Dosimetric study of RapidArc plans and conventional intensity modulated radiotherapy for prostate cancer involving seminal vesicles and pelvis lymph nodes

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Abstract

Purpose: The main purpose of this study is to (1) identify the continual diversity between conventional fixed field intensity modulation radiotherapy (IMRT) and RapidArc (RA) for high-risk prostate cancer; and (2) determine potential benefits and drawbacks of using for this type of treatment. Methods: A cohort of 20 prostate cases including prostate, seminal vesicles and pelvic lymph nodes was selected for this study. The primary planning target volume (PTV1) and boost planning target volume (PTV2) were contoured. The total prescription dose was 75.6 Gy (45 Gy to PTV1 and an additional 21.6 Gy to PTV2). Two plans were generated for each PTV: multiple 7-fields for IMRT and two arcs for RA. Results: A Sigma index (IMRT: 2.75 ± 0.581; RA: 2.8 ± 0.738) for PTV1 and (IMRT: 2.0 ± 0.484; RA: 2.1 ± 0.464) for PTV2 indicated similar dose homogeneity inside the PTV. Conformity index (IMRT: 0.96 ± 0.047; RA: 0.95 ± 0.059) for PTV1 and (IMRT: 0.97 ± 0.015; RA: 0.96 ± 0.014) for PTV2 was comparable for both the techniques. IMRT offered lower mean dose to organ at risks (OARs) compared to RA plans. Normal tissue integral dose in IMRT plan resulted 0.87% lower than RA plans. All the plans displayed significant increase (2.50 times for PTV1 and 1.72 for PTV2) in the average number of necessary monitor units (MUs) with IMRT beam. Treatment delivery time of RA was 2 – 6 minutes shorter than IMRT treatment. Conclusion: For PTV including pelvic lymph nodes, seminal vesicles and prostate, IMRT offered a greater degree of OARs sparing. For PTV including seminal vesicles and prostate, RA with two arcs provided comparable plan with IMRT. RA also improved the treatment efficiency due to smaller number of MUs required.

Keywords: IMRT, RapidArc, Sigma-Index, Conformity Index, Normal Tissue Integral Dose

1. Introduction

Cancer is the major cause of leading deaths in the 21st century in world with 14.1 million cases and 8.2 million deaths in 2012. Among them, prostate cancer stands as important due to risk of secondary malignancies associated with intensity modulation radiation therapy (IMRT) with conventional 3-dimensional conformal radiotherapy (3D-CRT). Among the different technologies adopted to cure the prostate cancer, external radiotherapy is recognized as one of the important treatment options. The technology aims to destroy cancer cells by minimal damaging (due to risk of secondary malignancies) to the surrounding normal tissues. It creates the best possible balance between maximizing dose to prostate cancer cells and minimization of side effects. IMRT technology for prostate tumor/cancer allows less toxicity in comparison to 3D-CRT. The development of IMRT technique has enabled the delivery of highly conformal
dose distribution to the target while limiting radiation damage to the critical organs within the tolerance limit. For the prostate cancer treatment, IMRT has been an ideal technique with the geometric relationship of the target volume to bladder, rectum and surrounding normal tissue.

RapidArc (Varian Medical Systems, Palo Alto, California, USA) is a volumetric radiotherapy technology that delivers a fast and precise sculpted 3D dose distribution with a single 360-degree rotation of the linear accelerator gantry to improve the standard of care and treatments. It is made possible by a treatment planning algorithm that simultaneously changes 3-parameters during treatment by tuning rotation speed of the gantry; shape of the treatment aperture using the movement of multileaf collimator leaves and delivery dose rate. Volumetric modulated arc therapy differs from existing techniques like helical IMRT or intensity-modulated arc therapy (IMAT) because it delivers dose to the whole volume, rather than segment by segment. The treatment-planning algorithm ensures the treatment precision and helping to spare normal healthy tissue.

