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Volumetric modulated arc therapy: a dosimetric comparison with dynamic IMRT and step-and-shoot IMRT

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Abstract

Aim: The aim of this study was to compare volumetric modulated arc therapy (VMAT) with dynamic intensity-modulated radiation therapy (dIMRT) and step-and-shoot IMRT (ssIMRT) for different treatment sites.

Materials and methods: Twelve patients were selected for the planning comparison study. This included three head and neck, three brain, three rectal and three cervical cancer patients. Total dose of 50 Gy was given for all the plans. Plans were done for Elekta synergy with Monaco treatment planning system. All plans were generated with 6 MV photons beam. Plan evaluation was based on the ability to meet the dose volume histogram, dose homogeneity index, conformity index and radiation delivery time, and monitor unit needs to deliver the prescribed dose.

Results: The VMAT and dIMRT plans achieved the better conformity ($CI_{98\%} = 0.965 \pm 0.023$) and ($CI_{98\%} = 0.939 \pm 0.01$), respectively, while ssIMRT plans were slightly inferior ($CI_{98\%} = 0.901 \pm 0.038$). The inhomogeneity in the planning target volume (PTV) was highest with ssIMRT with HI equal to 0.097 ± 0.015 when compared to VMAT with HI equal to 0.092 ± 0.0369 and 0.095 ± 0.023 with dIMRT. The integral dose is found to be inferior with VMAT 105.31 \pm 53.6 (Gy L) when compared with dIMRT 110.75 \pm 52.9 (Gy L) and ssIMRT 115 38 \pm 55.1(Gy L). All the techniques respected the planning objective for all organs at risk. The delivery time per fraction for VMAT was much lower than dIMRT and ssIMRT. *Findings*: Our results indicate that dIMRT and VMAT provide better sparing of normal tissue,

homogeneity and conformity than ssIMRT with reduced treatment delivery time.

Introduction

Intensity-modulated radiation therapy (IMRT) is an advanced type of radiation therapy used to treat cancer and non-cancerous tumours. It uses advanced technology to manipulate photon beams of radiation to conform to the shape of a tumour.¹ The development in modern imaging procedures, such as ultrasound, computed tomography (CT), magnetic resonance imaging, positron emission tomography (PET) and PET/CT, and the advent of advanced delivery techniques have made it more reliable treatment modality for cancer patients.² The complex shapes achieved using IMRT are made possible by considering each radiation beam as multiple rays, or beamlets, and assigning different beam strengths to the individual rays.³ These beamlets treat small areas of tissue, called voxels, which are a cubic millimetre of space. The beamlets are designated to satisfy the predetermined dose specifications to the tumour site and surrounding normal tissues.⁴ By modulating both the number of treatment fields and the intensity within each field, there is a greater control of dose distribution around the target and the dose homogeneity within the target. Volumetric modulated arc therapy (VMAT) is more advanced technique than IMRT in which the gantry speed, multileaf collimator (MLC) leaf position and dose rate vary continuously during delivery. This study was undertaken to compare the plan quality, delivery efficiencies and performance of two different IMRT delivery techniques, dMLC and ssMLC, with VMAT in four major anatomical sites. This included head and neck, brain, rectal and cervical cancer which are most common tumour sites at our institution.

Historical background

After discovery of X-rays in 1895, radiation was being used for the treatment of various malignant diseases.⁵ From 1950s to the late 1980s, the approach to radiation therapy was based on two-dimensional (2D) approach. In 2D radiation therapy, plans were created manually and a single beam from one to four directions was used to be given.⁶ Beam collimation was done with the manually applied shielding blocks. Technological advances in image acquisition since the early 1990s have changed the practice of radiation therapy significantly, and radiation therapy transitioned from the 2D method to a three-dimensional conformal therapy which conforms the

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Figure 1. Multileaf collimators (MLCs).

high radiation dose with uniform intensity to tumour. The MLC system was developed, which allows precise shaping of the treatment beam to the target volume.⁷ Since the mid-1990s and early 2000s, there has been a development of an advanced form of radiation therapy called IMRT. IMRT can provide an improved dose distribution and increased dose homogeneity when compared with three dimensional conformal radiotherapy.⁸

Methods of Delivering IMRT Treatment

IMRT treatments are primarily delivered by linear accelerators (linacs) with MLC systems. MLCs consist of pairs of highly absorbing tungsten leaves that can block the incident radiation and move against each other on each sides of the treatment fields, in such way that produce irregular fields similar in shape to the target tumour as shown in Figure 1. MLCs are computer-controlled motor-driven devices that could produce intensity modulations.⁹ MLCs are mostly operated in two modes: dynamic and step–shoot mode.

