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Multiparametric Magnetic Resonance Imaging of the Prostate
Repeatability of Volume and Apparent Diffusion Coefficient Quantification

Andriy Fedorov, PhD,* Mark G. Vangel, PhD,† Clare M. Tempany, MD,*, and Fiona M. Fennessy, MD, PhD*‡

Objectives: The aim of this study was to evaluate the repeatability of a region of interest (ROI) volume and mean apparent diffusion coefficient (ADC) in standard-of-care 3 T multiparametric magnetic resonance imaging (mpMRI) of the prostate obtained with the use of endorectal coil.

Materials and Methods: This prospective study was Health Insurance Portability and Accountability Act compliant, with institutional review board approval and written informed consent. Men with confirmed or suspected treatment-naïve prostate cancer scheduled for mpMRI were offered a repeat mpMRI within 2 weeks. Regions of interest corresponding to the entire prostate gland, the entire peripheral zone (PZ), normal PZ, and suspected tumor ROI (tROI) on axial T2-weighted, dynamic contrast-enhanced subtract, and ADC images were annotated and assessed using Prostate Imaging Reporting and Data System (PI-RADS) v2. Repeatability of the ROI volume for each of the analyzed image types and mean ROI ADC was summarized with repeatability coefficient (RC) and RC%.

Results: A total of 189 subjects were approached to participate in the study. Of 40 patients that gave initial agreement, 15 men underwent 2 mpMRI examinations and completed the study. Peripheral zone tROIs were identified in 11 subjects. Tumor ROI volume was less than 0.5 mL in 8 of 11 subjects. PI-RADS categories were identical between baseline-repeat studies in 11/15 subjects and differed by 1 point in 4/15. Peripheral zone tROI volume RC% was 233 mm³ (71%) on axial T2-weighted, 422 mm³ (112%) on ADC, and 488 mm³ (119%) on dynamic contrast-enhanced subtract. Apparent diffusion coefficient ROI mean RC% were 447 × 10−6 mm²/s (42%) in PZ tROI and 471 × 10−6 mm²/s (30%) in normal PZ. Significant difference in repeatability of the tROI volume across series was observed (P < 0.005). The mean ADC RC% was lower than volume RC% for tROI ADC (P < 0.05).

Conclusions: PI-RADS v2 overall assessment was highly repeatable. Multiparametric magnetic resonance imaging sequence differences in volume measurement repeatability. The mean tROI ADC is more repeatable compared with tROI volume in ADC. Repeatability of prostate ADC is comparable with that in other abdominal organs.

Key Words: magnetic resonance imaging, prostate, apparent diffusion coefficient, dynamic contrast-enhanced imaging, repeatability, quantitative imaging

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Table 1. Ranges of the Acquisition Parameters for the Analyzed MRI Image Types

<table>
<thead>
<tr>
<th>Repetition Time, ms</th>
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</tr>
</thead>
<tbody>
<tr>
<td>T2AX</td>
<td>3350–5109</td>
<td>84–107</td>
<td>140–200</td>
<td>3</td>
</tr>
<tr>
<td>DCE</td>
<td>3.68–4.1</td>
<td>1.3–1.42</td>
<td>260–280</td>
<td>5–6</td>
</tr>
<tr>
<td>DWI</td>
<td>2500–8150</td>
<td>76.7–80.6</td>
<td>160–280</td>
<td>3–4</td>
</tr>
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MRI indicates magnetic resonance imaging; T2AX, T2-weighted axial image; DCE, dynamic contrast-enhanced image; DWI, diffusion-weighted image.

The difference between the phase corresponding to the contrast bolus arrival and the baseline phase were computed using the scanner software.