Some of the studies were performed for the prostate cancer treatment; however, they included prostate only or prostate with seminal vesicles. The present study aims to expand such studies to identify the main and continual diversity between two techniques for prostate cancer cases that involve the seminal vesicles and pelvic lymph nodes. The dosimetric results and treatment delivery efficiency using the RapidArc technique were compared to those using the conventional static-gantry IMRT technique.

2. Methods and Materials

A cohort of 20 high-risk prostate cancer patients including prostate, seminal vesicles, and pelvic lymph nodes were selected for our study. Computed tomography (CT) images were acquired with an empty rectum and full bladder for all patients. The attending radiation oncologist manually segmented prostate, seminal vesicles, and nodes based on the CT images. The primary planning target volume (PTV) was defined to include a 1.0 cm margin around the pelvic lymph nodes in all directions plus a 0.7 cm margin around prostate, and seminal vesicles in all direction except the posterior direction, where 0.5 cm margin was added. The boost planning target volume (PTVs) was defined to include 0.7 cm margin around the prostate and seminal vesicle in all direction except the posterior; where 0.5 cm margin was added.

Rectum, bladder, bowel, left and right femoral head were contoured as organ at risk (OARs) based on CT images. In all the patients, anterior-posterior (AP) and lateral separation were very close. The average AP diameters were 21 cm and the lateral diameters were 34 cm. The average volume of PTVr and PTVs were 534.47 cc (range, 860.10 cc-360.00 cc) and 163.50 cc (range, 181 cc-148 cc), respectively. The average volume of bladder, rectum, right femoral head, left femoral head and bowel were 223.51 cc (range, 87.21 cc-360.32 cc), 68.61 cc (range, 42.91 cc-111.60 cc), 53.51 cc (range, 68.13 cc-40.31 cc), 53.47 cc (range, 37.90 cc-63.81 cc) and 1243.90 cc (range, 380 cc-1680 cc), respectively. For this study, we have followed Radiation Therapy Oncology Group (RTOG-0521) protocol. The total prescription dose was 75.6 Gy with a daily dose of 1.8 Gy. The prescription dose of the primary plans was 45 Gy and an additional 30.6 Gy to boost plan. Two arcs (182° to 178°; 178° to 182°) were used in both clockwise and anti-clockwise directions for all patients in RapidArc treatment planning. For the clockwise arc, the collimator was rotated 20°, whereas for the anti-clockwise arc, the collimated was rotated 340° in order to reduce the effect due to inter-leaf leakage. Seven fixed fields with angulations of 0°, 51°, 102°, 153°, 207°, 208°, and 309° with dynamic leaf were used for IMRT. All plans were generated using Eclipse treatment planning system (version 8.0.15) by a single user only and the volumetric dose optimization method followed the same systematic strategy regarding the objective and priorities. Dose grid sizes of (2.5×2.5×2.5) mm³ and the anisotropic analytical algorithm (AAA) were used in this study. The slice thickness of the CT images used for planning purpose was 3 mm. Both IMRT & RapidArc plans were developed for each patient using 6 MV photons with maximum dose rate of 600 MU/min in a Varian made Novalis Tx machine; having dynamic capability of 120 high definition Multi-leave collimator (MLC). The optimization constraints for all plans are listed in Table 1. These constraints and weightings were set initially and then modified by either relaxing or tightening during the optimization process based on the real-time updated dose-volume histograms (DVHs) of structures.

| PTVs and OARs       | Dose-Volume constraints                                                                 | Relative weighting |
|---------------------|----------------------------------------------------------------------------------------|--------------------|
| PTVr and PTVs       | D10 ≥ 98%; D5 ≤ 102%                                                                  | 120-130            |
| Rectum and          | V70% ≤ 30%; V50% ≤ 50%; V50% ≤ 70%; D5-D10 total of 45 Gy                             | 60-80              |
| Bladder             | 60-70 Gy; Dmax ≤ 100%                                                                  |                    |
| Bowel               | V50% ≤ 30%; V30% ≤ 50%; V10% ≤ 70%; Dmax stotal of 45 Gy                             | 50-60              |
| Femoral heads       | V20% V30% ≤ 30%; Dmax stotal of 45 Gy                                                 | 40-50              |

