
APT SI _{mean} (%)	2.81	0.693 (0.566–0.803)	75.0%	66.7%	0.006
APT SI _{max} (%)	5.40	0.731 (0.606–0.834)	64.3%	75.0%	<0.001
ADC _{min} (×10 ^{−3} mm ² /s)	0.52	0.740 (0.615–0.841)	75.0%	69.4%	<0.001
D _{mean} (×10 ^{−3} mm ² /s)	0.70	0.652 (0.522–0.767)	67.9%	63.9%	0.031
D _{min} (×10 ^{−3} mm ² /s)	0.48	0.727 (0.601–0.831)	92.9%	47.2%	<0.001
MK _{max}	1.52	0.712 (0.585–0.818)	60.7%	86.1%	0.002
MD _{min} (×10 ^{−3} mm ² /s)	0.61	0.810 (0.693–0.897)	100.0%	52.8%	0.002
Combined diagnosis [‡]	NA	0.903 (0.803–0.963)	78.6%	88.9%	<0.001
LVSI (+) vs. (−)					
MK _{mean}	0.77	0.689 (0.547–0.809)	55.6%	76.9%	0.010

[†], the combined diagnosis represents APT SI_{max} + APT SI_{min} + MK_{mean} + MK_{max}; [‡], the combined diagnosis represents APT SI_{mean} + APT SI_{max} + ADC_{min} + MK_{max} + MD_{min}. ROC, receiver operating characteristic; APT, amide proton transfer; DWI, diffusion-weighted imaging; AUC, area under the curve; CI, confidence interval; SCC, squamous cell carcinoma; AC, adenocarcinoma; SI, signal intensity; MK, mean kurtosis; ADC, apparent diffusion coefficient; D, pure molecular diffusion; MD, mean diffusivity; LVSI, lymphovascular space invasion.

the adverse histological features. We found that MK metrics showed significant differences between patients with different subtypes, grades and LVSI status. Moreover, diffusion metrics (ADC_{min}, D_{mean}, D_{min}, and MD_{min}) showed significant differences between high-grade and low-grade tumor, and no DWI metrics showed significant differences between FIGO stages. These results are partially consistent with previous studies. Winfield *et al.* reported that perfusion metrics (D*, f), and MK were significantly different between cervical SCC and AC, and diffusion coefficients (ADC, D, and MD) were significantly different between tumor grades (11). Wang *et al.* found that MD and ADC exhibited significant differences between histological subtypes and FIGO stages of cervical cancer, but not with tumor grades (29). Another study demonstrated that ADC, D, D*, and f were significantly different among the 3 grades of cervical cancer (30). These mixed results could be related to a variety of reasons, such as the reproducibility of DWI metrics, the choice of b values, and different research

subjects. According to a previous study, the reproducibility of IVIM metrics, especially D* and f, was significantly lower than that of ADC calculated from the mono-exponential model (31). Different b values might also cause a bias in DWI metrics. Rosenkrantz *et al.* suggested acquiring more than 3 b values for body DKI analysis, including at least 2 b values both above and below 1,000 s/mm² to allow robust estimates of MK (32). In the study of Winfield *et al.*, the highest b value was 800 s/mm², which may not have been sufficiently high enough to capture the non-Gaussian behavior (11). Perucho *et al.* suggested setting at least 6 b values for reliable estimation of IVIM metrics, with at least 2 b values greater than 200 s/mm² to estimate D and at least 2 b values less than 200 s/mm² to calculate D* and f (33). In this study, 11 b values were acquired, among which 9 b values (0, 10, 20, 50, 100, 200, 400, 800, and 1,200 s/mm²) were selected for IVIM fitting and 8 b values (0, 100, 200, 400, 800, 1,200, 1,600, and 2,000 s/mm²) were selected for DKI fitting. The sets of b values were sufficient to estimate

