Dosimetric Importance of the Implementation of Daily Image Guidance in Radiotherapy Practice

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Abstract

Background: The purpose of this study is to show the dosimetric importance of using daily image guidance in radiotherapy treatment. Methods: A sample of 30 patients with various head and neck cancers were retrospectively selected for the studies. The prescribed dose, 66 Gy/33 fractions was used for all patients who received 7 to 9 beams, 6 MV Intensity Modulated Radiation Therapy (IMRT) plan delivered by Varian Truebeam STx. Before the first fraction of the treatment, the patient’s shift corresponds to isocenter was noted and corrected. Subsequently, the images were taken daily for all 30 patients and the maximum, mode, median and mean of all shifts were recorded and applied to the base plan and recalculated for quantitative analysis of tumour coverage and Organ at Risk’s (OARs) doses using various dosimetric parameters such as Homogeneity Index (HI), Conformation Number (CN), Conformity Index (CI), Coverage Index (COVI), Dose Gradient Index (DGI) and Unified Dosimetry Index (UDI) of shift plan. Results: The results showed the Planning Target Volume (PTV) and the OARs values deteriorated from its base plan values in the various shift plans created by applying the patient setup errors analyzed using image guidance. Mean dose of maximum shift plan with a significant P value of 0.002, D2% of maximum shift plan with a significant P value 0.028, the D98% Values of maximum and mode plans with the significant P value 0.004 each, the D50% and D80% values of the maximum shift plans with their significant P values 0.001 and 0.002 respectively. Also, HI95%, CN95% and COVI values of the maximum shift plan showed much variation with significant P values of 0.004, 0.040 and 0.0004 respectively from their base plan values. There were significant changes observed in OARs values between base plan and shift plans. Conclusion: Implementation of daily image guidance in radiotherapy is mandatory taking into account of its dosimetric importance and to achieve the goal of radiotherapy practice of sparing the critical organs without compromising the target coverage.

Keywords: Head and Neck cancers- IGRT imaging- shift in IGRT and Imaging in Radiotherapy

Introduction

The main aim of radiotherapy is to relieve the cancer cells from their multiplication potential which itself makes it an important factor of cancer treatment (Baskar et al., 2012). And it is very significant in radiotherapy to maximize radiation exposure and dose deposition in the cancer cells and at the same time, to reduce the radiation exposure over the adjacent normal cells to its minimum (Van’t et al., 1997). Radiotherapy is a technology-oriented treatment modality and it has evolved into an extremely complex technique from the conventional treatment modes to highly advanced conformal radiotherapy techniques (Garibaldi et al., 2017). Advancements in radiotherapy have occurred in all sectors of radiotherapy workflow, starting from the different imaging modalities like Computed Tomography [CT] simulation, Magnetic Resonance Imaging [MRI] or Positron Emission Tomography [PET] for the accurate identification and delineation of target volumes (PTV) and OARs (Metcalfe et al., 2013; Ashamalla et al., 2005), then with the advanced registration and fusion algorithms in Treatment Planning Systems (TPS) to aid in precise contouring work as well as the highly sophisticated and advanced radiobiology-oriented treatment planning optimization tools (Nahun et al., 2012). In the treatment execution process great strides have been made by the emergence of new image-guided techniques with which the tumor can be localized very accurately before and during the treatment procedure (Jaffray et al., 2012; Nabavizadeh et al., 2016).

Intensity-modulated radiotherapy (IMRT) is a radiotherapy technique that allows the delivery of steep dose gradients with multiple beam angles to modulate the intensity of the dose. This technique helps in delivering a highly conformal dose to the tumour along with the advantage of sparing the adjacent OARs to a higher extent.
as compared to the old techniques used (Cheung et al., 2006). IMRT is the most suitable treatment modality compared to 3 Dimensional Conformal Radiotherapy (3DCRT) for heterogeneous head and neck cancers because these techniques take into account of the complex shape of the target and the close proximal OARs. The IMRT technique addresses the discrepancies like under dosage of target volumes and overexposure of OARs as well as giving better dose hotspot, improved target volume dose distribution and reduced radiation dose to OARs (Hummel et al., 2010; Murshad et al., 2004).

