Retrospective study comparing MR-guided radiation therapy (MRgRT) setup strategies for prostate treatment: repositioning vs. replanning

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Abstract
Background: This study compared adaptive replanning and repositioning corrections based on soft-tissue matching for prostate cancer by using the magnetic resonance-guided radiation therapy (MRgRT) system.

Methods: A total of 19 patients with prostate cancer were selected retrospectively. Weekly magnetic resonance image (MRI) scans were acquired for 5 weeks for each patient to observe the anatomic changes during the treatment course. Initial intensity-modulated radiation therapy (IMRT) plans (iIMRT) were generated for each patient with 13 coplanar ⁶⁰Co beams on a ViewRay™ system. Two techniques were applied: patient repositioning and replanning. For patient repositioning, one plan was created: soft-tissue (prostate) matching (Soft). The dose distribution was calculated for each MRI with the beam delivery parameters from the initial IMRT plan. The replanning technique was used to generate the Adaptive plan, which was the reoptimized plan for the weekly MRI. The dose-volumetric parameters of the planning target volume (PTV), bladder, and rectum were calculated for all plans. During the treatment course, the PTV, bladder, and rectum were evaluated for changes in volume and the effect on dosimetric parameters. The differences between the dose-volumetric parameters of the plans were examined through the Wilcoxon test. The initial plan was used as a baseline to compare the differences.

Results: The Adaptive plan showed better target coverage during the treatment period, but the change was not significant in the Soft plan. There were significant differences in D₉₈%, D₅₉%, and D₂% in PTV between the Soft and Adaptive plans (p < 0.05) except for Dmean. There was no significant change in Dmax and Dmean as the treatment progressed with all plans. All indices for the Adaptive plan stayed the same compared to those of iIMRT during the treatment course. There were significant differences in D₅₅%, D₃₁%, D₂₁%, and D₁₅% in the bladder between the Soft and Adaptive plans. The Adaptive plan showed the worse dose sparing than the Soft plan for the bladder according to each dosimetric index. In contrast to the bladder, the Adaptive plan achieved better sparing than the Soft plan during the treatment course. The significant differences were only observed in D₁₅% and D₃₁% between the Soft and Adaptive plans (p < 0.05).

Conclusions: Patient repositioning based on the target volume (Soft plan) can relatively retain the target coverage for patients and the OARs remain at a clinically tolerance level during the treatment course. The Adaptive plan did not clinically improve for the dose delivered to OARs, it kept the dose delivered to the target volume constant. However, the Adaptive plan is beneficial when the organ positions and volumes change considerable during treatment.