Image Annotation and Quantitation

Imaging studies were de-identified and presented in a randomized fashion (reader was blinded to the given study being a baseline or repeat study, and the studies were presented in a permuted order) to an abdominal radiologist (F.F., 12 years in prostate mpMRI interpretation) using 3D Slicer software (http://slicer.org). A consistent hanging protocol was applied to present all of the imaging series required for mpMRI assessment, as specified by PI-RADS v2. Results of the prior PSA tests and systematic transrectal ultrasound (TRUS) biopsy (when available as part of the medical record) were presented to the reader as an aid in tumor localization, per standard clinical interpretation workflow.

For each study, the reader confirmed its diagnostic quality, and that SUB was based upon subtraction of baseline precontrast images from first bolus arrival phase. Regions of interest were manually outlined on the T2AX, ADC, and SUB series for the whole gland (WG), peripheral zone (PZ), normal PZ (nPZ), and (when present) tumor-suspicious ROI (tROI). The reader could see the matching slice in all other series for the given study but was blinded to the ROIs contoured on other sequences of that study and to contours of the paired imaging study. Overall, all PI-RADS v2 assessment category between 1 and 5 was assigned for each study. The sector(s) containing the lesion were noted. Annotated ROIs identifying voxels corresponding to the suspected lesions were saved as rasterized segmentations using 3D Slicer software for subsequent extraction of ROI statistics.

Agreement in locations of tROI areas was assessed by a separate reader by comparing agreement of the noted lesion sector(s) between baseline and repeat studies. We did not attempt to validate the location of the annotated lesion with either biopsy or WG pathology. Because the data were collected prospectively under standard-of-care conditions, neither targeted biopsy sampling nor whole mount prostatectomy pathology were available to us. Spatial voxel-wise correspondence between the contoured regions was not attempted due to expected misregistration between the baseline and repeat examinations. Volumes of the WG, PZ, and tROI regions, and mean ADC value for the WG, PZ, nPZ, and tROI areas were automatically calculated from the ROIs.

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MRI indicates magnetic resonance imaging; T2AX, T2-weighted axial image; DCE, dynamic contrast-enhanced image; DWI, diffusion-weighted image.
due to ROI and image type, for each structure using 2-way analysis of variance mixed model regression, whereas adjusting for multiple comparisons using the approach of Bretz et al.33 Standard error and confidence interval for RC and RC% were estimated using the delta method. RC% of ADC volume and mean ADC were compared using 2-tailed $z$ test. Results were tabulated and summarized using Bland- Altman plots.34 Analysis was performed using R 3.1.0.35

RESULTS

Population
A total of 189 patients (the 189 patients we approached amounts to approximately 7.5% of the estimated total volume of the prostate MRI collected at our institution over the course of the study enrollment, considering typical volume of 5 prostate MRI patients per day) were offered to undergo repeat mpMRI examination with endorectal coil, out of which 40 patients agreed. The repeat study was performed in 15 of those who agreed. Sequence acquisition parameters are summarized in Table 1, and clinical indications for the imaging examination are listed in Table 2. The remaining 25 of 40 subjects did not complete the repeat study due to conflicts in patient’s schedule or withdrawal from the study (see the flow diagram in Fig. 1). The mean ± SD (range) age of the 15 subjects enrolled in the study was 61 ± 7 (47–69) years. Mean interval between the 2 mpMRIs was 10 (3–14) days. The mean ± SD (range) serum PSA in 14 of 15 subjects was 6.4 ± 2.2 (3.15–9.8) ng/mL. Prostate-specific antigen was not available in 1 subject, who was referred for imaging from an outside practice. Twelve of 15 patients had TRUS-guided sextant biopsy, of which 8 had pathology-confirmed PCa (3 + 3 Gleason score [GS]: 4; 3 + 4 GS: 3; 4 + 5 GS: 1), and 4 did not have confirmed PCa. Three of the patients did not undergo prostate biopsy before imaging, and thus no pathological grading was available. None of the patients had prostate biopsy performed between the baseline and repeat imaging examinations. Based on our experience, the population reported is typical of the prostate MR referrals for our institution. It consists primarily of the patients with confirmed PCa for staging or assessment of changes, or referred for imaging assessment due to elevated PSA in absence of histological confirmation of the disease, as shown in Table 2.