Dynamic IMRT

In dynamic technique, the beam is kept on while the MLC leaves move to produce the desired intensity modulation throughout the treatment delivery. Starting with a certain field opening and then by narrowing or widening of MLC leaves, any field shape can be created with the varying intensity maps. Mostly by keeping one leaf stationary and moving the opposite leaf towards it, a wedge-shaped field can be obtained.¹⁰

Step-and-shoot IMRT

One of the most commonly used modes of operation is the stepand-shoot or segmental MLC technique in which the modulation of intensity of beam in a treatment field is created by the exposure of a series of MLC-shaped discrete segmental fields.¹¹ The radiation beam is turned off when the MLC leaves are moving from one field segment to another, and is turned on only when the leaves reach and stop at the designated segment positions.¹²

Intensity-modulated stereotactic radiotherapy

In 1994, Cyberknife was introduced for stereotactic radio surgery. The system consists of a small light weight X-band linac with six degrees of freedom supported by a robotic arm coupled with an orthogonal pair of X-ray cameras.¹³ The cameras are able to image any change in patient setup and target motion during treatment, which are then fed to the robot to deliver radiation accordingly.¹⁴ As Cyberknife does not have any fixed isocentre, radiation can be delivered in any direction in step–shoot manner.¹⁵

IMAT

Yu has developed an IMAT technique that uses the MLC dynamically to shape the fields as well as rotate the gantry in the arc therapy mode.¹⁶ The method is similar to the step-and-shoot in which each field is subdivided into subfields of uniform intensity and subfields are superimposed to produce the desired intensity modulation. However, the MLC moves dynamically to shape each subfield, while the gantry is rotating and the beam is on all the time.

VMAT

IMAT has been improved with the addition of variable gantry rotation speeds and dose rates, and was introduced as VMAT in 2007 to describe rotational IMRT delivered in a 'single arc'.¹⁷ VMAT can provide highly conformal dose distributions and can significantly improve the IMRT delivery efficiency. The faster treatments reduce the effects of intra-fractional motion on both tumours and organs, and, of course, the shorter treatment times also increase patient throughput. The high plan quality and fast treatment delivery of VMAT are attractive, and it has been widely applied to many disease sites.

Materials and Methods

More than 450 patients have been treated with IMRT at our centre since July 2014 to till date. For this study, we had selected four major subsets. These included: (1) brain-3, (2) head and neck-3, (3) rectal-3 and (4) cervical-3 cancer patients. IMRT and VMAT plans were generated for comparison on Monaco planning system version 3.1 (Elekta Ltd, Crawley, UK) for Elekta synergy linac with clinically relevant planning constraints. Plan was created using a 6 MV photon beam. Planning parameters were kept same for both IMRT and VMAT. Plans were generated in such a way that 95% of planning target volume (PTV) receives 100% of prescribed dose. Critical organ dose kept as low as possible at the same time does not exceed the tolerance dose.

Case 1: brain cancer

Figure 2 shows an axial plan of a patient with meningioma planned with ssIMRT, dIMRT and VMAT. Total prescribed dose was 50 Gy in 25 fractions. The tolerance doses for organs at risk (OARs) were given in Table 1.

Case 2: head and neck cancer

Patients with primary head and neck cancers were treated at our centre. Figure 3 shows an axial treatment plan of a nasopharynx.

Table 1. Planning objective for the critical structures

Normal structure	Radiation thresholds (Gy)	Normal structure	Radiation thresholds (Gy)
Brain stem	Max dose < 54 Gy	Larynx	Mean dose < 45Gy
Spinal cord	Max dose < 45 Gy	Parotid glands	Mean dose < 26 Gy or V_{30Gy} < 50% (at least one side)
Lens	Max dose < 5 Gy		
Optic nerve	Max dose < 55 Gy	External ear	Max dose < 35 Gy
Optic chiasm	Max dose < 54 Gy	Cochlea	Mean dose < 45 Gy
Eyes	Max dose < 50 Gy	Mandible	Max dose < 70 Gy
Rectum	D _{60%} < 40 Gy	Femoral head	$D_{15\%} < 35 \text{ Gy}$
Bladder	D _{35%} < 45 Gy		
Optic chiasm Eyes Rectum Bladder	Max dose < 54 Gy	Cochlea Mandible Femoral head	Mean dose < 45 Gy Max dose < 70 Gy D _{15%} < 35 Gy



Figure 2. The isodose distribution of meningioma patient planed by (a) ssIMRT, (b) dIMRT, and (c) VMAT. Yellow colour was for isodose curve of 47.50 Gy.





