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Quantitative Assessment of Diffusion-Weighted MR Imaging in Patients with Primary Rectal Cancer: Correlation with FDG-PET/CT

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Abstract

Purpose: The aim of the study was to assess correlations between parameters on diffusion-weighted imaging and 2-deoxy-2-[18F]fluoro-D-glucose–positron emission tomography/computed tomography (FDG-PET/CT) in rectal cancer.

Procedures: Thirty-three consecutive patients with pathologically confirmed rectal adenocarcinoma were included in this study. Apparent diffusion coefficient (ADC) maps were generated to calculate ADC mean (average ADC), ADC min (lowest ADC), tumor volume, and total diffusivity index (TDI). PET/CT exams were performed within 1 week of magnetic resonance imaging. Standardized uptake values (SUVs) were normalized to the injected FDG dose and body weight. SUV max (maximum SUV), SUV mean (average SUV), tumor volume, and total lesion glycolysis (TLG) were calculated using a 50% threshold.

Results: Significant negative correlations were found between ADC min and SUV max ($r = -0.450$, $p = 0.009$), and between ADC mean and SUV mean ($r = -0.402$, $p = 0.020$). A significant positive correlation was found between TDI and TLG ($r = 0.634$, $p < 0.001$).

Conclusion: The significant negative correlations between ADC and SUV suggest an association between tumor cellularity and metabolic activity in primary rectal adenocarcinoma.

Key words: ADC, DWI, SUV, PET/CT, Primary rectal adenocarcinoma, TDI, TLG

Introduction

Colorectal cancer is the second leading cause of cancer death for men and women combined [1], and rectal cancer constitutes about one third of cases. Magnetic resonance imaging (MRI) has the advantages of superior soft tissue contrast and multiplanar imaging capability, thus often considered particularly useful for local staging. On the other hand, 2-deoxy-2-[18F]fluoro-D-glucose–positron emission tomography/computed tomography (FDG-PET/CT) is being more and more commonly used for detecting distant metastases and providing systemic staging of patients with rectal cancer.

Recent studies show that both MRI and PET have the capability of providing important functional information for tumor, in addition to the above-mentioned staging capacity. For example, with advances in MR technology and the use of faster, more robust sequences, diffusion-weighted imaging (DWI), an established diagnostic technique in neuroimaging, has shown great potential for evaluation of tumors in the body as well [2]. DWI is used to measure the Brownian motion of water molecules in tissue, which has been shown to be inversely proportional to cellular density [3], presumably because...
increased cellular density limits water diffusion in the interstitial space. The apparent diffusion coefficient (ADC), a quantitative parameter measured on DWI, has been shown to be useful for evaluating solid tumors in the abdomen and pelvis [4, 5]. It has been suggested that ADC may provide useful information regarding tumor cellularity [6], tumor aggressiveness [7], subtype characterization [8], and cancer treatment response [2, 9, 10]. However, the experience of applying DWI in evaluation of rectal cancer is relatively limited. It has been shown that DWI can improve detection of rectal cancer when combined with T2-weighted images [11] and may have a potential role in predicting tumor response to therapy [12, 13].

FDG-PET is capable of imaging tumors based on their increased glucose metabolism. FDG uptake on PET, quantified by the standardized uptake value (SUVs), is a useful marker for the level of tumor metabolic activity. It has been reported that high SUV is correlated with rapid cellular proliferation in breast, lung, and ovarian cancers [14–16]. A number of studies demonstrated the relationship between higher FDG uptake and a more aggressive course of the malignancy [17, 18]. Meanwhile, the role of PET/CT to assess treatment response has been established in multiple types of tumors [19, 20]. In rectal cancer imaging, PET/CT has been suggested as an accurate technique in the detection or staging of newly diagnosed or recurrence rectal cancer [21, 22]. Furthermore, qualitative assessment of FDG-PET provides useful information of treatment response and prognostic information in patients with rectal cancer [23, 24].