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2.1 Sigma index (S-index) and conformity index (CI)

Treatment planning was done based on the radiotherapy objective of delivering a therapeutic dose to a well-defined target while minimizing dose to the surrounding normal tissue and critical organs. It requires optimization of conformity of the prescription dose to the planning target volume, dose homogeneity within the PTV, dose to the surrounding normal tissue and critical organs. The following parameters were evaluated for comparing cumulative DVH of the patient such as Sigma-index, conformity index, and mean doses to the bladder, rectum, bowel, right femoral head, and left femoral head.

A better representation of homogeneity could be defined in term of sigma index (S-index). This uses the differential DVH, unlike other indices, which is cumulative DVH. S-index is a measure of the standard deviation of the doses about the mean dose. The S-index is expressed as:

\[ S\text{-index} = D_{SD} = \sum (D_i - D_{mean}) \times \frac{V_i}{V} \]

D_{mean} is the mean dose of the target (PTV in this study) curve. D_i is dose to the i-th bin having a volume v_i. V = total volume of the target.

The conformity index (CI) was defined to compare the treatment plans. The conformity index is defined as the ratio of the 95% isodose volume divided by the PTV that is enclosed by the 95% of the isodose line. From this definition, as the conformity index approaches 1, the more conformal is the treatment plan.

2.2 Dose distribution, PTV coverage and organs at risk

In addition to providing information on the homogeneity of radiation doses, DVHs can be used to assess the target coverage index, defined as the percentage of the tumor volume that received the prescribed dose. Ideally, tumor DVH would be a step function, with 100% of the target receiving the exact prescribed dose. However, actual DVH curves are not step functions, because of constraints imposed by tumor volume and other OAR. Rules for the PTV was set such that the prescribed dose covered at least 95% of the PTV (D95) and the PTV volume receiving >107% of the prescription was limited to 2% in line with ICRU report 50 and ICRU report 62. To achieve this objective, a constraint for D100 was set to receive ≥98% of prescription and constraint for maximum dose (D_{max}) was set to receive ≤102% of the prescription in the optimization process for both plans. The volumes that received a minimum of 70 Gy, 60 Gy, 40 Gy, 40 Gy, 40 Gy, 15 Gy, 20 Gy (V_{70Gy}, V_{60Gy}, V_{40Gy}) for rectum, 70 Gy, 60 Gy, 40 Gy, 40 Gy, 20 Gy, 15 Gy (V_{70Gy}, V_{60Gy}, V_{40Gy}) for bladder and 45 Gy, 15 Gy (V_{45Gy}, V_{15Gy}) for bowel, respectively, were selected to evaluate volumes of high dose,^{12-13}

2.3 Normal tissue integral dose and MU

The integral dose (in unit of liter-Gy) was defined as the absorbed dose integrated over the voxels in the entire volume excluding the PTV. The integral dose (ID) (mean dose × tissue volume) received by normal tissue (NTID) was calculated from dose-volume histograms. For comparison, the integral dose ratio was obtained by dividing the integral dose from IMRT plan by the integral dose from RapidArc plan. MU was analyzed for IMRT and RapidArc plans.

2.4 Statistical analysis

Statistical analysis was performed with Statistical Analysis Software (SAS), version 9.3. Mean values and standard deviation of the mean (SD) were collected. Relative dosimetric changes were compared applying the paired t-test. A two-sided p-value ≤ 0.05 was considered statistically significant. Confidence intervals included 95% of the measured data.