Keywords: Soft-tissue repositioning, Adaptive replanning, Prostate radiotherapy, MRI-guided treatment
Background
The goal of radiation therapy is to deliver an accurate prescription dose to the target while minimizing the dose to normal tissue. External beam radiotherapy (EBRT) is a common definitive treatment option for localized prostate cancer [1–7]. In the last decade, significant technological advances in EBRT have been achieved. Intensity-modulated radiation therapy (IMRT) is a type of conformal radiation therapy and is widely used to treat prostate cancer [1, 7, 8]. However, the complexity of the treatment delivery and inter- and intra-fraction variations are a concern when using IMRT for prostate cancer [9, 10]. Adding margins around the clinical target volume (CTV) can account for these variations to ensure target coverage. Gill et al. reported that when daily cone beam computed tomography (CBCT) was used for soft-tissue alignment of the prostate, a PTV margin of 3 mm allowed sufficient coverage for CTV [11]. On the other hand, in Engels’s study, margins of 6 mm LR and 10 mm AP and CC were used in patients who did not have markers implanted in them. Margins of 3 mm LR and 5 mm AP and CC were used for patients with implanted markers [12]. However, expanding the irradiated volume consequently increases normal organ toxicity, which is limited by the tolerances of the bladder and rectum [8, 13]. In recent years, advances in image guidance have allowed for better localization of the prostate [14–20]. Using image guidance together with IMRT delivers the dose to the prostate more precisely with smaller margins to decrease the doses to the rectum and bladder. Image-guided radiation therapy (IGRT) technology has been widely adopted to provide anatomic information with the patient in the treatment position. With image guidance (typically X-ray imaging such as kilovolt (kV), megavolt (MV), and CBCT), the patient positions in the daily images are registered to the planning images based on either bony landmarks or soft tissues [16–18, 21, 22]. Furthermore, fiducial markers for prostate IGRT have been in use since the 1990s and O’Neill et al. reviewed the evidence for the use of fiducial markers in clinical practice [23]. Image guidance corrects not only for patient position variations but also for changes in the target volume and shapes. Adaptive radiation therapy (ART) has been introduced to compensate for variations in patient treatment during a radiotherapy course [24–26]. ART techniques can be categorized as offline or online adaptive planning [27]. Offline adaptive planning uses a feedback strategy for current treatment by incorporating the obtained daily treatment images. This adaptive planning reflects patient-specific anatomic variations and provides a reoptimized treatment plan for the remaining treatment fractions. Online adaptive planning is the optimal strategy for each fraction delivery because the treatment plan is reoptimized while the patient is waiting on the treatment table. However, clinical implementation of both offline and online adaptive techniques with a common imaging modality based on X-rays is still in the development phase because it is not sufficient to modify the treatment plan according to anatomic changes on inter- and intra-fraction bases. In 2009, Thongphiew et al. compared three online IGRT techniques (bony-anatomy matching, soft-tissue matching, and online replanning) for prostate IMRT treatment using daily CBCT [18]. They demonstrated that CBCT can feasibly be used for reoptimizing the treatment plan online, which has significant benefits when a high degree of deformation or differential organ position displacement occurs. Clinical implementation of ART for prostate cancer has been greatly promoted by image guidance with onboard CBCT [28–31]. Onboard CBCT allows the target volume and adjacent anatomies to be localized before treatment of a patient in the treatment position. However, CBCT has insufficient image quality compared to planning computed tomography (CT) to provide an accurate delineation of structures and dose calculation owing to the higher noise and lower contrast [29, 32]. Furthermore, the inconsistency between the CT number to electron density curves of CT and CBCT images can result in a discrepancy in the dose calculations.

Methods
Patient data and study design
After institutional review board approval, a total of 19 patients with prostate cancer were selected for this study in a retrospective manner. Low-risk patients did not receive androgen deprivation therapy (ADT). Intermediate and high risk patients received neoadjuvant ADT. Table 1 summarizes the patients’ clinical characteristics. All patients were planned with IMRT using ViewRay. A CT image set and MRI image set of each patient in the supine position were acquired once before treatment. The Brilliance Big Bore™ CT simulator (Philips, Cleveland, OH, USA) was used for CT scanning with a slice thickness of 1.5 mm. Each patient was immobilized with Smart Rest™ (Chunsung, Seoul, Republic of Korea), which combines...
the knee fix and feet fix. The initial CT image (iCT) sets were only used to calculate the dose for the MRIdian™ system (ViewRay Inc., Cleveland, OH, USA); no CT images were taken during treatment. Initial MRI (iMRI) was used to draw the contours and the reference image. All patients underwent daily MRI scans for setup verification prior to treatment during the treatment period. Of these, only weekly MRI images were included in the study. All MRIs (six MRI sets per patient) were acquired on a 0.35 T ViewRay scanner system (ViewRay Inc., Cleveland, OH, USA). A non-contrast true fast imaging with steady-state precession (TRUFI) sequence was used for all MRI scanning [41]. The MRI resolution was 1.5×1.5×3.0 mm³ with a typical imaging time of 25 s and a field of view of 54×47×43 cm³. The CTV was defined to include prostate and seminal vesicles. The planning target volume (PTV) was generated by adding margins of 3 mm and 5 mm in all directions to the prostate and seminal vesicles, respectively. The rectum and bladder were not controlled for filling. These structures (six RT structure sets per patient) were delineated on each MRI by a physician to ensure that the contouring practice was consistent. The prescription dose to the PTV was 70 Gy in 28 fractions for all patients. Two strategies for MRgRT plans were considered. One plan used the repositioning technique, which was called as the Soft plan. The repositioning technique finds the best matches of the target volume between the iMRI and weekly MRI (nMRI). Patient repositioning was performed by translational correction because of the motion limitations of the ViewRay couch. The other plan used the replanning technique for every nMRI, which was called as the Adaptive plan. Thus, each patient had one initial IMRT plan (iIMRT), five Soft plans, and five Adaptive plans.