Even though both DWI and PET/CT have been used in various aspects of tumor evaluation, including detection, characterization, and treatment response assessment, to our knowledge, the relationship between ADC and SUV has not been explored in rectal cancer. Since both ADC and SUV values have been associated with biological aggressiveness and treatment response in certain tumors including rectal cancer, we hypothesize that SUV values which reflect metabolic activity may have correlation with ADC values which reflect cellular density in rectal cancer. Thus, the aim of this study was to assess the correlations between quantitative parameters on DWI and PET in primary rectal cancer.

Materials and Methods

Patients

Between November 2008 and June 2009, 33 consecutive patients (19 men, 14 women, 45–88 years old) with newly diagnosed and pathologically confirmed rectal adenocarcinoma and having undergone both MRI and PET/CT examinations were included in this study. All patients underwent both MRI and PET/CT scans within 1 week (mean time interval, 2±1 day). This study was approved by Institutional Review Board, and informed consent was obtained from all patients. This study was also compliant with the patient confidentiality regulations.

Acquisition of MR Images

All MR examinations were performed on a 3-T scanner (Achieva; Philips Healthcare) with the patients in supine position. Routine T2-weighted, T1-weighted, DWI, and contrast-enhanced sequences were obtained. Turbo spin echo T2-weighted images were obtained in three planes using the following parameters: transverse, TR/TE=1,862/99 ms, field of view (FOV)=19×23 cm, matrix size=272×318, slice thickness=6 mm, gap=0, number of acquisition=1, sense factor=1.5; coronal, TR/TE=2,800/100 ms, FOV=25×26 cm, matrix size=416×373, slice thickness=5 mm, gap=0.5 mm, number of acquisition=1, sense factor=1.5; sagittal, TR/TE=2,800/100 ms, FOV=23×24 cm, matrix size=328×341, slice thickness=5 mm, gap=0, number of acquisition=1, sense factor=1. Axial TSE T1-weighted images were obtained using the following parameters: FOV=23×38 cm, matrix size=236×314, slice thickness=7 mm, gap=1 mm, number of acquisition=1, sense factor=1. Transverse free-breathing DWI was obtained by using a single-shot multi-slice echo planar imaging (EPI) sequence with short TI inversion recovery (STIR) fat-suppression and slice-selection gradient reversal technique [25] with the following parameters: TR/TE=7,036/48 ms, FOV=40×33 cm, matrix size=188×159, slice thickness=5 mm, gap=0, number of acquisitions=4, sense factor=2, b value=0 and 1,000 s/mm². The acquisition time for DWI was approximately 4 min. Transverse single-shot TSE T2-weighted images were then obtained with slice locations identical to those of transverse DWI for image fusion (OsimirX Medical Imaging Software, version 3.5, Switzerland), using the following parameters: TR/TE=891/98 ms, FOV=40×33 cm, matrix size=332×125, slice thickness=5 mm, gap=0, number of acquisitions=1. Subsequently fast field echo (FFE) T1-weighted images were obtained before and after administration of intravenous contrast. Flex-L coil with using SENSE technique was placed over the pelvis to reduce gas effect to imaging quality. Patients were asked to fast for 4 h before MR.

After image acquisition, pixel-to-pixel ADC map was reconstructed using the standard software on the imaging console (Achieva; Philips Healthcare).

Acquisition of PET/CT Images

All PET/CT examinations were performed on a PET/CT scanner (Discovery VCT, GE Healthcare Bio-Sciences Corp.). All patients fasted with hydration 6 h before receiving intravenous injection of FDG at 4.8 MBq/kg body weight. PET/CT scans were performed 60 min after the injection of the FDG. A whole body emission PET scan with a 70-cm axial FOV, a 218×218 matrix, and 3.27 mm thickness was obtained with five bed positions within 20 min. CT images were performed using the following scan parameters: FOV=60 cm, matrix=512×512, collimation=0.625 mm×64, pitch=0.984, gantry rotation speed=0.5 s, tube voltage=120 kVp, and tube current=200–400 mA. The use of intravenous contrast for CT was at the discretion of the ordering physician. LASIX (4 mg) was administered intravenously to fill the bladder to reduce artifacts from high 18F-FDG activity in urine. Attenuation correction was performed on PET images with CT data using an ordered-subset expectation maximization
iterative reconstruction algorithm (14 subsets and two iterations) [26]. The CT images were then reconstructed at 2.5-mm intervals to fuse with the PET images (Advanced Workstation 4.3, GE Healthcare).

