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Analyzing the performance of ArcCHECK diode array detector for VMAT plan



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ABSTRACT

Aim: The aim of this study is to evaluate performance of ArcCHECK diode array detector for the volumetric modulated arc therapy (VMAT) patient specific quality assurance (QA). VMAT patient specific QA results were correlated with ion chamber measurement. Dose response of the ArcCHECK detector was studied.

Background: VMAT delivery technique improves the dose distribution. It is complex in nature and requires proper QA before its clinical implementation. ArcCHECK is a novel three dimensional dosimetry system.

Materials and methods: Twelve retrospective VMAT plans were calculated on ArcCHECK phantom. Point dose and dose map were measured simultaneously with ion chamber (IC-15) and ArcCHECK diode array detector, respectively. These measurements were compared with their respective TPS calculated values.

Results: The ion chamber measurements are in good agreement with TPS calculated doses. Mean difference between them is 0.50% with standard deviation of 0.51%. Concordance correlation coefficient (CCC) obtained for ion chamber measurements is 0.9996. These results demonstrate a strong correlation between the absolute dose predicted by our TPS and the measured dose. The CCC between ArcCHECK doses and TPS predictions on the CAX was found to be 0.9978. In gamma analysis of dose map, the mean passing rate was 98.53% for 3% dose difference and 3 mm distance to agreement.

Conclusions: The VMAT patient specific QA with an ion chamber and ArcCHECK phantom are consistent with the TPS calculated dose. Statistically good agreement was observed between ArcCHECK measured and TPS calculated. Hence, it can be used for routine VMAT QA.

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1. Background

Intensity modulation in radiation beam improves the treatment plan quality in terms of tumor control probability (TCP) and normal tissue complication probability (NTCP). Intensity modulation can be achieved by fixed gantry delivery as well as arc based delivery.¹ Yu² proposed intensity modulated arc therapy (VMAT) with dynamic MLC as an alternative to tomotherapy in 1995. Although the concept of VMAT was developed much earlier, it took more time for clinical implementation. This is because of limited scientific work, lack of appropriate and efficient planning algorithm, delivery technique and weak commercial interest. Several researchers developed different techniques to realize single arc VMAT.^{3–12} Karl Otto developed progressive resolution optimization (PRO) for single VMAT. The clinical applications and comparisons between existing delivery techniques of VMAT were studied and published in various research works.^{13–27} Stringent quality assurance (QA) is required for assessment of any new equipment and technique before its clinical implementation. VMAT delivery is complex in nature, it involved simultaneous changes in gantry speed, multileaf collimator speed and dose rate. VMAT patient specific QA is important to assess the coordination of these parameters and delivery accuracy. One of the novel equipment intended for patient specific QA is ArcCHECK phantom. This equipment is specially designed for VMAT delivery. More reliable, robust and simple method of patient specific QA is absolute point dose measurement using an ion chamber. Ion chambers carry the calibration traceable to the primary standard dosimetry laboratory (PSDL). The aim of this study is to evaluate performance of ArcCHECK diode array detector for the volumetric modulated arc therapy (VMAT) patient specific quality assurance (QA).

2. Materials and methods

2.1. ArcCHECK

The ArcCHECK, (Model 1220, Sun Nuclear, Melbourne, FL), is a 3-dimensional beam dosimetry QA system intended for the measurement of radiotherapy dose distributions that are delivered, as defined by a planning system, and compared to the dose distribution, as calculated by the planning system. It is a cylindrical water-equivalent phantom with a three-dimensional array of 1386 diode detectors, arranged in a spiral pattern, with 10 mm sensor spacing. The center of the phantom (15 cm diameter) is designed to accommodate various accessories such as a solid homogeneous core, a dosimetric core with ion chamber(s) or diode arrays, an imaging QA core, a core with heterogeneous materials for dose studies, etc. The ArcCHECK also features two inclinometers to measure the angle of rotation about the cylinder axis and to measure the tilt of the axis. A temperature sensor measures the ambient temperature of the detector area. Dose measurements from each sensor are updated every 50 ms; there is no time limit or dose limit for a measurement (1386 precision diode detectors [size 0.8 × 0.8 mm]) (Fig. 1).

2.2. Linear accelerator

The measurements were carried out on a Clinac iX linear accelerator (Varian Medical Systems, Palo Alto, USA), capable of generating 6 MV and 15 MV photon beams. Dose rates available for photon are 100 MU/min to 600 MU/min in steps of 100 MU/min for VMAT delivery the dose rate range 0–600 MU/min (continuous). It is equipped with Millennium 120 MLC with central 40 pairs of 5 mm and peripheral 20 pairs of 10 mm leaf width at the isocentre. By design round tip multi-leaf collimators are at the tertiary level.