Image-guided radiotherapy [IGRT] is a step ahead of IMRT and it has played an important role throughout radiotherapy history as its introduction and subsequent innovations have revolutionized cancer treatment. The IGRT uses advanced imaging modalities for accounting and thereby adjusting the target motion uncertainties and positional uncertainties (Kearney et al., 2020).

The need for an image guidance in radiotherapy practice is very important due to many reasons like tumor regression with treatment response, weight loss due to less nutrition or perhaps growth itself inspired by treatment. There could be interfractional changes in treatment between each radiation fraction due to variations in patient positioning. The patient setup in each radiotherapy fraction is affected by different factors such as variations in patient setup, dosimetric uncertainties, mechanical uncertainties and human errors (Weygand et al., 2016). There are many reports that have studied the impact of these errors on PTV coverage and OARs. A positional error can increase dosimetric error and it leads to the reduction of therapeutic index. Recent studies have examined how the setup errors in head and neck cancer patients treated with IMRT have influenced the customized immobilization systems and other patient factors (Contesini et al., 2017). A study which was carried out on the effect of setup errors and organ motion on prostate cancer treated with IMRT showed that in most of the recalculated plans, the rectal wall dose values were very much altered from the planned one despite the small variations seen in the Clinical Target Volume (CTV) coverage (Landoni et al., 2006).

The IGRT uses 2D or 3D images which allow for correcting the patient position to radiotherapy treatment position before radiation delivery and paves way for precise treatment delivery. The radiotherapy treatment has advanced from the early conventional methods of skin markings on the patient body to recent advancements in IGRT such as CT, Cone Beam CT (CBCT), MV and KV On-Board Imaging (OBI) that helps in precise internal anatomy based patient positioning during treatment. It is also significant to manage the imaging dose exposure to the patients as it may result in excessive dose to the OARs and increase the chance of inducing secondary cancers (Ding et al., 2018).

The practice of radiotherapy in hospitals is very much related to the workload or work quality in the respective department. Hence this present study is intended to find out the dosimetric effect on non-IGRT practices. The main aim of this study is to find out the level of dosimetric error that arises from the actual plan i.e. whether the approved IMRT delivery plan delivers the same intensity-modulated beam plan at the treatment position by the image guidance treatment procedures. Moreover in this study, the possible degrees of positional shift errors and its impact on the plan execution results has been verified by taking the maximum, mode, median and mean of the daily shift values.

Materials and Methods

Patient Selection

Thirty patients with various head and neck cancers were retrospectively selected for this study. The patients selected were in the age group of 45 to 75 years after getting approval from the institutional ethical committee.

Treatment Planning

The patients selected for the study were simulated in the headfirst supine position using a thermoplastic mould and shoulder retraction, for which 3mm slice thickness CT images were acquired. The acquired images were then transferred to the Eclipse TPS version 15.06. The MRI images taken in the same position was sent to the TPS which was later registered and fused with the CT images for better localization of tumour and OARs and based on the treatment protocol, the target volumes, as well as the critical organs, were delineated. Initially, the Gross Tumour Volume GTV) was delineated including all positive lymph nodes with the aid of fused images. Following the GTV, the Clinical Tumour Volume CTV was contoured taking into account of the primary tumour size, its involved nodes and then a margin for the microscopic spread. Planning Target Volume PTV was contoured by the expansion of the CTV by 5mm in all directions [International Commission on Radiation Units and Measurements (ICRU) 50 and 62]. Then the critical organs or the OARs such as spinal cord, left parotid, right parotid, larynx, oral cavity, left eye, right eye, left optic nerve, right optic nerve, brainstem, mandible, left cochlea, right cochlea and optic chiasm were contoured. An additional healthy tissue was also defined which is the patient CT volume excluding the PTV volume.