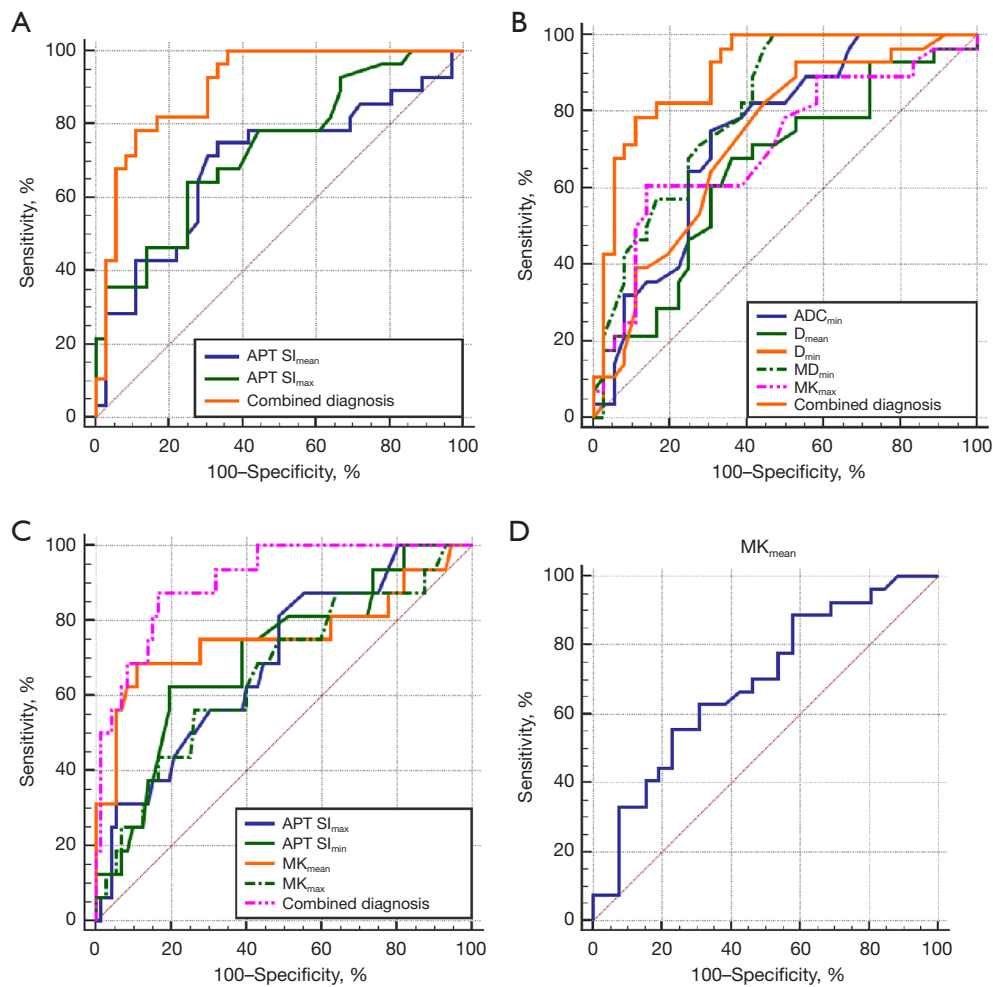


Figure 4 ROC curves of APT SI_{mean} , APT SI_{max} , ADC_{min} , D_{mean} , D_{min} , MD_{min} , MK_{max} , and the combined model (APT SI_{mean} + APT SI_{max} + ADC_{min} + MK_{max} + MD_{min}) for differentiating high-grade and low-grade cervical cancer (A,B). ROC curves of APT SI_{max} , APT SI_{min} , MK_{mean} , MK_{max} , and the combination of all four metrics for differentiating squamous cell carcinoma and adenocarcinoma (C). ROC curve of MK_{mean} for differentiating LVS-positive and LVS-negative cervical cancer (D). APT SI, amide proton transfer signal intensity; ADC, apparent diffusion coefficient; D, pure molecular diffusion; MD, mean diffusivity; MK, mean kurtosis; ROC, receiver operating characteristic; LVS, lymphovascular space invasion.

IVIM and DKI metrics according to the previous studies (32,33). Additionally, our study had a high proportion of SCC, whereas other studies had a more balanced distribution of SCC and AC cases (11,29); this could have influenced our results. Therefore, further studies with large samples and multiple centers are needed to confirm the role of different DWI metrics in cervical cancer evaluation.