Figure 3. The isodose distribution of nasopharyngeal cancer planed by (a) ssIMRT, (b) dIMRT, and (c) VMAT. Brown colour was for isodose curve of 47.50 Gy.



Figure 4. The isodose distribution of rectal cancer planed by (a) ssIMRT, (b) dIMRT, and (c) VMAT. Brown colour was for isodose curve of 47.50 Gy.

Total prescribed dose was 50 Gy in 25 fractions for primary tumour volume and bilateral neck nodal volume. The tolerance doses for OARs were given in Table 1.

Case 3: rectal cancer

Total prescribed dose was 50 Gy in 25 fractions. Figure 4 shows an axial treatment plan of a patient treated with IMRT and VMAT. Critical organs were rectum, bladder and left and right femoral heads. The tolerance doses for OARs were given in Table 1.

Case 4: cervical cancer

Patients of cervical cancer were planned with ssIMRT, dIMRT and VMAT (Figure 5). Prescribed dose was 50 Gy in 25 fractions. The OARs were bowel, bladder, rectum and femoral heads. The tolerance limits for OARs are given in Table 1.

Evaluation tools

Evaluation of the plans was performed by means of standard dose volume histogram (DVH). For PTV, $D_{98\%}$ and $D_{2\%}$ (doses received

Table 2. Dosimetric results for the PTV

Parameter	ssIMRT (mean ± SD)	dIMRT (mean ± SD)	VMAT (mean ± SD)
D _{98%} (Gy)	47.87 ± 0.64	48·36 ± 1·03	$46{\cdot}01\pm2{\cdot}06$
D _{2%} (Gy)	$52{\cdot}74\pm0{\cdot}15$	52.50 ± 0.36	$52{\cdot}73\pm0{\cdot}38$
НІ	0.097 ± 0.015	0.095 ± 0.023	0.092 ± 0.0369
CI	0.901 ± 0.038	0.939 ± 0.01	0.965 ± 0.023
MU	893·16 ± 340·6	849·22 ± 191·5	868·70 ± 234·01
Beam ON time (minute)	11±5	6 ± 2	2 ± 1
Integral dose (Gy L)	115.38 ± 55.1	110·75 ± 52·9	105·31 ± 53·6

Table 3. Case1: head and neck

Parameter	ssIMRT (mean ± SD)	dIMRT (mean ± SD)	VMAT (mean ± SD)
Brainstem D _{max} (Gy)	$29{\cdot}11\pm1{\cdot}09$	$25{\cdot}81\pm1{\cdot}12$	24·56 ± 1·21
Spinal cord D _{max} (Gy)	$24{\cdot}84\pm1{\cdot}23$	24.52 ± 1.11	$24{\cdot}56\pm1{\cdot}19$
Left lens D _{max} (Gy)	5.71 ± 0.3	5.20 ± 0.5	4·8 ± 0·3
Right lens D _{max} (Gy)	4.5 ± 0.4	3.78 ± 0.5	4.02 ± 0.2
Right parotid D _{mean} (Gy)	16.3 ± 1.02	15.8 ± 1.01	14.9 ± 0.9
Larynx D _{mean} (Gy)	$44{\cdot}58\pm0{\cdot}4$	44.07 ± 0.5	40.18 ± 0.6

Table 4. Case2: brain

Parameter	ssIMRT (mean ± SD)	dIMRT (mean ± SD)	VMAT (mean ± SD)
Left lens D _{max} (Gy)	2.74 ± 0.83	$2 \cdot 23 \pm 0 \cdot 72$	1.85 ± 0.51
Right lens D _{max} (Gy)	2·76 ± 0·53	2.25 ± 0.42	1.84 ± 0.32
Brainstem D _{max} (Gy)	9.6 ± 10.23	9.5 ± 6.98	9.43 ± 6.86
Optic chiasm D _{max} (Gy)	8·1±8·23	7·6 ± 5·99	6·7 ± 5·67