PI-RADS v2 Assessment
Tumor-suspicious tROIs were localized and contoured in 11 of 15 subjects (PI-RADS v2 > 1 in all 11/15 cases). All focal tROIs were located in the PZ. In 2 subjects, a secondary lesion was identified in the central zone of the gland, which was excluded from the subsequent quantitative analysis. No suspected lesion was identified in 4 of 15 subjects. In the subjects where a tROI was identified, in all cases, tROI was

TABLE 2. Patient-Level Summary of the Clinical Indications for MRI, Histopathology, PSA, and PI-RADS v2 Assessment of the Disease in the Evaluated Cohort

<table>
<thead>
<tr>
<th>Subject</th>
<th>Indication for the MRI Examination</th>
<th>Maximum Gleason Grade at Biopsy</th>
<th>PSA, ng/mL</th>
<th>PI-RADSv2 Assessment Category, Baseline Study</th>
<th>PI-RADSv2 Assessment Category, Repeat Study</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Known PCa, staging</td>
<td>3 + 4</td>
<td>5.4</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>2</td>
<td>Known PCa, assess change</td>
<td>3 + 4</td>
<td>7.5</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>3</td>
<td>Known PCa, staging</td>
<td>3 + 3</td>
<td>8.2</td>
<td>2</td>
<td>4</td>
</tr>
<tr>
<td>4</td>
<td>Known PCa, staging</td>
<td>3 + 3</td>
<td>4.3</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>5</td>
<td>Suspected PCa, staging</td>
<td>NA*</td>
<td>NA*</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>6</td>
<td>Elevated PSA, staging</td>
<td>Benign</td>
<td>5</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>7</td>
<td>Elevated PSA, staging</td>
<td>4 + 5</td>
<td>6.2</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>8</td>
<td>Known PCa, assess change</td>
<td>3 + 3</td>
<td>4.8</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>9</td>
<td>Elevated PSA, staging</td>
<td>Benign</td>
<td>9.4</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>10</td>
<td>Known PCa, assess change</td>
<td>3 + 3</td>
<td>3.15</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>11</td>
<td>Known PCa, assess change</td>
<td>3 + 3</td>
<td>9.7</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>12</td>
<td>Elevated PSA, staging</td>
<td>Benign</td>
<td>5.5</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>13</td>
<td>Known PCa, staging</td>
<td>3 + 4</td>
<td>4.16</td>
<td>4</td>
<td>3</td>
</tr>
<tr>
<td>14</td>
<td>Elevated PSA, staging</td>
<td>No biopsy performed</td>
<td>7</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>15</td>
<td>Repeat negative systematic TRUS biopsy, rising PSA</td>
<td>Benign</td>
<td>9.8</td>
<td>1</td>
<td>2</td>
</tr>
</tbody>
</table>

*The subject was referred for imaging from an outside practice, and their Gleason and PSA were not present in our institution electronic health records system.

MRI indicates magnetic resonance imaging; PSA, prostate-specific antigen; PI-RADS, prostate imaging reporting and data system; PCa, prostate cancer; TRUS, transrectal ultrasound.
localized in both baseline and repeat studies. There was agreement in the location of the tROI for all 11 cases. The images and annotations of the tROI for 2 of the subjects are shown in Figures 2 and 3.

Agreement in the PI-RADS v2 category was observed in 11 of 15 subjects as follows (also summarized in Table 2): PI-RADS v2 = 4 in 7 of 15 patients whose pathology was GS 4 + 5 (n = 1), GS 3 + 4

FIGURE 2. Midgland level axial T2-weighted (panels A and D), SUB (B and E), and ADC (C and F) images for subject 8 from Table 2. Top row (panels A–C) shows baseline study images, bottom row (panels D–F) is the repeat study. A contour (white arrow) is placed around the tumor on each sequence. At the time of ROI placement for each sequence, previously annotated ROI contours for other sequences within the same study were not visible to the radiologist.