2.3. Performance tests

2.3.1. Linearity and reproducibility of dose

Linearity of dose measurement of ArcCHECK was evaluated. Monitor units ranging from 1 MU to 500 MU were delivered to ArcCHECK for 10 cm × 10 cm jaw setting. A range of monitor unit (MU) was chosen to cover the clinical VMAT plans. Phantom response to 100 MU was normalized to calculate linearity of dose. Similar measurement was carried out with an ion chamber in water phantom for comparison. Reproducibility of dose delivered is checked for ArcCHECK. It is evaluated in terms of coefficient of variation (COV). Ten repeated dose measurements were observed for 100 MU and 10 cm × 10 cm jaw setting. Similar measurement was carried out with an ion chamber in water phantom for comparison.

2.3.2. Field size and dose rate dependence

Response of ArcCHECK for various field sizes was evaluated. Field sizes ranging from 3 cm × 3 cm to 25 cm × 25 cm were exposed to a dose of 100 MU, and respective dose measured by ArcCHECK recorded. All readings of ArcCHECK phantom were normalized to the response of 10 cm × 10 cm field size. Similar measurement was carried out with an ion chamber in water phantom for comparison. The dose response of ArcCHECK was measured with different dose rates ranging from 100 MU/min to 600 MU/min. This measurement shows the Dose rate dependence of ArcCHECK. Similar measurement was carried out with an ion chamber in water phantom for comparison.

2.4. VMAT Plan QA

2.4.1. Treatment planning

All the plans in this study were performed in Eclipse treatment planning system (TPS), version 10.0 (Varian Medical Systems, Palo Alto, USA) using 6 MV photon beam. A dose rate of 600 MU/min was opted for all the arcs used, but the final dose rate was decided by the optimization algorithm. All the plans were optimized by a progressive resolution optimizer 3 (PRO3, Second generation).²⁸ In this algorithm full arc is optimized at 178 control points progressively in four phases. At every iteration level it optimizes multi-leaf collimator (MLC) position and monitor unit (MU) weight within limitations (MLC speed, gantry speed, dose rate, and mechanical limits) of the delivery unit.⁴ During optimization, dose calculation is performed with a simplified multi resolution dose calculation (MRDC) algorithm. In order to mitigate the leakage due to tongue and groove effect, collimator rotated to 30° from the

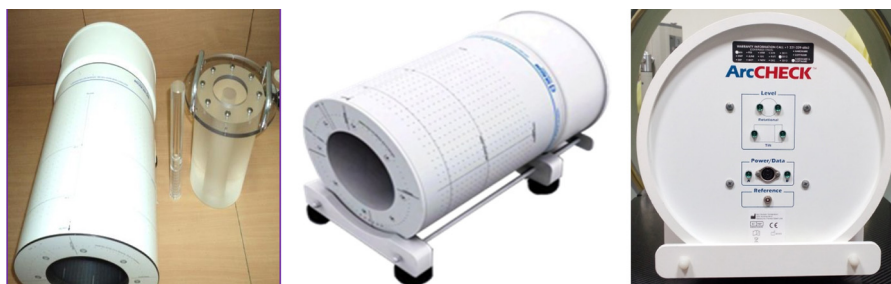


Fig. 1 – ArcCHECK phantom with cavity plug and chamber insert.

nominal angle. Number of full length arcs used ranged from 1 to 2 and the maximum number of partial ones was 3. Final dose calculations carried out with Analytical Anisotropic Algorithm (AAA). The Dose grid resolution for dose calculation was set to 2 mm. All the plan goals set to have 100 percent prescription dose to 95 percent of the target volume ($D_{95} = 100\%$ Rx dose). Objectives to other normal structures are set to clinically acceptable tolerance dose. Verification plans were created on ArcCHECK phantom using patient plans.

2.4.2. Phantom setup

A specially designed cradle supports the cylindrical ArcCA specially designed cradle supports the cylindrical ArcCHECK phantom. This stand was made up of low atomic number material to avoid beam attenuation. Electronic circuit of the phantom was placed away from the primary beam and in the direction away from the gantry. An inclinometer served to identify and correct the rotation (roll) and tilt (pitch). Coronal and sagittal lasers are matched with their corresponding line on the phantom. Then setup was verified by reading same SSD (~86.7 cm) on the phantom surface irrespective of the gantry angle.