**Image Analysis**

For ADC measurement, regions of interest (ROIs) were manually drawn along contours of each tumor on ADC maps on every slice covering the entire tumor by an investigator (J. G., with 3 years of experience in reading and performing volumetric tumor measurements on body MRI) who was blinded to PET/CT images and all clinical information other than that the patient was diagnosed with rectal cancer.

Mean ADC value (ADC$_i$) and the cross-sectional area (area$_i$) of the tumor ROI on each slice ($i$ representing the slice number) was calculated by Image J software (NIH, USA). Subsequently, ADC$_{mean}$ of the entire tumor was calculated as the weighted average for all ADC$_i$ values in each tumor by Eq. 1:

$$ADC_{mean} = \frac{\sum_i (ADC_i \times Area_i)}{\sum_i Area_i} \quad (1)$$

We calculated weighted averages because this would be mathematically identical to calculating averages of ADC values directly from all voxels within the entire tumor volume. ADC$_{min}$ was also determined as the lowest ADC value among all voxels in each tumor. Ratio of ADC (rADC) was then calculated as ADC$_{min}$/ADC$_{mean}$, which showed a significant correlation with SUV$_{max}$/SUV$_{mean}$ ratio (rSUV) in cervical cancers [27]. Volume of the rectal tumor on DWI images was calculated by Eq. 2:

$$V_{DWI} = \sum_i Area_i \times (\text{thickness} + \text{gap}) \quad (2)$$

We also introduced the concept of total diffusivity index (TDI), which is the sum of 1/ADC among all the voxels in a tumor calculated by home-made MATLAB script using Eq. 3, as a corresponding value to total lesion glycolysis (TLG) measured from PET:

$$TDI = V_{DWI} \times \frac{\sum_i (D_i \times Area_i)}{\sum_i Area_i} \quad (3)$$

where $V_{DWI}$ is calculated by Eq. 2; where $D$ is the mean 1/ADC for all voxels on each ROI area. This equation would be mathematically identical to calculating sums of 1/ADC values directly from all voxels within the entire tumor volume.

For SUV measurement, the PET, CT, and fused PET/CT images were displayed on a workstation (Advanced Workstation, 4.3, GE Healthcare), and a 3D ROI was placed over the entire tumor by an investigator (J. Z., with 2 years of experience in interpreting PET/CT) who was blinded to MRI images, and all clinical information other than that the patient was diagnosed with rectal cancer.

Maximum standardized uptake value (SUV$_{max}$), mean standardized uptake value (SUV$_{mean}$), volume of tumor, and TLG were automatically calculated by the workstation. SUV was defined by Eq. 4:

$$SUV = \frac{\text{measured radioactivity concentration [Bq/mL]}}{\text{injected radioactivity [Bq]/(bodyweight[kg] \times 1,000)}} \quad (4)$$

SUV$_{max}$ was defined as the highest value of SUV among all voxels within the 3D ROI placed over the rectal tumor. Subsequently, a fixed threshold value of 50% of the maximum uptake was used to determine tumor margins automatically, and the tumor volume ($V_{PET}$) was calculated by the workstation accordingly [22]. The SUV$_{mean}$ was then measured as the average of SUV values in all voxels within the threshold-defined tumor volume. rSUV was then calculated as SUV$_{max}$/SUV$_{mean}$. TLG was calculated using Eq. 5:

$$TLG = V_{PET} \times SUV_{mean} \quad (5)$$

**Clinical Correlation**

All patients had biopsy under colonoscopy that confirmed histopathological diagnosis of rectal cancer before the imaging examinations were performed (mean time interval, 15±4 days). Circulating plasma level of carcinoembryonic antigen (CEA) was obtained within 2 weeks of MRI scan (mean time interval, 9±3 days). Based on the references from our institution, CEA $>$ 5.0 mg/mL was regarded as abnormal.