FIGURE 3. Midgland level axial T2-weighted (panels A and D), SUB (B and E), and ADC (C and F) images for subject 14 from Table 2. Top row (panels A–C) shows baseline study images, bottom row (panels D–F) is the repeat study. No lesion was identified for this subject.
FIGURE 4. Bland-Altman plots summarizing repeatability of the volumetric measurements for the regions of interest (WG = whole gland, PZ = peripheral zone, tROI = tumor region of interest) across the 3 modalities (T2AX = axial T2-weighted image, SUB = dynamic contrast-enhanced MRI subtract map, ADC = apparent diffusion coefficient map) considered in this study.
We observed a gradual increase of measurement variability, and increasingly wider confidence intervals, going from large (WG) to small observed for the tROI volumes.

Repeatability of the WG and PZ volumetry estimation was obtained using data from all 15 subjects, whereas lesion repeatability was evaluated in 11 patients with identifiable lesions.

T2AX indicates T2-weighted axial image; SUB, DCE MRI subtract image computed as the difference between the phase corresponding to the contrast bolus arrival and the baseline phase; ADC, apparent diffusion coefficient; SD, standard deviation; PZ, peripheral zone; nPZ, normal PZ; tROI, tumor region of interest; RC, repeatability coefficient; CI, confidence interval.

ROI Volume and ADC Quantitation

Tumor-suspicious ROI volume, averaged between the measurements obtained in the baseline and repeat T2w scans, was less than 0.5 mL in 8 of the 11 subjects that had an identifiable lesion. Volumes of the considered structures across the baseline/repeat scans, and subjects were as follows (mean ± SD [range]): WG (T2AX, 46.5 ± 28.1 [19.1–115.9] mL); SUB, 48.2 ± 27.9 [19.6–110.7] mL; ADC, 60.5 ± 36.4 [21.3–141.8] mL; PZ (T2AX, 9.9 ± 3.9 [3.2–18.6] mL; SUB, 9.8 ± 3.7 [2.5–17.5] mL; ADC, 13.6 ± 5.5 [3.2–25.8] mL), and tROI (T2AX, 0.3 ± 0.2 [0.02–0.8] mL; SUB, 0.4 ± 0.2 [0.05–0.9] mL; ADC, 0.3 ± 0.2 [0.02–1.1] mL).

Linear mixed-effects model testing showed that volumes of the WG and PZ obtained from ADC maps were significantly larger than the values using T2w scans and SUB maps (P < 0.00005). However, no statistically significant differences were observed for the tROI volumes.

We observed a gradual increase of measurement variability, and increasingly wider confidence intervals, going from large (WG) to small (tROI) regions of interest, for both ROI volumes and mean ADC values. Repeatability of the volume measurements are summarized in Figure 4 and Table 3. Unequal variance testing identified no significant difference in the standard deviation of the measurements for either WG or PZ volumes across different sequences (see Table 4), whereas within-subject SD of the tROI volume was significantly different across sequences: 138 mm³ in ADC, 224 mm³ in SUB, and 68 mm³ in T2AX, P < 0.0005. Repeatability for the mean ADC measurements is summarized in Figure 5 and Table 5. RC% of mean ADC was significantly lower than that of ADC volume in PZ (P = 0.03) and tROI (P = 0.049). Coverage probability (CP) curves, intraclass coefficient (ICC), and concordance correlation coefficient (CCC) measures were also calculated and are included in the Supplemental Material, Supplemental Digital Content 1, http://links.lww.com/RLI/A319.

DISCUSSION

In this study, we evaluated repeatability of e-coil prostate mpMRI at 3 T, in a mixed cohort of 15 consecutive, consenting treatment-naive patients, being evaluated for PCa.