3. Results

3.1 Sigma index (S-index) and conformity index (CI)

The S-index and conformity indices for all the prostate patients in the both IMRT and RapidArc plans with 6 MV photons are analyzed. The relative efficacy (IMRT/RapidArc) values for, S-index and conformity index are 0.9832, 1.010 for PTVr and 1.0, 1.005 for PTVs respectively. Table 2 shows S-index and conformity index of both IMRT and RapidArc Plans in all type of PTVs.

| Technique       | SI       | CI       |
|-----------------|----------|----------|
| PTVr IMRT       | 2.35±0.581 | 0.96±0.047 |
| PTVr RapidArc   | 2.39±0.738 | 0.95±0.059 |
| PTVs IMRT       | 2.1±0.484  | 0.97±0.015  |
| PTVs RapidArc   | 2.1±0.464  | 0.97±0.014  |

3.2 Dose distribution, PTV coverage and organs at risk

In this study, we analyzed a total of 80 prostate cancer treatment plans of 20 patients. The clinical dosimetric impact of IMRT, in comparison with RapidArc, was assessed. Transversal view of comparative dose distribution between IMRT plan (left side) and RapidArc plans for PTVr (right side) for prostate cancer with lymph nodes involved is shown in Figure 1. A typical DVHs for 6-MV plans (IMRT and RapidArc) are compared in Figure 2.
Figure 1: Transversal view of comparative dose distribution between IMRT plan (Left Side) and RapidArc plans for PTVP (Right side) for prostate cancer with lymph nodes involved.

Figure 2: Comparison of the dose-volume histograms for IMRT (square) and (triangle) for RapidArc plan for the case shown in Figure 1.

Table 3: Mean dose statistics (with SD) of organ at risk for composite plan.

| Structure          | Dose-volume constraints | Volume achieved in % |
|--------------------|-------------------------|----------------------|
|                    |                         | IMRT | RA               |
| Rectum             | $V_{70} < 30\%$         | 63.0±5.9% | 69.4±4.4%       |
|                    | $V_{40} < 50\%$         | 50.0±4.1% | 57.4±3.0%       |
|                    | $V_{40} < 70\%$         | 38.2±3.3% | 39.5±2.6%       |
| Bladder            | $V_{70} < 30\%$         | 63.3±10.2% | 69.4±9.4%      |
|                    | $V_{40} < 50\%$         | 45.0±7.4% | 49.5±7.7%       |
|                    | $V_{40} < 70\%$         | 33.0±5.6% | 38.0±5.8%       |
| Bowel              | $V_{45\text{Gy}} < 195\text{cc}$ | 4.49±1.08% | 5.00±1.09%     |
| Rt. femoral head   | $V_{50\text{Gy}} < 5\%$ | 0.0±0.0% | 0.0±0.0%         |
| Lt. femoral head   | $V_{50\text{Gy}} < 5\%$ | 0.0±0.0% | 0.0±0.0%         |
For the rectum, there were dose differences between the two treatment modalities in the volume exposed to 70 Gy, 60 Gy, and 40 Gy are 10%, 14.8% and 3.4%, respectively, which is statistically significant ($p < 0.0001$). In case of bladder, there were dose differences for the volume exposed to 70 Gy, 60 Gy and 40 Gy are 9.63%, 10% and 15%, respectively, which is statistically significant ($p < 0.0001$). For the bowel, there were no significant dose differences between two treatment modalities in the volume exposed to 45 Gy or 15 Gy, and femoral heads were within the required constraints for all patients. Table 3 shows mean dose statistics with standard deviation of organ at risk for composite plan.

Table 4 shows the average maximum dose statistics (with SD) and average dosimetric indices with standard deviation of PTV and organ at risk for composite plan. There were significant differences between IMRT and RapidArc plans for the average doses delivered to the bladder and rectum. A significant increase in mean dose of 12% ($p < 0.0002$, 95% CI = 0.7714 to 2.2312) to the rectum and 15.4% ($p < 0.0001$, 95% CI = 4.294 to 6.771) for bladder was measured for RapidArc plans compared with IMRT plans. For the bowel, there was no significant dose difference between two treatment modalities in the average doses and femoral heads were within the required constraints for all patients. The average maximum dose of PTV is 106.7% and 105.7% of prescribe dose for both IMRT and RapidArc plans, respectively and different was significant ($p < 0.0002$, 95% CI = 45.49 to 127.5). No significant differences between two modalities were observed with respect to mean average maximum dose to bladder, rectum, femoral heads, and bowel.