**Statistical Analysis**

All results were expressed as mean±standard deviation (SD). Pearson’s correlation test was used to detect the relationships between quantitative indices of DWI and PET. Bland–Altman plot was performed to assess agreement between volumes measured on DWI and PET. One-way ANOVA test was used to analyze differences of SUV and ADC values in terms of well, moderate, or poorly differentiated rectal cancer. A $p$ value of $<$0.05 was considered to indicate statistical significance. All statistical analyses were performed using SPSS software package (SPSS, Version 16.0.1, Chicago, IL, USA).

**Results**

**Patient and Lesion Characteristics**

Among the 33 patients included in this study, 19 were men and 14 were women. They had a mean age of 67 years (standard deviation, 11 years; range, 45–88 years). All patients were confirmed to have rectal adenocarcinomas by pathological evaluations. Characteristics of study subjects are shown in Table 1.

| ADC$_{min}$ and ADC$_{mean}$ for all tumors were 0.35±0.15×10$^{-3}$ mm$^2$/s (range, 0.10–0.75×10$^{-3}$ mm$^2$/s) and 0.85± |
0.24×10\(^{-3}\) mm\(^2\)/s (range, 0.43~1.5×10\(^{-3}\) mm\(^2\)/s), respectively. rADC was 0.42±0.15 (range, 0.12~0.67). TDI was 27.25±30.64×10\(^3\) ms (range, 2.54~175.01×10\(^3\) ms). Tumor volume measured by DWI was 22.6±24.5 cm\(^3\) (range, 2.87~135.8 cm\(^3\)). Fig. 1 demonstrates the measurement of ADC values and tumor volume on DWI in a typical case.

SUV\(_{\text{max}}\) and SUV\(_{\text{mean}}\) values were 10.55±5.26 (range, 4.10~26.10) and 6.94±3.56 (range, 2.70~17.5), respectively. rSUV was 0.65±0.04 (range, 0.54~0.73). TLG of all tumors was 99.8±98.7 g (range, 8.5~386.5 g). Tumor volume measured by PET/CT was 15.0±12.0 cm\(^3\) (range 2.4~64.5 cm\(^3\)), using 50% of SUV\(_{\text{max}}\) as threshold. Fig. 2 demonstrates the measurement of SUV, TLG, and tumor volume on PET/CT in a typical case.

**Correlations Between ADC and SUV**

Significant negative correlations were found between ADC\(_{\text{min}}\) and SUV\(_{\text{max}}\) (r=−0.450, p=0.009, SUV\(_{\text{max}}\)=−15.825ADC\(_{\text{min}}\)+16.081), and ADC\(_{\text{mean}}\) and SUV\(_{\text{mean}}\) (r=−0.402, p=0.020, SUV\(_{\text{mean}}\)=−5.884ADC\(_{\text{mean}}\)+11.948) (Fig. 3). The correlation between rADC and rSUV was weak (r=−0.161, p=0.370).

**Correlations Between TDI and TLG**

Significant correlation was found between TDI and TLG (r=0.634, p<0.001, TLG=2.044TDI+44.114).