The dose prescribed for the tumour volume of these patients taken for this study was 66Gy in 33 fractions (2Gy per fraction). For all the patients 7-9 beam IMRT plan was generated with gantry angles 0°, 50°, 100°, 150°, 200°, 250°, 300°. The plan was optimized for 6 MV X-ray photon delivered by Varian Truebeam STx linear accelerator equipped with an HD Multi-Leaf Collimator (MLC) with 120 leaves using Photon Optimizer (15.6.05) algorithm to achieve the given planning objectives. Dose calculation was done using the Anisotropic Analytical Algorithm (AAA) at 2.5mm calculation grid size. The plan objectives for PTV were that at least 95% of the PTV should cover 95% of the dose prescribed and the dose maximum should not exceed a value of 107%. The dose constraints followed for OARs like spinal cord, bilateral parotids, larynx, optic nerve, mandible, cochlea, brainstem, eye were D1% < 46 Gy, Dmean < 26 Gy; V30Gy < 45%, Dmean<45Gy, D1%<50Gy, D1%<61Gy, Dmean < 15Gy, D1% < 54Gy, Dmax < 45Gy respectively where D1% - Dose to 1 % of volume, Dmean-Mean dose,
Volume. The ideal values of CN should be 1. The DGI was noted down using the formula PI/D50% whose ideal value is 1, where PI is the prescribed isodose volume and D50% is the volume of 50% of prescribed isodose volume. The coverage index (COVI) as TVpi/TV and Conformity Index (CI) as PI/TV were calculated and noted down whose ideal value is 1. Unified Dosimetry Index (UDI) is tool used to compare and evaluate any given treatment plans by its deviations in terms of dose coverage, conformity, homogeneity, and dose gradient. It is calculated using the formula:

$$ UDI = UDI[CI] \times UDI[CF] \times UDI[HI] \times UDI[DG] $$

whose ideal value is 1, where the CI-coverage index, CF-conformity index, HI -Homogeneity Index and DG-gradient index. The mathematical logic-based UDI formula is:

$$ UDI = \left( \prod_{k=1}^{4} W_k \left[ \left| D_{1.0} - D_{Lk} \right| + 0.1 \right] \right) \times 10^4 \quad (1) $$

Where $D_{Lk}$ is dosimetry index, each of the four indices and $W_k$ is the weighting factors as per the relative importance of all the four components. For the OARs, the mean dose, the maximum dose and appropriate values of volume receiving xGy were noted. The calculated difference of the patients were analysed statistically using one sample ‘t’ test and considered significant if the P value was less than 0.05.

**Results**

The dose distribution of base plan and all other compared plans of Maximum, Mode, Median and Mean shift plans are shown in Figure 1.

Similar observations were plotted in other graphs generated. And the results show that the base plan parameters got deteriorated from their actual base plan values in the shift plans created by applying the maximum, V30Gy -Volume receiving 30 Gy and Dmax - Maximum dose respectively.

The plans were analysed and approved based on Dose Volume Histograms (DVH), tumour coverage and OAR dose. Before delivering the approved plan (base plan), a set of verification images were taken to ensure that the patient position is the same as the simulated position which was used for planning. The observed shifts in the patient position by matching methods was taken and recorded and as Mohandass et al., (2020) stated in his study any of the matching methods can be used during CBCT to check patient translation errors for the delivery of the VMAT head and neck patients. After verifying the position and making necessary corrections, the plan was delivered. The maximum, mode, median and mean of all observed shifts were calculated. Then the shift values were intentionally applied to the approved base plan, recalculated and was named as shift plans, to assess the dosimetric deviation from the original approved plan if the image verification procedure was not carried out continuously. The DVH and tumour coverage of these shift plans were also found and recorded. Various parameters such as Homogeneity Index (HI), Conformation Number (CN) and Dose Gradient index (DGI) were used for evaluating the shift plan as compared to base plan. The homogeneity index is defined as;