Table 5. Case 3: rectal

Parameter	ssIMRT	dIMRT	VMAT		
i arameter	(mean ± 3D)	(mean ± 5D)	(mean ± 5D)		
Left femoral hea	ad				
D _{mean}	16.56 ± 2.36	14.41 ± 2.16	12.64 ± 2.10		
D _{15%}	21.77 ± 2.18	16.76 ± 2.09	16.06 ± 2.11		
Right femoral head					
D_{mean}	16.56 ± 2.46	14.28 ± 1.45	14.01 ± 1.34		
D _{15%}	21·77 ± 2·35	16.62 ± 1.42	17.17 ± 1.31		
Bladder					
D _{mean}	26.65 ± 0.61	28·34 ± 0·52	29·30 ± 0·26		
D _{35%}	37.23 ± 0.54	37·74 ± 0·48	37·76 ± 0·18		



Figure 5. The isodose distribution of cervical cancer planned by (a) ssIMRT, (b) dIMRT, and (c) VMAT. Brown colour was for isodose curve of 47.50 Gy.

by the 98% and 2% of the volume) were defined for minimum and maximum doses. The homogeneity of the treatment plans was expressed in terms of $(D_{2\%} - D_{98\%})/D_{50\%}$ [homogeneity index (HI)] according to International Commission on Radiation Units and Measurements 83.¹⁸ An HI of zero indicates that the absorbed-dose distribution is almost homogeneous. The conformity of the plans was measured with a conformity index, $CI_{98\%}$, defined as the ratio between the patient volume receiving at least 98% of the prescribed dose and the volume of the PTV.¹⁹ For OARs, the analysis included the mean dose (D_{mean}) and the maximum dose (D_{max}). The integral dose (ID) of radiation delivered to the whole patient body was defined as ID [Gy...L] = D (Gy)...V (L), where D (Gy) is the mean dose delivered to volume *V* (L) (where L—litre).²⁰ Doses relevant to OARs were evaluated according to the specific treatment sites.

Results and Discussion

All plans sufficiently respected the planning objectives and can be clinically accepted. Table 2 provide an overview of the numerical

findings from an average DVH analysis on PTV, which are reported as mean values \pm standard deviation (SD) to assess for the relative inter-patient variability. Dosimetric results for OAR for all the case sites were reported independently as the analysis was done for different sites (Tables 3–6). The results are very homogeneous within the different approaches with small SDs. Both delivery techniques, if considered from a clinical perspective, appear to be equivalent.

PTV

Target coverage was almost similar for all the techniques. VMAT plans resulted in a better conformity ($CI_{98\%} = 0.965 \pm 0.023$) than dIMRT ($CI_{98\%} = 0.939 \pm 0.01$) and ssIMRT plans ($CI_{95\%} = 0.901 \pm 0.038$). The dose inhomogeneity in PTV was higher for ssIMRT plans with HI equal to 0.097 ± 0.015 when compared with dIMRT 0.095 ± 0.023 and VMAT 0.092 ± 0.0369 . The average monitor unit (MU) (\pm SD) needed to deliver the dose of 200 cGy per fraction was $893 \cdot 16 \pm 340.6$ MU for ssIMRT, $849 \cdot 22 \pm 191.5$ MU for

Table 6. Case 4: cervical

Parameter	ssIMRT (mean ± SD)	dIMRT (mean ± SD)	VMAT (mean ± SD)
Left femoral he	ead		
D_{mean}	23.27 ± 1.15	$22{\cdot}27\pm1{\cdot}11$	22.46 ± 1.02
D _{15%}	$28{\cdot}67\pm1{\cdot}09$	$27{\cdot}96\pm1{\cdot}07$	$28{\cdot}09\pm1{\cdot}01$
Right femoral h	nead		
D_{mean}	$27{\cdot}43\pm1{\cdot}16$	23.72 ± 1.05	$24{\cdot}10\pm1{\cdot}02$
D _{15%}	$32{\cdot}97\pm1{\cdot}11$	$28{\cdot}96\pm1{\cdot}01$	27·87 ± 0·98
Bladder			
D_{mean}	30.51 ± 0.67	$28{\cdot}81\pm0{\cdot}56$	29·53 ± 0·45
D _{35%}	$41{\cdot}86\pm0{\cdot}58$	$39{\cdot}51\pm0{\cdot}51$	38·95 ± 0·42
Rectum			
D _{mean}	33·00 ± 2·04	29·97 ± 2·18	28·94 ± 2·14
D _{60%}	28.63 ± 2.11	23.52 ± 2.12	20·71 ± 2·13

dIMRT and 868.70 ± 234.01 for VMAT. The average beam on time for VMAT plans (2 ± 1) was found to be significantly lower when compared with dIMRT (6 ± 2) and ssIMRT (11 ± 5).