We considered a variety of repeatability measures, including those recommended by the consensus guidelines developed by Quantitative Imaging Biomarker Alliance,23,25 along with their confidence intervals. RC quantifies the absolute repeatability of the measurements in the same units as the measurement itself. It is defined as a 95% upper confidence bound on the absolute difference between the 2 replicate measurements and is directly related to the limits of agreement.

Repeatability of the Volumetric Measures for the Segmented Structures on T2-Weighted Axial Images, and ADC and SUB Maps

<table>
<thead>
<tr>
<th>Structure/Modality</th>
<th>RC% (95% CI)</th>
<th>RC (95% CI), mm³</th>
<th>Mean Difference (% Mean Difference), mm³</th>
<th>SD of Difference (% SD of Difference), mm³</th>
</tr>
</thead>
<tbody>
<tr>
<td>WG ADC</td>
<td>20.1 (10.33–29.87)</td>
<td>12166 (6253–18078)</td>
<td>5067 (8.37%)</td>
<td>3711 (6.13%)</td>
</tr>
<tr>
<td>SUB</td>
<td>39.8 (20.81–58.79)</td>
<td>19109 (9992–28226)</td>
<td>7364 (15.34%)</td>
<td>6613 (13.77%)</td>
</tr>
<tr>
<td>T2AX</td>
<td>38.44 (19.8–57.08)</td>
<td>17869 (9205–26534)</td>
<td>5817 (12.51%)</td>
<td>7266 (15.63%)</td>
</tr>
<tr>
<td>PZ ADC</td>
<td>45.98 (26.54–65.42)</td>
<td>6270 (3619–8921)</td>
<td>2773 (20.33%)</td>
<td>1651 (12.11%)</td>
</tr>
<tr>
<td>SUB</td>
<td>74.98 (44.31–105.65)</td>
<td>7351 (4343–10358)</td>
<td>3049 (31.11%)</td>
<td>2260 (23.05%)</td>
</tr>
<tr>
<td>T2AX</td>
<td>42.26 (24.43–60.09)</td>
<td>4201 (2428–5973)</td>
<td>1444 (14.53%)</td>
<td>1639 (16.49%)</td>
</tr>
<tr>
<td>PZ tROI ADC</td>
<td>112.22 (44.73–179.72)</td>
<td>422 (168–676)</td>
<td>159 (42.48%)</td>
<td>151 (40.27%)</td>
</tr>
<tr>
<td>SUB</td>
<td>119.43 (52.34–186.51)</td>
<td>488 (213–762)</td>
<td>187 (42.89%)</td>
<td>171 (42.05%)</td>
</tr>
<tr>
<td>T2AX</td>
<td>70.57 (29.05–112.09)</td>
<td>233 (96–371)</td>
<td>84 (25.56%)</td>
<td>88 (26.6%)</td>
</tr>
</tbody>
</table>

ROI indicates region of interest; ADC, apparent diffusion coefficient; SD, standard deviation; PZ, peripheral zone; tROI, tumor region of interest; T2AX, T2-weighted axial image.

Comparison of the Within-Subject Standard Deviation in ROI Volume Measurements Across Modalities Estimated With Unequal Variance Analysis

<table>
<thead>
<tr>
<th>Structure</th>
<th>ADC Volume SD, mm³</th>
<th>Subtract Volume SD, mm³</th>
<th>T2AX Volume SD, mm³</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>WG</td>
<td>8458</td>
<td>9030</td>
<td>7722</td>
<td>0.81</td>
</tr>
<tr>
<td>PZ</td>
<td>2617</td>
<td>2882</td>
<td>2316</td>
<td>0.70</td>
</tr>
<tr>
<td>PZ tROI</td>
<td>138</td>
<td>224</td>
<td>68</td>
<td>0.0004</td>
</tr>
</tbody>
</table>

ROI indicates region of interest; ADC, apparent diffusion coefficient; SD, standard deviation; PZ, peripheral zone; tROI, tumor region of interest; T2AX, T2-weighted axial image.
Based on our review of the literature, repeatability evaluations are often limited to absolute and relative percent differences, and RC is often not reported. This motivated our inclusion of other, more commonly used measures, together with those recommended by the consensus guidelines.