2.4.3. Measurement

Calibrated CC 13S (IBA Dosimetry, Germany) ion chamber with Dose 1 (IBA Dosimetry, Germany) electrometer was used for absolute point dose measurements. Ion chamber was placed in the central core of the ArcCHECK at the isocenter of the linear accelerator. Diodes are kept at 2.9 cm below the surface of the ArcCHECK; this thickness of the buildup material is

equivalent to 3.3 cm of water. Dose maps were acquired at the diode level (89.6 cm). It is measured in cumulative mode for all the fields of the plan. Dose map and absolute point dose are measured simultaneously. This eliminates any minor difference in setup. Ion chamber readings were corrected for temperature and pressure variation.

3. Results

The most robust, reliable and simple technique for patient specific quality assurance is point dose measurement of IMRT plans. Ion chambers are recommended for absolute dose measurement of high energy photon beam. Point dose measurements are usually performed at one or more points in the phantom and compared with the calculation for the same measurement points. Twelve different patient plans were used in this study. Six head and neck plans, one breast plan, two esophagus plans, one abdomen plan and two pelvis plans.

3.1. Performance tests

3.1.1. Linearity and reproducibility of dose

The response of the ArcCHECK as a function of dose delivered (MU) shows that linearity is better than 0.25% for monitor units ≥ 5 MU. In lower monitor units linearity response is within 0.25–0.68%, except for 1 MU (3.5%). The slope of the trendline along the ArcCHECK phantom response curve is 0.998 and R^2 value is 1.000. The response observed with an ion chamber shows linearity better than 0.25% for monitor units ≥ 3 MU

Table 1 – Point dose measured and calculated at the isocenter/ion chamber.

Patient	Site	Ion chamber (ArcCHECK) (cGy)	TPS (cGy)	Dose diff (cGy)	% Dose diff
Patient 1	H & N	158.63	159	-0.37	-0.23
Patient 2	ESOPHAGUS	182.93	184.1	-1.17	-0.64
Patient 3	H & N	130.37	129.7	0.67	0.52
Patient 4	H & N	159.98	154.9	5.08	3.17
Patient 5	H & N	110.31	109.6	0.71	0.64
Patient 6	ABDOMEN	191.86	193.3	-1.44	-0.75
Patient 7	H & N	145.98	145.4	0.58	0.39
Patient 8	H & N	141.32	140.9	0.42	0.3
Patient 9	PELVIS	198.87	199.6	-0.73	-0.37
Patient 10	BREAST	158.21	159.2	-0.99	-0.63
Patient 11	PELVIS	173.14	174.1	-0.96	-0.55
Patient 12	ESOPHAGUS	169.66	170.4	-0.74	-0.44

Table 2 – Point dose measured and calculated at the diode plane along the CAX.

Patient	Site	ArcCHECK (Diode plane) (cGy)	TPS (cGy)	Dose diff (cGy)	% Dose diff
Patient 1	H & N	60.36	60.7	-0.34	-0.56
Patient 2	ESOPHAGUS	79.74	80.5	-0.76	-0.95
Patient 3	H & N	58.42	62.51	-4.09	-7
Patient 4	H & N	50.43	52.2	-1.77	-3.51
Patient 5	H & N	112.08	115.2	-3.12	-2.78
Patient 6	ABDOMEN	92.86	94.86	-2	-2.15
Patient 7	H & N	66.09	67.75	-1.66	-2.51
Patient 8	H & N	44.13	44.66	-0.53	-1.2
Patient 9	PELVIS	149.16	151.5	-2.34	-1.57
Patient 10	BREAST	81.64	79.54	2.1	2.57
Patient 11	PELVIS	126.93	129.2	-2.27	-1.79
Patient 12	ESOPHAGUS	59.01	59.8	-0.79	-1.34

(Fig. 2). In lower monitor units linearity response is within 0.63–0.84%. The slope of the trendline along the ion chamber response curve is 1.000, R^2 value is 1.000. Reproducibility observed for ArcCHECK phantom is 0.028% and for an ion chamber it is 0.038%.