**Table 1. Characteristics of patients and lesions**

<table>
<thead>
<tr>
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<tr>
<td>Age</td>
<td>67±11 years</td>
</tr>
<tr>
<td>Gender</td>
<td>Female (n=14), male (n=19)</td>
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<tr>
<td>Serum CEA level</td>
<td>34.4±69.7 mg/mL</td>
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<td>Patients with normal CEA (≤5.0 mg/mL)</td>
<td>20 (60.6%)</td>
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<tr>
<td>Patients with abnormal CEA (&gt;5.0 mg/mL)</td>
<td>13 (39.4%)</td>
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<tr>
<td>Pathological diagnosis</td>
<td>Number of lesions (%)</td>
</tr>
<tr>
<td>Well-differentiated adenocarcinoma</td>
<td>10 (30%)</td>
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<tr>
<td>Moderately differentiated adenocarcinoma</td>
<td>14 (42%)</td>
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<tr>
<td>Poorly differentiated adenocarcinoma</td>
<td>9 (27%)</td>
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<tr>
<td>Clinical staging</td>
<td>Number of patients (%)</td>
</tr>
<tr>
<td>Stage I</td>
<td>1 (3%)</td>
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<tr>
<td>Stage II</td>
<td>5 (15%)</td>
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<tr>
<td>Stage III</td>
<td>24 (73%)</td>
</tr>
<tr>
<td>Stage IV</td>
<td>3 (9%)</td>
</tr>
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</table>

**Fig. 1.** MRI images of a 55-year-old man with moderately differentiated rectal adenocarcinoma. **a** Axial single-shot TSE T2-weighted image of the pelvis shows a large rectal mass (arrow). **b** Axial DWI image of the pelvis at the same slice location as a is shown with inverted gray scale to demonstrate a PET-like image of the rectal mass (arrow). **c** Fused image from **a** and **b** can be performed for easy viewing if desired. **d** On the ADC map generated from **b**, an ROI was manually drawn along the contour of the tumor (black line). Subsequently, ADC\(_{\text{mean}}\), ADC\(_{\text{min}}\), and the cross-sectional area of the tumor on this image were calculated by ImageJ software.
Agreement Between Tumor Volumes Measured on MR and PET

A Bland–Altman plot analysis, which is used to compare the mean of the differences and limits of agreement (average difference±1.96 standard deviation of the difference) between two methods, showed good agreement between tumor volumes measured on DWI and PET (Fig. 4).

However, by average, by using the standard 50% threshold, PET underestimated the rectal cancer volume compared with DWI.

Correlations Between Imaging Parameters and Cell Differentiation of Rectal Cancer

Table 2 shows SUV and ADC values in well-differentiated, moderately differentiated, and poorly differentiated rectal cancers respectively. Although there was a trend of having higher SUV and lower ADC_min values in the poorly differentiated tumors, these differences did not reach statistical significance. However, this could be related with a suboptimal statistical power associated with the relatively small patient population in this study.

Discussion

Both DWI and PET/CT are established imaging modalities in tumor assessment. It has been shown that tumors often demonstrate decreased ADC on DWI, and increased SUV on FDG-PET. In addition, ADC and SUV have both been shown to correlate with tumor characteristics and response to treatment. Therefore, we undertook this study to assess the relationship between quantitative parameters measured on DWI and PET in primary rectal cancer.