### 3.3 Normal tissue integral dose (NTID) and MU

NTID was used as an index to compare techniques for radiation induced second cancers. The NTID (mean ± SD) was 182.07 ± 46.07 liter-Gy for IMRT plan and 183.65 ± 47.26 liter-Gy for Rapid Arc plans. RapidArc plans produced 0.87% ($p < 0.044$, 95% CI = 0.045 to 3.113) more dose compared with the IMRT beam, and this difference is not significant. Comparison of the integral dose for IMRT and in RapidArc composite plans is shown in Figure 3. We found that MU (mean ± SD) was 1542 ± 239 for IMRT plans and 618 ± 71 for RapidArc plans in PTV. MU (mean ± SD) was 1041 ± 195 for IMRT plans and 605 ± 78 for RapidArc plans in PTV. Compared with RapidArc plans, the MU significantly increased by 2.50 times ($p < 0.0001$, 95% CI = 368.749 to 502.3) in PTV and 1.72 times ($p < 0.0001$, 95% CI = 846.4 to 998.2) in PTV compared with IMRT plans shown in Figures 3-5. The beam–on time of PTVp was 4.2 min to 4.3 min for IMRT and 2.1 min to 2.3 min for RapidArc; whereas for PTVb, it was 2.5 min to 2.6 min for IMRT and 1.0 min to 2.3 min for RapidArc. Treatment time of PTVp was 8.1 min to 8.2 min for IMRT and 3.1 min to 3.2 min for RapidArc. The treatment time of RapidArc was 1.5 times shorter than IMRT for PTVp but approximately same in PTVb plans.

**Table 4:** Average maximum dose statistics (with SD) and average dosimetric indices (with SD) of PTV and organ at risk for composite plan.

| Structure   | IMRT Mean dose in Gy | IMRT Maximum dose in Gy | RA Mean dose in Gy | RA Maximum dose in Gy |
|-------------|----------------------|-------------------------|-------------------|-----------------------|
| PTV         | 76.50±0.82           | 80.89±1.31              | 76.50±0.72        | 80.13±0.93            |
| Rectum      | 37.9±4.75            | 76.44±6.81              | 42.7±4.59         | 77.77±7.75            |
| Bladder     | 35.7±11.86           | 79.46±0.56              | 41.2±12.0         | 78.92±0.66            |
| Bowel       | 23.0±2.89            | 45.6±2.42               | 23.08±2.84        | 45.7±23.6             |
| RT FH       | 24.5±1.88            | 47.47±4.44              | 25.3±1.82         | 49.35±5.13            |
| LT FH       | 24.4±1.85            | 47.86±4.60              | 25.2±1.97         | 50.51±6.02            |

![Figure 3: Comparison of the integral dose for IMRT and RapidArc composite plans.](image)
4. Discussion

It clearly represents that there is no significant difference of homogeneous index for both the PTVs in IMRT and RapidArc plans. From our data between IMRT and RapidArc plans, IMRT plans are showing lower S-indices for both PTV_P and PTV_B plans indicating improved dose homogeneity compared to RapidArc plans. However, target inhomogeneity (defined as Dmax-Dmin) was increased by 12.3% (p < 0.0001, 95% CI = 2.256 to 3.677) for IMRT compare with RapidArc composite plans. For IMRT plans, target inhomogeneity is significantly increased for PTV_P compared to RapidArc plans because of lymph nodes involvement with PTV_P. However, in PTV_B, inhomogeneity is comparable for both plans.

Miften et al.\textsuperscript{15} have demonstrated the use of target conformity index (TCI) and normal tissue-sparing index (NTSI) to assist in the process of judging the merit of a clinical treatment plan. However, in this work, the widely accepted conformity index was used to evaluate the conformity of the treatment plans (Table 2) for all the patients using both techniques. The mean conformity index was 0.96 and 0.95 for PTV_P and 0.979 and 0.975 for PTV_B in both IMRT and RapidArc plans, respectively. These small differences indicate that these plans are nearly identical in their conformity of dose to the target. However, there is difference observed in between PTVs. For PTV_P, conformity index is lower values than PTV_B plans due to involvement of lymph nodes. Sua et al.\textsuperscript{16} reported similar results in their planning study for PTVs involved with pelvic lymph nodes.