Techniques and treatment plans
The iCT was deformed to iMRI to generate the deformed CT (dCT) for dose calculation. The initial structure set (iRS) contoured on iMRI was also included in the treatment planning. A total of 13 fields (five beam groups) were used to generate an iIMRT plan for each patient with the tri-60Co system. The gantry angles of the fields were 0° (group 1); 24°, 144°, and 264° (group 2); 48°, 168°, and 267° (group 3); 72°, 192°, and 312° (group 4); and 96°, 216°, and 336° (group 5). The values for the IMRT efficiency that smoothed the fluence map intensity pattern (could be set from 0 to 1) and maximized the deliverable beam-on segments per field were 0.5 and 10, respectively. The optimization and dose calculation were performed with an imaging surface coil in the presence of a magnetic field and using a calculation grid of 3 mm. The Monte Carlo calculation algorithm developed by the manufacturer (ViewRay Inc., Cleveland, OH) was used to calculate the dose. The number of histories in the system was set to $2.4 \times 10^6$. This setting achieves about 1% statistical uncertainty for a dose grid of 3 mm cubed voxels. The optimization was performed according to National Comprehensive Cancer Network (NCCN) guidelines, and each plan was normalized to cover 100% of the PTV with 95% of the prescription dose. The MRIdian™ system use a convex nonlinear programming model for dose optimization. Table 2 shows the dose optimization parameters for the target and OARs. For a relatively fair comparison, all plans used these optimization parameters. Finally, the iIMRT plan was determined according to several beam delivery parameters. For each patient, six sets of MRIs (nMRI) and structure sets (nRS) were scanned during

| Table 1 | Summary of patients’ clinical characteristics |
|---------|---------------------------------------------|
| Age (years) | Mean (range) | 77 (65–86) |
| T stage | | |
| T1 | 2 (10.5%) |
| T2 | 11 (57.9%) |
| T3 | 2 (10.5) |
| T4 | 4 (21.1%) |
| GS score | | |
| 6 (3 + 3) | 3 (15.8%) |
| 7 (3 + 4) | 3 (15.8%) |
| 7 (4 + 3) | 5 (26.3%) |
| 8 (4 + 4) | 7 (36.8%) |
| 9 (4 + 5) | 1 (5.3%) |
| PSA | Median (range) | 18.89 (2.19–161.95) |
| | Mean (SD) | 27.78 (35.63) |
| ADT use | 10 (52.63%) |

*PSA prostate-specific antigen; ^ADT androgen deprivation therapy

| Table 2 | Summary of MRIdian™ optimization parameters for target volume and OARs |
|---------|------------------------------------------|
| Structure | Importance | Power | Threshold | Prescription |
|          | Lower | Upper | Lower | Upper | (Gy) | (Gy) | Offset |
| Skin | – | 1 | – | 1 | 21 | – | – |
| Rectum | – | 1 | – | 1 | 56 | – | – |
| Bladder | – | 1 | – | 1 | 56 | – | – |
| aPTV | 2 | 2 | 2 | 2 | – | 70 | 0.5 |

*PTV planning target volume
the treatment course and used in this study. For the Soft plan, the isocenter from iIMRT was relocated to the center of the PTV from nRS with nMRI. After the patient was repositioned based on target volume, the dose (from the iIMRT) was recalculated on nMRI with nRS. In contrast, the Adaptive plan was generated from the Soft plan with reoptimization by using the same constraints and associated weights for iIMRT. The dCT was generated by deforming dCT to nMRI and used to calculate the dose for each plan. Each plan was compared at a prescription dose of 70 Gy.