We have also applied multivariable analysis to the identification of SCC and AC and the identification of high- and low-grade cervical cancer. Our results revealed that among APT and DWI metrics, APT SI_{max} , APT SI_{min} , and

MK_{mean} were the independent predictors for tumor subtype. APT SI_{max} and MD_{min} were the independent predictors for the grade of cervical cancer. The combined diagnostic model (APT SI_{max} + APT SI_{min} + MK_{mean} + MK_{max}) revealed additional value in comparison to any individual metric for distinguishing between SCC and AC with an AUC of 0.908. The combination of APT and DWI metrics also significantly improved the diagnostic performance of high-grade cervical cancer than that of individual metrics with an AUC of 0.903. The results of additional value of APT to DWI were basically consistent with the conclusions of

Choi *et al.* and Meng *et al.* (25,34). Choi *et al.* found that APT imaging added value to the ADC for discriminating high-grade from low-grade gliomas (25). Meng *et al.* demonstrated that the AUC of the combination of D, MK, and APT SI was significantly higher than that of any single metric for risk stratification in early-stage endometrial carcinoma (34). These findings support the hypothesis that APT imaging complement the multi-model DWI for cervical cancer typing and grading, when the APT signal reflects a different aspect of the tumor microenvironment in comparison to the DWI metrics, namely the protein and peptide concentrations.

In addition to the benefit to cervical cancer patients from quantitative MR parameters, the practicability of APT imaging in the clinical real-world setting should also be considered. 3D APT MRI in our study required a scan time of about 6 minutes. Although the fast-imaging acquisition technique (3D TSE readout) has been used to accelerate the sequence, the use of multiple radio-frequency (RF) saturation frequencies (to correct for the B0 inhomogeneity) and multiple acquisitions (to increase the signal-to-noise ratio) extend the scan time (35). A next step in APT sequence may focus on the development of a fast 3D acquisition technique by combining some novel undersampling acquisition and reconstruction approaches (including keyhole, compressed sensing, and deep learning). It is known that quantification of APT signal is sensitive to accuracy of saturation pulse (B1), B0 inhomogeneity, and lipid artifacts (36). We used the APT imaging protocol that employed the multi-transmit RF system for the specific-frequency saturation with amplitude of 2 μ T and duration of 2 seconds with sufficient B1 homogeneity. Additionally, B0 homogeneity correction was performed and the Spectral Attenuation with Inversion Recovery module was used to suppress strong lipid artifacts in the pelvic APT imaging. A previous study evaluated the image quality and clinical feasibility of 3D APT MRI for cervical cancer and showed the excellent agreement in both image quality assessment and APT values measurement (37). Lee *et al.* found that APT MRI showed high repeatability under the condition of the same acquisition protocol and the same MRI machine (38). However, little is known about the repeatability of APT signal measurement in different platforms. In this respect, it is encouraging to see that the recent 8th International Workshop on chemical exchange saturation transfer (CEST) Imaging [2020] provided consensus recommendations for APT imaging of brain tumors involving the three main MRI vendors. They indicated that APT imaging methods would be optimized

and standardized separately for other body regions in a few years (39).

Several limitations must be considered when interpreting the present results. First, the sample size of AC patients was small due to the relatively low incidence (40). Some rare subtypes of cervical cancer such as adenosquamous carcinoma and neuroendocrine carcinomas were excluded. Our observations need to be validated in multicenter studies with maximum AC cases and other rare subtypes. Second, we only evaluated the relationship between the imaging metrics and traditional prognostic factors of cervical cancer. Several recent studies have clarified the importance of the molecular status of the tumor (40). We should conduct further studies to investigate the potential associations between imaging metrics and some molecular prognostic factors of cervical cancer (41,42). Third, the histological reference of some cases recruited in our study was based on biopsy samples; therefore, although the results are promising especially for tumor subtype characterization, further investigation with larger study cohorts confirmed by surgical histopathology should be conducted to validate our preliminary study in the future.

Conclusions

In conclusion, our findings indicate that 3D-APT imaging and multi-model DWI are useful for cervical cancer typing and grading. DKI-derived MK_{mean} may provide helpful information to assist in the prediction of LVSI. In combination with multi-model DWI, 3D-APT showed the potential to significantly improve the performance in differentiating tumor subtypes and grades.

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Footnote

Reporting Checklist: The authors have completed the STARD reporting checklist. Available at <https://qims.amegroups.com/article/view/10.21037/qims-23-324/rc>

Conflicts of Interest: All authors have completed the ICMJE

uniform disclosure form (available at <https://qims.amegroupp.com/article/view/10.21037/qims-23-324/coif>). LL, an MR collaborating scientist from Philips Healthcare, provided technical support under the Philips collaboration regulations and had no financial or other conflicts for this study. The other authors have no conflicts of interest to declare.

Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013). The study was approved by the institutional ethics board of The First Affiliated Hospital of Zhengzhou University, and all patients provided written informed consent prior to enrollment in the study.

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