$$ Homogeneity \ Index \ (HI) = \frac{D_{2\%} - D_{98\%}}{D_{50\%}} $$

Where D2%, D98% and D50% are dose received by 2%, 98% and 50% volumes respectively. The ideal value of HI is 0. The CN was calculated for each plan to evaluate the conformity of dose to target by the following formula;

$$ CN_{95\%} = \frac{TV_{pi}/TV}{TV_{pi}/V_{pi}} $$

Where TVpi is Target Volume within the 95% prescribed isodose volume, Vpi is the Volume of 95% of prescribed isodose volume and TV is the tumour volume. The ideal values of CN should be 1. The DGI was noted down using the formula PI/D50% whose ideal value is 1, where PI is the prescribed isodose volume and D50% is the volume of 50% of prescribed isodose volume. The coverage index (COVI) as TVpi/TV and Conformity Index (CI) as PI/TV were calculated and noted down whose ideal value is 1. Unified Dosimetry Index (UDI) is tool used to compare and evaluate any given treatment plans by its deviations in terms of dose coverage, conformity, homogeneity, and dose gradient. It is calculated using the formula:

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Table 1. PTV Parameters for Base Plan and Shift Plans

| Parameters | Base Plan | Maximum | Mode | Median | Mean | P value (Max vs Base) | P value (Mode vs Base) | P value (Median vs Base) | P value (Mean vs Base) |
|------------|-----------|---------|------|--------|------|-----------------------|-----------------------|------------------------|-----------------------|
| Mean dose  | 62.092±3.900 | 61.196±4.140 | 61.639±3.960 | 61.723±4.050 | 61.952±3.900 | 0.002 | 0.012 | 0.021 | 0.052 |
| D2%        | 65.062±3.900 | 65.982±4.120 | 65.244±3.930 | 65.262±3.890 | 65.139±3.900 | 0.028 | 0.057 | 0.059 | 0.148 |
| D98%       | 55.068±3.790 | 47.784±7.010 | 52.846±4.200 | 53.452±4.570 | 54.259±3.900 | 0.004 | 0.004 | 0.009 | 0.01 |
| D95%       | 62.634±3.900 | 62.294±4.000 | 62.465±3.910 | 62.634±3.930 | 62.541±3.900 | 0.001 | 0.073 | 1 | 0.09 |
| CI95%      | 60.723±3.960 | 59.443±4.640 | 60.057±4.070 | 59.560±3.890 | 60.542±4.000 | 0.002 | 0.017 | 0.091 | 0.078 |
| HI95%      | 1.268±0.270 | 1.203±0.260 | 1.168±0.170 | 1.169±0.170 | 1.186±0.190 | 0.516 | 0.246 | 0.247 | 0.354 |
| COV1       | 0.160±0.020 | 0.294±0.110 | 0.199±0.036 | 0.189±0.036 | 0.174±0.020 | 0.004 | 0.005 | 0.006 | 0.008 |
| CN95%      | 0.691±0.120 | 0.599±0.100 | 0.674±0.098 | 0.697±0.103 | 0.696±0.100 | 0.04 | 0.504 | 0.86 | 0.891 |
| DGI        | 0.920±0.040 | 0.837±0.050 | 0.897±0.050 | 0.894±0.045 | 0.900±0.047 | 0 | 0.005 | 0.002 | 0.013 |
| UDI        | 98.904±68.670 | 123.631±107.070 | 99.559±75.990 | 95.323±74.890 | 96.500±73.960 | 0.269 | 0.959 | 0.795 | 0.87 |