Rectum and bladder are showing high dose for RapidArc plans comparable to IMRT plans. This dose variation was significantly increased in PTV_p compared to PTV_B. Sua et al.\textsuperscript{16} reported that IMRT plans with PTV_P showed better sparing of bladder, rectum, and small bowel than 2 arcs plans. Kjaer-kristoffersen et al.\textsuperscript{17} noted larger OAR doses with RapidArc than IMRT which is not confirmed in our study. Palmer et al.\textsuperscript{20} noted larger integral doses with IMRT than with IMAT based on MU values for prostate.

Because radiation could induce second malignancies\textsuperscript{18, 19}, integral dose was used as an index to evaluate treatment plans. In our study, integral dose index (IMRT/RapidArc) is 0.99, which is very negligible. Sua et al.\textsuperscript{16} reported that 7% to 8% greater integral dose in RapidArc than in IMRT which is not confirmed in our study. Palmer et al.\textsuperscript{20} noted larger integral doses with IMRT than with IMAT based on MU values for prostate.
cancer which is conflict with our findings. Integral dose is related not only to MUs but also to other complicated factors. The combination of MUs, corresponding aperture sizes and shapes, target volumes and shapes contributes to integral dose.\textsuperscript{16} In the primary plans, MUs was approximately 60% greater than in RapidArc plans. In boost plans, the average values of total MUs in IMRT plans was approximately 42% greater than in RapidArc plans.

The IMRT planning was performed in Eclipse treatment planning system using Novalix Tx Linac and 120HD MLC, which have a limit that MLC cannot travel beyond 14 cm during beam delivery. To overcome this limit, a large IMRT beam is split into two or three subfields, which are planned and calculated as one beam but delivered as two or three separate beams.\textsuperscript{16} Without this limitation, IMRT delivery time could be reduced by approximately 1.2 min for a prostate cases that include the lymph nodes. The mean delivery time of IMRT was longer owing to the larger number of beams, the dead time in the gantry rotation from field to field and the larger number of MUs delivered. The shorter treatment time of RapidArc is beneficial to the patient, as it would help reduce the time the patients required to maintain a full bladder status. Shorter treatment time does not only reduce the probability of the intra-fractional motion of patients, but it can also decrease the impact of internal organ motion on the treatment delivery.\textsuperscript{24} The shorter treatment time of RapidArc would be welcomed by busy centers and shorten the patient waiting list.

Although our study focused on photon therapy, proton therapy is also another option to treat the prostate cancer in external beam radiotherapy.\textsuperscript{22, 23} Proton therapy can produce excellent dose distribution because protons have finite range and sharp distal fall-off at the end of proton beam path. Vargas et al.\textsuperscript{24} and Rana et al.\textsuperscript{25} demonstrated that proton therapy is better at sparing rectum and bladder when compare to the photon therapy.

5. Conclusion

In this study, the clinical dosimetric impact of IMRT, in comparison with RapidArc, was assessed for prostate cancer patients. For complicated and large PTVs that included prostate, seminal vesicles and lymph nodes, conventional IMRT spared the bladder, rectum doses better than did RapidArc. For simple and small PTVs that included prostate and seminal vesicles, RapidArc plans were comparable to those achieved with conventional IMRT plans. The integral dose was more in RapidArc but not significant. The treatment delivery efficiency improved with RapidArc plans. RapidArc plans should be compared with IMRT plans for individual cases to measure gains and losses before selecting one over the other. Clinical studies are required to evaluate its clinical benefits.

Conflict of interest

The authors declare that they have no conflicts of interest. The authors alone are responsible for the content and writing of the paper.

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