First, we found significant negative correlations between ADC and SUV values in rectal cancer. SUV values are quantitative indices provided by PET/CT and represent the activity concentration in tumor tissue normalized to the injected FDG dose and body weight of the patient. Intravenous injected FDG is transported across the cell membrane by glucose transporters (Gluts) and accumulated in metabolically active tumor cells. SUV values have correlation with tumor cellular density as well as the grade and the differentiation of some tumors and can aid in detection, characterization, prognostication, and monitoring treatment response of malignancy [17, 28–33]. SUV correlated significantly with cellular density in non-small cell lung cancer [32]. In astrocytomas, Herholz et al. [31] found that glucose consumption correlated significantly with cell density which is a major determinant of glucose consumption in astrocytomas. In rectal cancer, tumor size and depth of invasion of rectal cancer were significantly correlated with SUV [30]. On the other hand, ADC is a quantitative parameter provided by DWI and reflects water diffusion in tissue. There is decreased ADC in tumor tissue due to increased diffusion barrier from tumor cell membrane. ADC has been regarded as a useful imaging marker to reflect tumor cell density and to distinguish different tissue compartments in early, intermediate, and advanced tumor stages [34, 35]. Variation of ADC has been used to monitor changes in the biological structure of tumor tissue during tumor progression. In early stages of tumor development, tumors appeared homogeneous and have lower ADC values. In contrast, there is an increased ADC that correlated well
with areas of necrosis (reduced cell density) in intermediate and advanced tumor stage [34]. Although a significant correlation between tumor FDG uptake and tumor cellular density was found in some studies [16, 36], there is limited information about the relationship between SUV and ADC. Ho et al. made an important attempt but did not find significant correlations between SUVmax and ADC values in cervical cancer [27]. There may be several potential causes for this discrepancy. One is that imaging parameters may very well have different diagnostic performances for different types of tumors (cervical versus rectal cancer). Another potential cause is that mixed pathological types were present in Ho’s study on cervical cancer patients, including squamous cell carcinoma, adenocarcinoma, and adenosquamous carcinoma. The mixed pathologies may conceivably lead to obscuration of an underlying correlation. In addition, Ho et al. calculated ADCmean by taking averages of mean ADC values for each image, regardless of the cross-sectional size of the tumor on each image [27]. We feel that a weighted average taking into consideration of the tumor cross-sectional size on each image is a more accurate calculation to reflect true mean ADC value of all voxels in the entire tumor.

We also found that TDI values, defined as the sum of 1/ADC among all the voxels in a tumor, showed a significant correlation with TLG, which is the sum of SUV among all the voxels in a tumor. TLG reflects the total amount of glycolysis in a given tumor and has been used clinically as a surrogate biomarker for monitoring treatment response [37–39]. For example, Guillem et al. found that a reduction of 30.5% of TLG in rectal cancer could predict no-evidence-of-disease status and freedom from recurrence with a sensitivity of 90% and specificity of 80% [38]. The correlation between TDI and TLG is not surprising, as the ADC and SUV values, as well as the tumor volumes measured by MR and PET, are correlated. Since tumor’s response to treatment may be reflected in changes of both size and function, we feel that TDI values may have a potential role in monitoring treatment response similar to TLG. However, longer-term studies with bigger patient populations would be helpful to validate this.

In terms of correlation between the imaging parameters and tumor pathologic characteristics, we did see a trend of higher SUV values and lower ADC values in the more poorly differentiated rectal tumors. This is not surprising given that less differentiated tumors metabolize more glucose for energy production and grow more rapidly.
However, the differences between poorly, moderately, and well-differentiated rectal tumors did not reach statistical significance. It is possible that the relative small patient population in our study may have lead to suboptimal statistical powers. Larger studies in the future would be helpful to further evaluate these potential correlations. Another possible reason is that FDG-PET might be insensitive in showing mucinous carcinomas because of the low cellularity of these tumors caused by presence of mucin which is commonly found in colonic tumors [40]. Significantly lower FDG uptake was found in bronchioloalveolar cell carcinoma, which often contains abundant mucin, compared with adenocarcinoma, in which positive correlation was found between FDG uptake and the degree of cell differentiation [41, 42]. On the other hand, one study showed high ADC value in poorly differentiated rectal adenocarcinoma, including mucinous carcinoma and signet ring cell carcinoma [43]. And no overt correlation was found between ADC and cell differentiation due to a lot of mucin contained in the rectal adenocarcinoma [43]. In terms of measuring tumor volume on MR, although T2-weighted images are considered to provide superior anatomic details, we measured tumor volume on ADC maps, so that the calculation of TDI would be solely based on parameters generated from DWI images. A high correlation of volumes detected by DWI and PET/CT was demonstrated in the Bland–Altman plot. In Fig. 4, the outlying data point on the right upper corner represented a large tumor with MRI volume of 135 cm$^3$, which was vastly underestimated in volume on PET (64.5 cm$^3$) when 50% threshold was used, presumably due to an unusual distribution of SUV values in the tumor voxels. If a 30% threshold was used, the tumor volume on PET would have been 142 cm$^3$. Moreover, by average, tumor volumes measured on DWI was 7.6 cm$^3$ greater than those measured on PET in this study. Similar findings have been reported in the literature. Daisne et al. compared the diagnostic accuracy of measuring head and neck tumor volumes by different imaging modalities including CT, PET, and MRI [44]. They found that PET provided the smallest tumor volumes among the three modalities. The smallest tumor volumes on PET may be related to the value of threshold used for automated tumor volume measurement. In this study, we used a threshold of 50% SUV$_{max}$ to automatically determine tumor boundary on PET. Usually, 40% or 50% threshold is suggested to evaluate rectum cancer by PET [22]. However, currently, there is no clinical consensus as to what threshold can provide accurate information of tumor boundary [45, 46], and it is possible that different types of tumors may require different thresholds. It has been suggested that using an optimal SUV threshold on PET may provide more accurate estimation of tumor volume, with the optimal threshold being defined as the threshold at which PET shows a similar tumor volume compared with CT or MRI [27].