Figure 2. The DVH Comparison of PTV of a Patient for All the Plans

Figure 3. The Graphical Comparison of V30 and Mean Dose Values of Both Left Parotid and Right Parotid
Table 2. OARs Parameters for Base Plan and Shift Plans

| Parameters     | Base plan | Maximum | Mode | Median | Mean | Maximum | Mode | Median | Mean | P Value |
|----------------|-----------|---------|------|--------|------|---------|------|--------|------|---------|
| Spinal cord    | D1%       | 37.67±2.20 | 40.59±4.60 | 38.66±4.20 | 38.34±2.90 | 40.51±6.50 | 0.06 | 0.25 | 0.14 | 0.26 |
|                | Dmax      | 40.77±2.10 | 44.74±5.90 | 42.77±4.90 | 41.62±2.20 | 41.21±1.70 | 0.04 | 0.12 | 0.04 | 0.06 |
| Lt parotid     | Mean dose | 32.01±7.10 | 42.06±6.90 | 33.87±11.10 | 34.90±13.40 | 32.02±6.90 | 0.00 | 0.26 | 0.26 | 0.99 |
|                | V10       | 50.86±16.80 | 70.40±15.40 | 54.20±23.40 | 51.91±19.30 | 50.57±16.70 | 0.00 | 0.33 | 0.71 | 0.82 |
| Rt parotid     | Mean dose | 38.09±11.30 | 36.81±11.30 | 37.70±11.40 | 38.38±12.80 | 38.19±12.10 | 0.22 | 0.66 | 0.8 | 0.91 |
|                | V10       | 63.51±24.20 | 61.88±26.30 | 61.82±26.20 | 63.54±26.90 | 63.20±25.80 | 0.48 | 0.44 | 0.99 | 0.86 |
| Larynx         | Mean dose | 42.12±12.00 | 41.24±4.90 | 39.74±7.50 | 39.76±7.40 | 41.5±10.70 | 0.83 | 0.46 | 0.46 | 0.48 |
|                | V10       | 51.21±38.00 | 53.84±23.80 | 48.30±31.90 | 48.39±31.60 | 49.80±37.40 | 0.81 | 0.52 | 0.53 | 0.05 |
| Lt eye         | Mean dose | 2.35±1.60 | 3.34±4.20 | 3.01±4.30 | 2.26±2.10 | 2.08±1.60 | 0.33 | 0.53 | 0.75 | 0.19 |
|                | V10       | 22.0±1.20 | 3.24±3.20 | 2.61±2.90 | 2.16±1.60 | 2.02±1.20 | 0.18 | 0.55 | 0.89 | 0.22 |
| Rt optic nerve | D1%       | 3.15±2.50 | 8.55±14.20 | 6.93±13.90 | 2.33±0.90 | 3.02±2.40 | 0.21 | 0.36 | 0.29 | 0.28 |
|                | Dmax      | 3.62±3.40 | 10.15±16.04 | 5.80±9.80 | 3.77±3.90 | 3.29±2.70 | 0.16 | 0.44 | 0.85 | 0.32 |
| Brainstem      | Maximum dose | 40.44±8.82 | 41.33±10.02 | 38.53±11.75 | 39.08±10.02 | 39.54±9.78 | 0.58 | 0.24 | 0.29 | 0.44 |
| Mandible       | Maximum dose | 62.94±4.00 | 65.30±4.90 | 63.96±4.40 | 63.98±4.50 | 63.31±4.30 | 0.05 | 0.08 | 0.08 | 0.28 |
| Lt cochlea     | Mean dose | 14.11±15.90 | 25.04±22.40 | 21.87±23.60 | 19.89±20.50 | 14.56±14.90 | 0.13 | 0.29 | 0.42 | 0.80 |
| Rt cochlea     | Mean dose | 17.97±26.30 | 27.44±26.70 | 23.51±28.30 | 23.07±27.50 | 18.25±25.50 | 0.20 | 0.45 | 0.49 | 0.87 |
| Healthy tissue | V10       | 27.46±8.30 | 26.64±8.20 | 27.46±8.70 | 27.53±8.70 | 27.63±8.50 | 0.00 | 0.99 | 0.36 | 0.13 |
|                | V10       | 11.25±3.40 | 10.99±3.40 | 12.84±6.90 | 11.16±3.40 | 11.22±3.40 | 0.00 | 0.38 | 0.12 | 0.34 |
| Mean dose      | 8.20±2.40 | 8.13±2.30 | 8.18±2.50 | 8.18±2.50 | 8.21±2.40 | 0.04 | 0.12 | 0.04 | 0.06 |