Tumor-suspicious ROI volume seems to be most stable when measured on T2-weighted axial images (~26% as the difference relative to the mean and ~71% RC%). RC and RC% of the tROI from ADC and SUB were about twice as large. This ordering is not unexpected, considering T2AX images have higher resolution, do not suffer from the DWI distortions, and are not dependent on the choice of the contrast uptake phases and organ motion in DCE analysis. Mean ADC (~16% as the relative difference and ~42% RC% for the PZ tROI) is more stable than volumetric measurements.

Changes in prostate gland volume may be useful preoperatively while evaluating response to hormonal therapy and in determining response of benign prostatic hyperplasia to androgen deprivation. Gland volume is also required for evaluating changes in PSA density and is sometimes considered as a potential marker for evaluating disease progression in patients under AS. These uses justify evaluation of the WG measurements repeatability. The WG and PZ regions can also serve as a measure of reference in evaluating the impact of region size

FIGURE 5. Bland-Altman plots summarizing repeatability of the mean apparent diffusion coefficient measurements for the regions of interest (WG = whole gland, PZ = peripheral zone, tROI = tumor region of interest, nPZ = normal peripheral zone).
on repeatability: our results show that measurements become less repeatable for smaller regions, as can be seen from Table 3. In general, we expect that the volume measurement error will be proportional to the surface area of the ROI, which in turn will depend on the shape of the region, because the segmentation error is concerned with the localization of the ROI boundary. This explains the increasing RC% and decreasing absolute RC for the regions we evaluated, going from large size WG ROI to the smaller size tROIs. Given the small number of subjects with lesions (n = 11), we could not make any statistically justified statements about the dependence of RC% or RC on the lesion size for tROIs.

We report both the absolute and relative repeatability measures. This choice is motivated by the lack of consensus on what is the best approach for assessing change in PCa lesions; Moore et al. suggested that small absolute changes in volume may appear as large percentage changes for small lesions. This is particularly relevant in the present study, because most of the lesions evaluated were below the clinically significant volume threshold. It is important to emphasize that the threshold of 0.5 mL was established based on the volume of the pathology estimated from the surgical specimen. There is evidence of high discordance between the MRI-based and histopathology volumes, especially for small volume disease, which is prevalent in indolent PCa cases. We observed imaging-based average volume of 0.3 mL, stressing the need for better understanding of repeatability in a low disease burden follow-up setting.

Repeatability of quantitative mpMRI measurements in the prostate has been investigated in the past. Alonzi et al. studied repeatability of DCE-derived metrics at 1.5 T without the use of e-coil. They did not evaluate repeatability of the parametric maps derived from multiphase DCE MRI, because it is recognized that DCE MRI does not play a primary role in detection of PCA and differentiation of aggressive cancer, as supported by PI-RADS v2 criteria for detection of PCa lesions. We are not aware of prior studies evaluating the repeatability of mpMRI-based volumetric measurements. However, similar repeatability studies have been performed for PET/CT in advanced gastrointestinal malignancies, reporting RC of ~36%.