Dose-volumetric evaluation and comparison

The dose-volumetric parameters of the PTV, bladder, and rectum were calculated for all plans. The dosimetric index \( D_{\text{vol}} \) is defined as the dose to \( x\% \) of volume of a structure. For the PTV, the mean dose \( (D_{\text{mean}}) \), \( D_{98\%} \), \( D_{95\%} \), and \( D_{2\%} \) were calculated. The conformity index (CI), homogeneity index (HI), and gradient index (GI) were calculated as follows [42, 43]:

\[
\text{Conformity index (CI)} = \frac{\text{Volume of reference isodose}}{\text{Volume of target volume}}
\]

\[
\text{Homogeneity index (HI)} = \frac{D_{98\%} - D_{95\%}}{D_{50\%}}
\]

\[
\text{Gradient index (GI)} = \frac{V_{50\%} \text{ of the prescription dose}}{V_{100\%} \text{ of the prescription dose}}
\]

For the bladder and rectum, \( D_{15\%} \), \( D_{25\%} \), \( D_{35\%} \), and \( D_{50\%} \) were calculated. During the treatment course, the PTV, bladder, and rectum were evaluated for changes in volume and their effect on the dosimetric parameters. Each technique was compared according to the aforementioned dose-volumetric parameters. Wilcoxon tests were performed at the 95% confidence level to examine the statistical significance of differences between the dose-volumetric parameters of the plans. The iIMRT (i.e., IMRT plan for the first fraction) was used as a criterion for comparing the differences. The given plan was defined as “reproducible” for both PTV and OARs, respectively, if the difference was less than 5% compared to the iIMRT.

**Results**

**Dose-volumetric evaluation and comparison of PTVs**

Table 3 lists the results of dose-volumetric evaluation and a comparison of the PTVs for each plan. There were significant differences in \( D_{98\%} \), \( D_{95\%} \), and \( D_{2\%} \) in PTV between the Soft and Adaptive plans \((p < 0.05)\), except for \( D_{\text{mean}} \). For each dosimetric index, one plan group including the iIMRT exhibited an average value relative to the prescription dose and its standard deviation \((\text{Mean} \pm \text{SD})\). With regard to the index related to target coverage, the Adaptive plan clearly demonstrated superior results for \( D_{98\%} \) (99.1 ± 0.3%). For \( D_{95\%} \), the average value for Adaptive plan was always 100% because each plan was normalized to cover 100% of the PTV with 95% of the prescription dose. The difference was statistically significant compared with that of the Soft plan \((p = 0.011)\). Compared with iIMRT, the average difference for the Adaptive plan was less than 1% for all dose-volumetric evaluations. The percentage of reproducible plans for \( D_{98\%} \) was 77.9% with the Soft plan.

Figure 1 shows the averaged dose-volumetric values of the PTV for each plan and all patients during treatment. The target coverage during the treatment course did change significantly with the Soft plan. There was no significant change in \( D_{\text{max}} \) and \( D_{\text{mean}} \) as the treatment progressed with all plans. Figure 2 shows the averaged CI, HI, and GI of the PTV for all plans during treatment. The average CI values during the treatment course were 1.38 ± 0.27 with the Soft plan, and significant changes were observed during the treatment course. In this analysis, the iIMRT was excluded because this was the reference plan. The CI value was lower in the Adaptive plan but not statistically significant compared to that in the Soft plans \((p = 0.08)\). However, HI and GI values were also lower in the Adaptive plan and statistically significant, compared to the Soft plans \((p < 0.05)\). The averaged HI and GI values for the Soft plan compared to those of

| Indices | Plan | Mean ± SD (%) | \( p \)-value | % of reproducible plan | Average difference (%) |
|---------|------|--------------|---------------|------------------------|------------------------|
| \( D_{\text{mean}} \) | Soft | 102.5 ± 0.6 | 0.099 | 100 | 0.1 ± 0.9 |
| | Adaptive | 102.2 ± 0.3 | 0.011 | 100 | −0.1 ± 0.5 |
| \( D_{98\%} \) | Soft | 96.6 ± 3.3 | 0.005* | 77.9 | −2.8 ± 4.1 |
| | Adaptive | 99.1 ± 0.3 | 0.011* | 100 | −0.2 ± 0.6 |
| \( D_{95\%} \) | Soft | 98.5 ± 2.6 | 0.011* | 92.6 | −1.8 ± 3.2 |
| | Adaptive | 100 | 0 | 100 | 0 |
| \( D_{2\%} \) | Soft | 105.5 ± 1.0 | <0.001* | 100 | 0.7 ± 1.0 |
| | Adaptive | 104.6 ± 0.4 | 0.011* | 100 | −0.4 ± 1.0 |

*Statistically significant comparisons; † Percentage of plans with a difference of < 5% compared to the initial plan
iIMRT were 0.09 ± 0.05 and 1.11 ± 0.18, respectively. All indices for the Adaptive plan stayed the same compared to those of iIMRT during the treatment course; the average values of CI, HI, and GI were 1.28 ± 0.22, 0.06 ± 0.01, and 1.04 ± 0.11, respectively.