With fast imaging techniques such as EPI and parallel imaging techniques [47], DWI can be performed with diagnostic quality at high $b$ values in the body [9], which lead to heightened interests in investigating its role in body and oncologic imaging. Takahara et al. developed the concept of diffusion-weighted whole body imaging with background body signal suppression (DWIBS) and introduced the feasibility of free-breathing DWI combined with STIR-EPI techniques [48]. In our study, a similar technique was used, with additional slice-selection gradient reversal technique to aid in more complete suppression of the fat signal. Due to good background body signal suppression, DWIBS can produce PET-like images, which has been widely used for oncologic imaging. However, with the longer survival rate seen in many tumor patients due to improved clinical care, patients will get more and more radiation exposure from surveillance imaging studies such as PET/CT. In addition, the contrast injections for both PET and CT are associated with certain complications, although relatively rare. Therefore, if DWI parameters such as ADC values can be validated clinically for accurate tumor assessment, whole-body MR techniques such as DWIBS may provide a safe alternative imaging modality for the oncologic patient.

There are certain limitations to our study. First, the number of subjects in our study is relatively small. However, we found some significant correlations between MR and PET parameters despite the relatively small sample size. Second, although imaging correlations were demonstrated in this current study, pathological evidence, such as measurements of cellularity, Gluts, and hexokinase, is lacking to support our findings. Thirdly, imaging parameters may have different diagnostic performances for different types of tumors. Therefore, the correlations we found in rectal adenocarcinomas may not be transferrable to other types of tumors, and each tumor type should be tested individually.

### Table 2. Imaging parameters and tumor differentiation

<table>
<thead>
<tr>
<th>Imaging Parameter</th>
<th>Well differentiated</th>
<th>Moderately differentiated</th>
<th>Poorly differentiated</th>
<th>$p$ value</th>
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<tbody>
<tr>
<td>$n=10$</td>
<td>$n=14$</td>
<td>$n=9$</td>
<td></td>
<td></td>
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<tr>
<td>SUV$_{max}$</td>
<td>9.01±4.44</td>
<td>11.18±6.00</td>
<td>11.30±5.08</td>
<td>0.55</td>
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<tr>
<td>SUV$_{mean}$</td>
<td>5.88±2.92</td>
<td>7.31±4.15</td>
<td>7.52±3.33</td>
<td>0.54</td>
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<tr>
<td>ADC$_{min}$</td>
<td>0.34±0.16</td>
<td>0.40±0.15</td>
<td>0.28±0.13</td>
<td>0.16</td>
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<tr>
<td>ADC$_{mean}$</td>
<td>0.80±0.25</td>
<td>0.84±0.30</td>
<td>0.82±0.13</td>
<td>0.83</td>
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Conclusion

Our results indicate that DWI parameters such as ADC and TDI are correlated with PET parameters such as SUV and TLG in primary rectal cancer. Future larger clinical trials in a variety of tumors would be helpful to confirm our preliminary findings and further determine the roles of ADC and TDI in oncologic imaging.

Conflict of interest. The authors declare that they have no conflict of interest.

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