3.1.2. Field size and dose rate dependence

Field factors calculated for the respective measurements from ArcCHECK and an ion chamber. Fig. 3 shows the normalized field factor calculated for ArcCHECK and an ion chamber. For field sizes less than 10 cm × 10 cm, ArcCHECK shows higher response than an ion chamber and vice versa for larger field

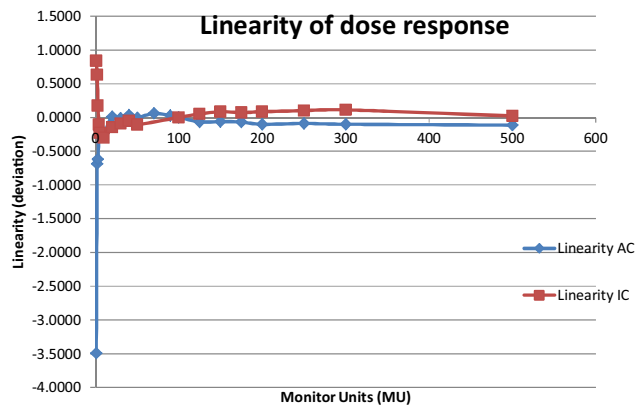


Fig. 2 – Linearity of dose response for Ion chamber and ArcCHECK phantom.

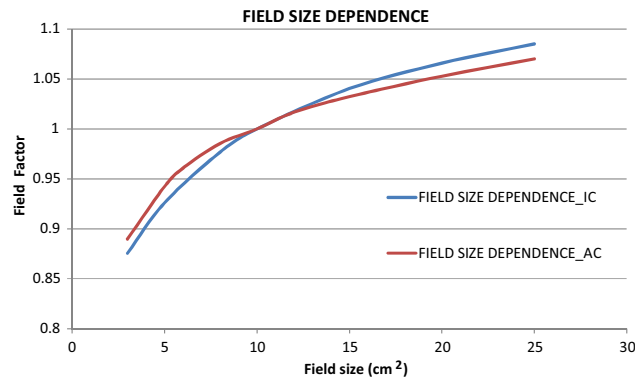


Fig. 3 – Field size dependent response of Ion chamber and ArcCHECK.

sizes. However, the deviation in the field factor is not more than 1.6% (SD 1.3%). Dose response of ArcCHECK phantom was identified at different dose rates measured. These measurements were normalized to the response at 300 MU/min. The dose responses observed for dose rates less than 300 MU/min were less than the response at 300 MU/min and vice versa for the larger dose rates. Maximum deviation observed for 600 MU/min is 0.39%.

3.2. VMAT plan QA

3.2.1. Ion chamber results

The dose values at the measurement point of an ion chamber were calculated and collected from Eclipse TPS. Ion chamber measured doses and TPS calculated doses are given in Table 1.

The point dose difference calculated between ArcCHECK measurement and TPS calculation was $0.50 \pm 0.51\%$. This satisfies the indispensable idea of intensity modulated beam QA guidelines.³⁰ Fig. 4a shows the linear relationship between measured and calculated doses and the slope of the trend line is equal to 1. Statistical analysis reveals a strong correlation between the ion chamber measured dose and the TPS predicted dose.

3.2.2. ArcCHECK results

Dose at a point (P_{AC}) where ArcCHECK diode plane intersects with the central axis was compared between TPS calculation and ArcCHECK measurement. This enabled us to make a direct comparison of point dose analogous to ion chamber measurement. No diode was present on the point of measurement (CAX intersection), so mean dose of adjacent diodes was taken as the measured dose. TPS calculated and measured doses are shown in Table 2.

The dose difference between ArcCHECK measurement and TPS prediction on the CAX ranged from 0.56% to 3.51%, except for one patient (7.00%). The mean dose difference in the central axis dose is 2.11% and standard deviation is 2.2%. The 95% confidence interval is from 0.9932 to 0.9995. In gamma analysis of dose/fluence map the mean passing rate was 98.53% for 3% dose difference and 3 mm distance to agreement (Table 3).

4. Discussion

Quality assurance becomes an integral part of complex radiation therapy delivery techniques. Patient specific quality