Dose-volumetric evaluation and comparison for the bladder
Figure 3 shows the averaged dose-volumetric values of the bladder for each technique and all patients during treatment. With the Soft and Adaptive plans, the average $D_{15\%}$ values excluding that of iIMRT during the treatment
course were 53.3 ± 7.6 and 55.5 ± 7.1 Gy, respectively. The average $D_{20\%}$ values were 44.8 ± 9.2 and 47.4 ± 9.0 Gy, respectively. The average $D_{35\%}$ values were 37.9 ± 10.1 and 40.7 ± 10.2 Gy, respectively. The average $D_{50\%}$ values were 29.7 ± 10.3 and 32.5 ± 10.4 Gy, respectively. The Adaptive plan did not achieve a better result than the Soft plan during the treatment course. However, all plans had similar values for the standard deviation and exhibited a tolerable dose distribution of the bladder. Table 4 lists the results of the dose-volumetric evaluation and compares the plans for the bladder. There were significant differences in $D_{15\%}$, $D_{25\%}$, $D_{35\%}$, and $D_{50\%}$ in the bladder between the Soft and the Adaptive plans. For each dosimetric index, one plan group including the iIMRT showed the average value relative to the prescription dose and its standard deviation (Mean ± SD). The Adaptive plan showed the worse dose sparing compared with the Soft plan for bladder according to each dosimetric index. The difference was statistically significant compared to that for the Soft plan ($p < 0.05$). Even though the Adaptive plan showed a higher dose in the bladder than the Soft plan, the average percentage of differences compared to iIMRT showed lower values than those of the Soft plan. However, the percentages of reproducible plans for $D_{15\%}$, $D_{25\%}$, and $D_{35\%}$ were 35.8, 28.4, and 25.3%, respectively, which were higher values than those of the Soft plan except for $D_{50\%}$.

### Dose-volumetric evaluation and comparison for the rectum

Figure 4 shows the averaged dose-volumetric values of the rectum for each technique and all patients during treatment. With the Soft and Adaptive plans, the average $D_{15\%}$ values excluding that of iIMRT during the treatment course were 62.4 ± 3.5 and 61.3 ± 1.9 Gy. The average $D_{25\%}$ values were 57.7 ± 3.6 and 56.5 ± 2.2 Gy; the average $D_{35\%}$ values were 52.8 ± 3.3 and 51.6 ± 2.0 Gy; the average $D_{50\%}$ values were 48.6 ± 2.9 and 47.3 ± 2.2 Gy; the average $D_{70\%}$ values were 44.1 ± 2.4 and 43.0 ± 2.1 Gy; the average $D_{85\%}$ values were 39.9 ± 2.2 and 38.5 ± 2.0 Gy.

### Table 4 Dose-volumetric evaluation and comparison of plans for the bladder during the treatment course

| Indices | Plan      | Mean ± SD (%) | p-value | % of reproducible plan* | Average difference (%) |
|---------|-----------|---------------|---------|------------------------|------------------------|
| $D_{15\%}$ | Soft     | 77.6 ± 7.8    | 0.01*   | 33.7                   | −9.14 ± 15.1           |
|         | Adaptive  | 80.2 ± 8.2    |         |                        | −5.4 ± 14.8            |
| $D_{25\%}$ | Soft     | 65.9 ± 9.8    | 0.01*   | 25.3                   | −12.5 ± 23.5           |
|         | Adaptive  | 69.0 ± 10.6   |         |                        | −7.2 ± 24.9            |
| $D_{35\%}$ | Soft     | 56.3 ± 11.2   | 0.016*  | 23.2                   | −14.6 ± 33.5           |
|         | Adaptive  | 59.6 ± 12.2   |         |                        | −8.0 ± 37.4            |
| $D_{50\%}$ | Soft     | 44.8 ± 11.5   | 0.02*   | 23.2                   | −16.4 ± 48.5           |
|         | Adaptive  | 48.1 ± 12.7   |         |                        | −8.0 ± 56.3            |