OARs

All the techniques satisfied the planning objective for all OARs. The different characteristics of patients prevent the possibility to present an average conclusion and therefore the analysis was done separately for all the cases. The results are displayed in Tables 3–6. This study demonstrates that treatment techniques differ in terms of the treatment delivery time and overall plan quality. In comparison with ssIMRT, dIMRT and VMAT performed better in sparing OAR, particularly the lens, parotid glands, spinal cord and brain stem.

Case 1

The D_{max} to the brain stem was reduced by 12% with dIMRT and 17% with VMAT. For spinal cord, the D_{max} was also reduced by 1.3% with dIMRT and 1.2% with VMAT. There was a remarkable decrease in maximum doses of VMAT to the lens. The average maximum dose to the lens was reduced by 12.8% with dIMRT and 14.6% with VMAT. The mean dose to the ipsilateral parotid (right parotid) was reduced by 3.1% with dIMRT and 9% with VMAT.

Case 2

The average maximum dose to the lens was reduced by 20.4% with dIMRT and 39.1% with VMAT. The D_{max} to the brain stem was reduced by 1% with dIMRT and 1.8% with VMAT. For optic chiasm, the D_{max} was also reduced by 6.3% with dIMRT and 18% with VMAT.

Case 3

The average mean dose to the femoral head was reduced by 14.3% with dIMRT and 21.8% with VMAT. Bladder mean dose was increased by 6% with dIMRT and 9.4% with VMAT.

Case 4

The average mean dose to the femoral head was reduced by 22.9% with dIMRT and 8.5% with VMAT. Bladder mean dose was reduced by 5.7% with dIMRT and 3.2% with VMAT. Also, the rectum mean dose was reduced by 9.6% with dIMRT and 13.1% with VMAT.

Discussion

This study compared VMAT with dynamic IMRT and step-andshoot IMRT for 12 patients of different sites. It indicated that given the same target dose coverage, VMAT is able to treat more efficiently with less damage to OAR. A study of head and neck cancer by Vanetti et al.²¹ demonstrated that, given the same coverage, arc therapy offered better protection to OAR compared to IMRT similar to our study. Cozzi et al.²² conducted a planning study comparing VMAT with five-field conventional fixed-field IMRT in eight patients with cervical cancer. The results show similar target volume coverage with improved homogeneity and conformity with VMAT. OAR sparing (bladder and rectum) was significantly improved with VMAT with lower mean doses similar to our study. The study of Canyilmaz et al.²³ shows that the OAR sparing with VMAT is better than IMRT in case of high-grade gliomas similar to our study. The study of Studenski et al.²⁴ also shows that the VMAT treatment delivery time is lower than the IMRT. The significant difference between VMAT and IMRT is the treatment delivery time.

There are some inherent limitations with this study. First, this study included very small sample size. To obtain more precise results and improve clinical decisions, studies with larger sample sizes are required. Second, it is extremely difficult to completely eliminate planner bias with the same strict planning objectives and calculation algorithms.

A major source of concern with VMAT and IMRT is the increase in low dose radiation to surrounding normal tissue, which potentially increases the risk of secondary malignancy.²⁵ It has been seen in a number of studies that induced risk of secondary malignancy with VMAT should be lower compared with IMRT as volume of normal tissues receives lower dose with VMAT.²⁶ Longer follow-up of patients treated with these techniques will be required to accurately quantify this risk. Finally, although there is evidence to show that VMAT gives better dose conformity and OAR sparing but cannot be considered as the universal solution for all clinical scenarios. Each case must be evaluated on an individual basis to select the most appropriate radiation technique that will give optimal results.

Conclusion

With the same PTV coverage, radiation therapy using VMAT and dIMRT can lower the dose to most OAR, minimise potential damage, and achieve better homogeneity and conformity compared with ssIMRT. ID to the patients with VMAT was also lower than the dIMRT and ssIMRT plans, which reduces the risk of secondary malignancy. The average treatment time for VMAT and dIMRT plans was shorter than that of ssIMRT, which can increase efficiency and relieve patient discomfort due to lack of movement for a prolonged time. **Acknowledgements.** The authors would like to acknowledge the team of Radiotherapy Department for their co-operation. We would also like to acknowledge RIMS administration for the facilities provided in the department.

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