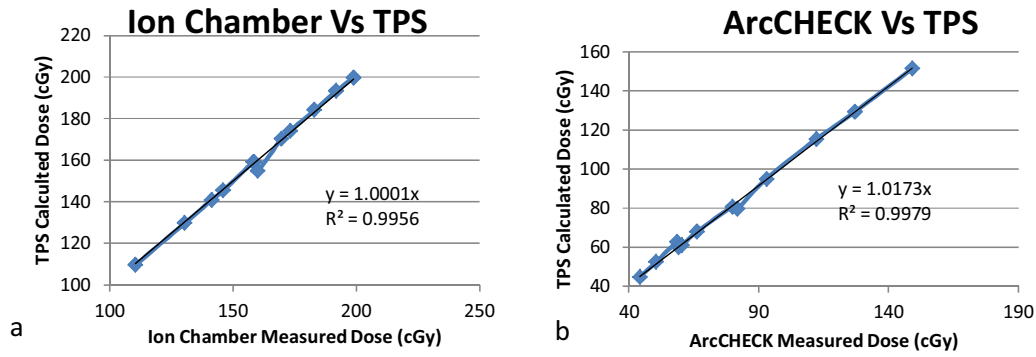


Fig. 4 – (a) Graphical plot of ion chamber measured dose and TPS calculated dose. (b) Graphical plot of ArcCHECK dose and TPS calculated dose.

assurance identifies the existence of manual error, mechanical inaccuracy and computerized faults. ArcCHECK phantom is an independent quality assurance tool. The diode detector’s response with respect to the gantry angle is corrected using Virtual inclinometer algorithm.³¹ Comparison was made of the device measured and TPS calculated dose for different patients studied. All the patients had field sizes <20 cm along the y-direction, so that the field fitted within the active area of the ArcCHECK phantom. Also the electronic circuits of ArcCHECK were protected against unwanted radiation damage. This phantom is very much suitable for hybrid plan verification¹⁵ where errors undetectable at 0° or any other

fixed gantry angle can be detected. Couch attenuation is accounted in the planning process by including couch modeling. Basic requirement of any detector used for hybrid plan verification is that its response should be independent of beam orientation, energy and dose rate. SNC Patient6® (SunNuclear Systems) software applied required correction to the diode response (Background correction, Angular correction, Field size correction). The spiral nature of diode placement virtually increases the measurement resolution (Fig. 5).

Tests relevant to the performance of the Arccheck phantom yielded a satisfactory outcome. Linearity of dose response of Arccheck phantom is very similar to that of Ion chamber.

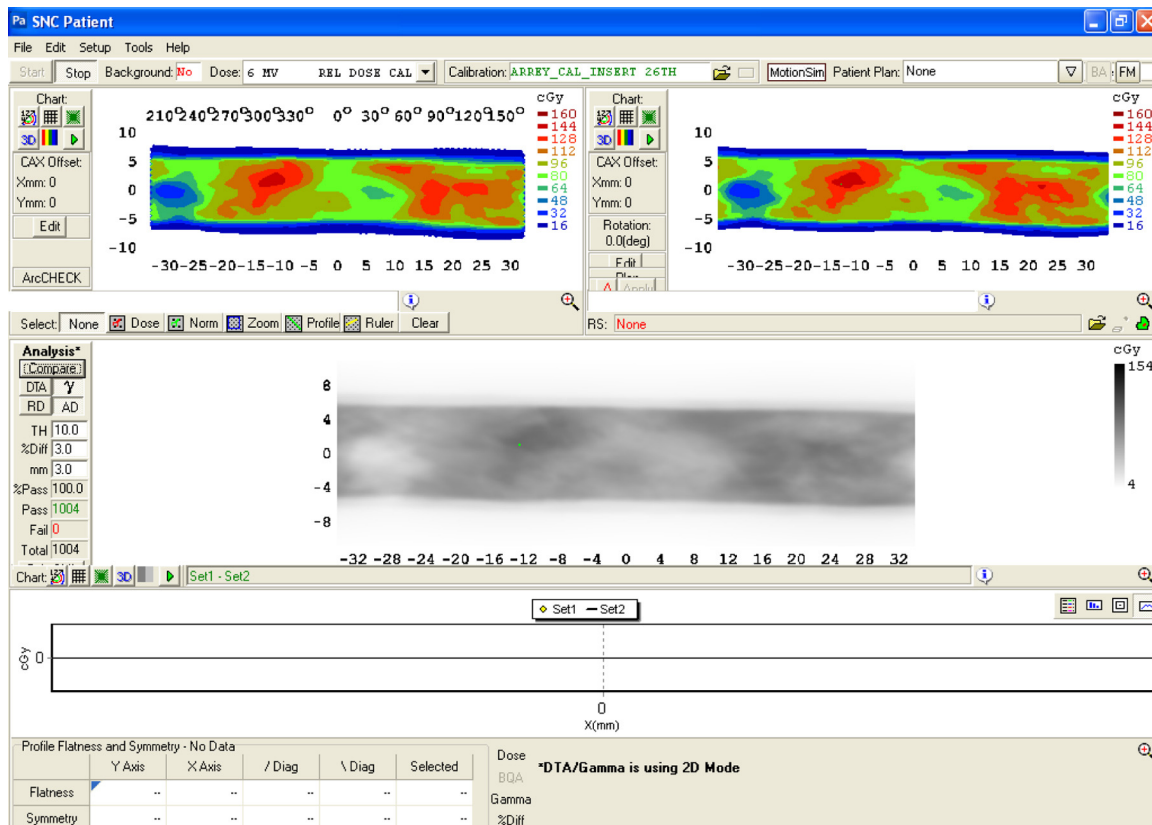
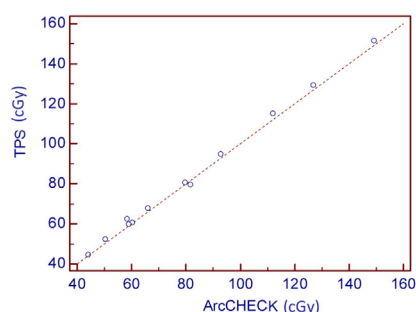