*Statistically significant comparisons; *Percentage of plans with a difference of < 5% compared to the initial plan.
D35% values were 53.2 ± 4.5 and 52.0 ± 5.4 Gy; and the average D50% values were 46.7 ± 4.8 and 45.6 ± 2.6 Gy. All plans were below the tolerance dose of rectum. In contrast to the bladder, the Adaptive plan achieved better sparing than that by the Soft plan during the treatment course and showed the smallest standard deviation. Table 5 lists the results of the dose-volumetric evaluation and compares the plans for the rectum. The differences were only observed in D15% and D35% between the Soft and Adaptive plans, statistically (p < 0.05). For each dosimetric index, one plan group including iIMRT showed the average value relative to the prescription dose and its standard deviation (Mean ± SD). The Adaptive plan shows less average value and lower standard deviation for the rectum than the Soft plan according to each dosimetric index. However, no differences were observed between the plans in D25% and D50% (p > 0.05). With the Soft plan, the percentages of reproducible plans for D15%, D25%, D35% and D50% were 87.4, 87.4, 87.4, and 86.3%, respectively. The average percentages of difference compared to iIMRT had positive mean values and large standard deviations: 1.1 ± 6.4%, 1.1 ± 7.7%, 1.2 ± 8.8%, and 1.2 ± 9.9% for D15%, D25%, D35% and D50%, respectively. With the Adaptive plan, on the other hand, the percentages of reproducible plans for D15%, D25%, D35% and D50% were 89.5, 86.3, 81.1, and 76.8%, respectively. The average percentages of difference compared to iIMRT had negative mean values and small standard deviations: −0.8 ± 3.0%, −1.0 ± 3.7%, −1.0 ± 4.4%, and −1.1 ± 5.8% for D15%, D25%, D35% and D50%, respectively.

Table 5 Dose-volumetric evaluation and comparison of plans for the rectum during the treatment course

| Indices | Plan      | Mean ± SD (%) | p-value | % of reproducible plan | Average difference (%) |
|---------|-----------|---------------|---------|------------------------|------------------------|
| D15%    | Soft      | 62.3 ± 2.8    | 0.009*  | 87.4                   | 1.1 ± 6.4              |
|         | Adaptive  | 61.3 ± 1.5    |         | 89.5                   |                        |
| D25%    | Soft      | 57.5 ± 3.2    | 0.16    | 87.4                   | 1.1 ± 7.7              |
|         | Adaptive  | 56.5 ± 1.8    |         | 86.3                   | −1.0 ± 3.7             |
| D35%    | Soft      | 75.7 ± 5.1    | 0.033*  | 87.4                   | 1.2 ± 8.8              |
|         | Adaptive  | 74.4 ± 2.7    |         | 81.1                   | −1.0 ± 4.4             |
| D50%    | Soft      | 66.5 ± 5.6    | 0.059   | 86.3                   | 1.2 ± 9.9              |
|         | Adaptive  | 65.2 ± 2.9    |         | 76.8                   | −1.1 ± 5.8             |

*Statistically significant comparisons; % Percentage of plans with a difference of < 5% compared to the initial plan