Discussion

Different planning studies emphasize the importance of image guidance in radiotherapy treatment for tumor coverage (Ariyaratne et al., 2016; deCrevoisier et al., 2018; Wertz et al., 2007). Ariyaratne et al., (2016) did a study in which they showed that how it enhanced target scope day by day online in contrast with week by week imaging along with the decrease in rectal measurements and the possibility of PTV edge lessen to 5mm with satisfactory target scope (Verma et al., 2014). From our study results it is very clear that in all the plans created by taking the maximum, mode, median and mean of the shift values inferred from image guidance, all the

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DOI:10.31557/APJCP.2022.23.1.355

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dosimetric parameters of PTV values deviated from the actual base plan. The dosimetric parameters which showed the maximum variations were a mean dose of maximum shift plan with a significant P value 0.002, D2% of maximum shift plan with a significant P value 0.028, the D98% values of maximum and mode shift plans with the significant P values 0.004 each, the D50% and D80% values of the maximum shift plan with their significant P values 0.001 and 0.002 respectively.

The HI95%, CN95% and COVI values of the maximum shift plan showed significant P values of 0.004, 0.040 and 0.0004 respectively from their base plan values. In a study conducted by Landoni et al., (2006), the effect of organ motion and setup errors on prostate cancer treatment with IMRT was analysed and it was concluded that by combining the patient motion and setup errors accurately, the treatment will be more defined. In another study by Wang et al., (2008) on the dosimetric effect of translational and rotational errors on spinal metastases patients treated with image-guided stereotactic body radiotherapy, it was stated that a 2mm error in translational patient positioning in any direction will cause a tumor coverage loss of >5% and an increase in maximal dose to the organs at risk by >25%. The study emphasized the importance of keeping the patient setup translational errors to ≤ 1mm and rotational errors to ≤ 2 degrees for obtaining the best dosimetric results in spinal stereotactic treatments.

It is clear from Table 2 of OARs that the shift plans showed a vast variation with an increase in its dosimetric parameter values from the actual base plan and the maximum variations was seen among the maximum shift plan and was mainly seen with the critical organs like the spinal cord, lt Parotid, right and left cochlea, mandible, right and left optic nerve. In a study, it was reported that for the spinal cord the maximum radiation dose that can be applied is 45Gy (Matzinger et al., 2009). The unique feature of this study from other studies is that, all the possible effective modes of shift errors and its dosimetric impact on the patient was studied. The study emphasized the importance of making image guidance a daily practice in radiotherapy treatment which is not followed in many centers as their protocol since they have to compromise with their patient work load. Image guidance is a mandatory procedure on daily basis in radiotherapy treatment for the precise treatment delivery and optimum results. Any amount of minimum to a maximum error in the patient setup will lead to a major missing of target and more dose exposure to critical organs which is very evident from the dosimetric results of the present study. In contrast to the studies of Tonndel et al., (2018) who didn’t demonstrate any advantage over daily imaging compared to weekly imaging, this study results dosimetrically reported the advantage of using daily imaging in radiotherapy which has to be mandatorily implemented in all radiotherapy centres for quality treatment.

Author Contribution Statement

The authors confirm contribution to the paper as follows: Study conception and design: Hridya V T, data collection analysis and interpretation of results: Hridya V T and Aswathi Raj, draft manuscript preparation: Hridya V T. All authors reviewed the results and approved the final version of the manuscript.

Acknowledgements

Ethical Standards

These articles don’t contain any studies on human participants or animals by any of the authors.

Data Availability

Research Data are not shared

Conflict of Interest

All authors declare no conflict of interest.

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