Fig. 5 – ArcCHECK measured does map compared with TPS calculated dose map in SNC Patient software (Sun Nuclear, Melbourne, FL).



Concordance correlation coefficient

Variable Y	TPS
Variable X	ArcCHECK
Sample size	12
Concordance correlation coefficient	0.9978
95% Confidence interval	0.9931 to 0.9993
Pearson ρ (precision)	0.9990
Bias correction factor C_b (accuracy)	0.9988

Fig. 6 – TPS calculated dose potted against ArcCHECK measured dose and CCC calculated.

Table 3 – Gamma analysis results.		
Patient	Site	Gamma (ArcCHECK) (DD 3%, DTA 3 mm)
Patient 1	H & N	99.1
Patient 2	ESOPHAGUS	97.8
Patient 3	H & N	98
Patient 4	H & N	96.1
Patient 5	H & N	99.9
Patient 6	ABDOMEN	100
Patient 7	H & N	98.9
Patient 8	H & N	97.2
Patient 9	PELVIS	99.3
Patient 10	BREAST	99.9
Patient 11	PELVIS	100
Patient 12	ESOPHAGUS	95.8

Monitor units of more than 5 MU show good linearity (within 0.25%). The deviation in linearity is more than 0.25% for lower MU, but all the clinical VMAT plans used in this study had MU in the range of 300–600 MU. Field size dependence test shows a closer agreement between the response of ArcCHECK phantom and Ion chamber. Absolute output of the linear accelerator was measured with default dose rate of 300 MU/min (Fig. 6). The default dose rate of the machine was used as reference dose rate to evaluate the dose rate dependence of ArcCHECK phantom. There are a number of QA devices and accessories available for VMAT patient specific QA. Many authors investigated the accuracy and implementation of these devices. Bedford et al.²⁹ evaluated Delta4 phantom (Scandidos, Uppsala, Sweden) for VMAT delivery and observed dose difference within 2.5%. Also, they demonstrated that Delta 4 phantom resulted in higher gamma passing rate than film. The absolute dose measured at the central axis diode plane is within $2.11 \pm 2.2\%$ of the TPS calculated dose. Concordance correlation coefficient (CCC) calculated between the ion chamber measured dose and the TPS calculated dose is 0.9996. These results again demonstrate a strong correlation between the absolute dose predicted by our TPS and the measured dose. CCC contains precision and accuracy of the data used. The advantage of using CCC over the Pearson’s correlation coefficient (r) (PCC) is that it indicates its components (accuracy and precision) in a single parameter (CCC = accuracy \times precision). The calculated accuracy coefficient and precision coefficient of the TPS prediction were 0.9999 and 0.9997, respectively. It means that the precision of the TPS model and calculation was 0.9999, and its accuracy was 0.9997. This method validates the

point dose calculation of the TPS software. Dose map and central axis diode dose were compared and evaluated with TPS predicted dose map and dose, respectively. Central axis diode dose measurement shows CCC of 0.9978.

5. Conclusion

This study evaluated the performance of ArcCHECK phantom and it is found to be clinically acceptable. The VMAT patient specific QA measurements with ion chamber and ArcCHECK phantom are consistent with the TPS dose calculation. Statistically good agreement was observed between the ArcCHECK measured dose and the TPS calculated dose. Hence, it can be used for routine VMAT patient specific QA. Also, it is a valid tool for acceptance and commissioning of the VMAT delivery technique.

Conflict of interest

None declared.

Financial disclosure

None declared.

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