![Fig. 4 Averaged dose-volumetric values of (a) D15%, (b) D25%, (c) D35%, and (d) D50% for the rectum during the treatment course](image-url)
Discussion
This study explored different MRgRT techniques to compensate for inter-fraction errors during prostate treatment. The Soft plan, which repositions the patient by using the target volume, may be sufficient for correcting anatomic variations. For OARs with the Soft plan, the dose to the bladder was relatively low, while the dose to the rectum was relatively consistent during treatment. The Adaptive plan, which is a replanning technique, may be suitable for correcting anatomic variations, and the doses to the OARs remained relatively consistent during treatment. The Adaptive plan performed very similarly to the initial plan and compensated for inter-fraction uncertainty. The large variations in the mean values and standard deviation indicate that substantial fluctuations in target coverage can occur weekly with IGRT techniques based on patient repositioning. Only the replanning technique (i.e., Adaptive plan) maintained a consistent quality during the treatment period in terms of target coverage. However, the Adaptive plan did not achieve the better results in terms of sparing the bladder during the treatment course. In this study, the enrolled patients were not controlled for bladder and rectum filling because they had been treated with gated radiotherapy using the ViewRay system. Thus, the bladder and rectum were filled with various volumes during the treatment course. Figure 5 shows the averaged volume variation for all patients during the treatment course. The changes in the bladder volume were significant with large standard deviations. Significant variations in the bladder volume affect both the bladder dose volume and positions of adjacent organs [44–46]. The Adaptive plan did achieve the better results for rectum sparing because the variation in the rectum volume was not significant. Figure 6 compares the example plans. In this case, the Soft plan had better OAR sparing than the Adaptive plan while missing target coverage. Thongphiew et al. used daily CBCT and also reported that the adaptive technique does not achieve the better OAR sparing for all fractions [18]. They discussed the limitations of CBCT. Compared to CT, CBCT images make it more difficult to distinguish the boundaries of soft-tissue organs (e.g., the prostate, bladder, and rectum) because of more noise and poorer soft-tissue contrast. In this study, however, a 0.35 T MRI was used with high soft-tissue contrast and real-time imaging capability to exactly define the soft-tissue organ variations. Even low-field-strength MRI gives improved anatomic visualization compared with CT and CBCT [47]. The MRI allowed for the most accurate delineation of soft tissues and observation of organs changing. Thus, we adopted PTV margins of 3 mm and 5 mm in all directions for the prostate and seminal vesicles, respectively. McPartlin et al. reported that the CTV- to-PTV margin can be safely reduced to less than 3 mm in MRI-guided prostate adaptive radiotherapy.
because of high soft-tissue contrast and real-time imaging capability [48]. On the other hand, the plan quality was not better than that of a treatment-based linear accelerator (linac) because of the large penumbra of the $^{60}$Co source and a large width of the multileaf collimator (MLC). The patient positioning techniques were also limited because only translational corrections were available. An MR-linac with 6 MV beam is available in a clinic, and this system provides a better plan quality than a system equipped with $^{60}$Co beams because of small penumbra and a small width of MLC [49, 50]. In addition, adaptive planning will increase the time between the imaging and the beam delivery. The intra-fractional motion and deformation also increase with time [51]. These could be handled by using the gating delivery technique in treatment delivery. Furthermore, the Adaptive plan is not the actual dose delivered and the actual delivered dose should be assessed in parallel after treatment.

Conclusions
In this study, the MRgRT technique using adaptive planning was compared to the technique based on patient repositioning for 19 prostate cancer patient with the MRI. Patient repositioning based on the target volume (Soft plan) can relatively retain the target coverage for patients and the OARs remain at a clinically tolerance level during the treatment course. Although the Adaptive plan did not clinically improve for the dose delivered to OARs, it kept the dose delivered to the target volume constant. However, the Adaptive plan is beneficial when organ positions and volumes change considerably during treatment.

Abbreviations
ADT: Androgen deprivation therapy; ART: Adaptive radiotherapy; CBCT: Cone-beam computed tomography; CI: Conformity index; CT: Computed tomography; CTV: Clinical target volume; dCT: Deformed CT; EBRT: External beam radiotherapy; GI: Gradient index; HI: Homogeneity index; ICT: Initial CT image; IGRT: Image-guided radiotherapy; iIMRT: Intensity-modulated radiation therapy plans; IMR: Initial MRI; IMRT: Intensity-modulated radiation therapy; IRS: Initial structure set; kV: Kilovolt; Linac: Linear accelerator; MLC: Multileaf collimator; MRgRT: Magnetic resonance-guided radiation therapy; MRI: Magnetic resonance imaging; MV: Megavolt; NCCN: National comprehensive cancer network; nMRI: Weekly MRI; nRS: Weekly structure set; OARs: Organs at risk; PTV: Planning target volume; TRUFI: True fast imaging tomography

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Authors’ contributions
JHK and JMP conceived the study concept, compiled and analyzed the data, drafted the manuscript, and participated in all aspects of the study. CHC and HJA helped measure the data. YJK and JHK analyzed the data from a clinical perspective and engaged in discussions to improve the significance of this study. JHK oversaw and verified the study completion. All authors read and approved the final manuscript.

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Availability of data and materials
Data sharing not applicable to this article because no datasets were generated or analyzed during the current study.

Ethics approval and consent to participate
An approval for this study was obtained from the institutional review board (IRB No. 1607–047–774).

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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