There have been prior studies investigating the repeatability of DWI-based measurements in the prostate. Gibbs et al. evaluated prostate ADC repeatability within 1 month of acquisition in healthy subjects (n = 8), reporting repeatability of 35% and Litjens et al. evaluated ADC measurement variability in a cohort of 10 subjects scanned at the interval of 6 to 12 months and observed average difference of ADC in PW at 68 ± 10^-6 mm^2/s (neither RC nor %RC was reported), as compared to 175 ± 10^-6 mm^2/s (see Table 5) observed in our cohort that underwent repeat scan within 2 weeks. Sadinski et al. investigated short-term repeatability of mean ROI ADC in 14 subjects with biopsy-proven PCA, with the same subject scanned twice without repositioning within the scanner bore. Their median tROI ADC variation was 36 ± 10^-6 mm^2/s (4.2%), lower than the 170 ± 10^-6 mm^2/s (15.9%) we report (see Table 5). This is not unexpected, considering there was no change in patient positioning and minimal time between the scans. Several recent studies led by Jambor et al investigated repeatability of DWI-derived parameters using multi-b acquisition with e-coil. Those studies evaluated short-term (10–15 minutes) repeatability but did not consider lesion volumetrics and were conducted under clinical trial conditions, which are typically different from clinical standard-of-care settings. Overall, we note that repeatability values we report are comparable with those for other abdominal organs. Our study has several limitations. First, our cohort is small, although we should recognize that we approached 189 patients to participate in the study. Only 40 patients agreed to the study, and a repeat MR was possible in 15 subjects, underscoring the challenges faced in MR repeatability studies enrollment, especially those that utilize e-coil. Of note, the size of our cohort exceeds that reported by Litjens et al. and Gibbs et al. Second, most of the lesions analyzed are below the pathology-based clinical significance volume threshold of 0.5 mL. However, we also note that there is no clear consensus on the imaging-based definition of the significant disease, and that quantification of changes is of particular importance in longitudinal follow-up of the low-volume disease in AS patients. Third, histological confirmation with prostatectomy could not be performed considering the disease characteristics of the enrolled patient population and the prospective nature of the study. We do not consider this a major limitation because the main goal was the evaluation of repeatability of the measurements derived from the ROIs consistently identified in mpMRI studies and not their pathological correlation.

In conclusion, we report repeatability measures for standard-of-care single-center e-coil prostate mpMRI at 3 T. Our results indicate that PI-RADS v2 category assignment is highly repeatable. Volumetric measurement of changes in tROI may be considered significant at the 95% confidence level if they exceed 70% (233 mm3) on T2AX, or 112% to 119% (422–488 mm3) in subtract and ADC imaging. Peripher-eral zone tROI mean ADC is more stable than ADC tROI volume, and its changes may be considered significant if they exceed 40% (447 ± 10^-6 mm^2/s). As with any small study, these results should be interpreted cautiously. Further investigation of mpMRI repeatability is warranted.

**REFERENCES**


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**TABLE 5. Repeatability of the Mean ADC Measurements for the Segmented Structures**

<table>
<thead>
<tr>
<th>Structure</th>
<th>RC% (95% CI)</th>
<th>RC (95% CI), ×10^-6 mm^2/s</th>
<th>Mean Difference (% Mean Difference), ×10^-6 mm^2/s</th>
<th>SD of Difference (% SD of Difference), ×10^-6 mm^2/s</th>
</tr>
</thead>
<tbody>
<tr>
<td>WG</td>
<td>29.53 (18.23–40.83)</td>
<td>359 (221–497)</td>
<td>83 (6.85%)</td>
<td>169 (13.89%)</td>
</tr>
<tr>
<td>PW</td>
<td>22.31 (13.91–30.71)</td>
<td>305 (190–419)</td>
<td>88 (6.45%)</td>
<td>132 (9.71%)</td>
</tr>
<tr>
<td>nPW</td>
<td>30.24 (18.86–41.61)</td>
<td>471 (294–649)</td>
<td>175 (11.27%)</td>
<td>170 (10.91%)</td>
</tr>
<tr>
<td>tROI</td>
<td>41.83 (23.12–60.49)</td>
<td>447 (247–647)</td>
<td>170 (15.93%)</td>
<td>159 (14.88%)</td>
</tr>
</tbody>
</table>

ADC indicates apparent diffusion coefficient; SD, standard deviation; PW, peripheral zone; nPW, normal PW; tROI, tumor region of interest; T2AX, T2-weighted axial image; RC, repeatability coefficient; CI, confidence interval.


