Automated calculation of point A coordinates for CT-based high-dose-rate brachytherapy of cervical cancer

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Abstract

Purpose: The goal is to develop a stand-alone application, which automatically and consistently computes the coordinates of the dose calculation point recommended by the American Brachytherapy Society (i.e., point A) based solely on the implanted applicator geometry for cervical cancer brachytherapy.

Material and methods: The application calculates point A coordinates from the source dwell geometries in the computed tomography (CT) scans, and outputs the 3D coordinates in the left and right directions. The algorithm was tested on 34 CT scans of 7 patients treated with high-dose-rate (HDR) brachytherapy using tandem and ovoid applicators. A single experienced user retrospectively and manually inserted point A into each CT scan, whose coordinates were used as the “gold standard” for all comparisons. The gold standard was subtracted from the automatically calculated points, a second manual placement by the same experienced user, and the clinically used point coordinates inserted by multiple planners. Coordinate differences and corresponding variances were compared using nonparametric tests.

Results: Automatically calculated, manually placed, and clinically used points agree with the gold standard to < 1 mm, 1 mm, 2 mm, respectively. When compared to the gold standard, the average and standard deviation of the 3D coordinate differences were 0.35 ± 0.14 mm from automatically calculated points, 0.38 ± 0.21 mm from the second manual placement, and 0.71 ± 0.44 mm from the clinically used point coordinates. Both the mean and standard deviations of the 3D coordinate differences were statistically significantly different from the gold standard, when point A was placed by multiple users (p < 0.05) but not when placed repeatedly by a single user or when calculated automatically. There were no statistical differences in doses, which agree to within 1-2% on average for all three groups.

Conclusions: The study demonstrates that the automated algorithm calculates point A coordinates consistently, while reducing inter-user variability. Point placement using the algorithm expedites the planning process and minimizes associated potential human errors.

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Key words: brachytherapy, automation, dose calculation point, image-guided brachytherapy, cervical cancer.

Purpose

Locally advanced cervical cancer is routinely treated with concurrent chemotherapy and external beam radiation treatment (EBRT) followed by high-dose-rate (HDR) brachytherapy using a tandem and either ovoid pair or ring applicators [1]. For disease involving lymph nodes or parametrial extension, an additional EBRT boost is prescribed [2]. The American Brachytherapy Society (ABS) recommends reporting point A dose for treatment planning for HDR cervical brachytherapy [3]. When 3D imaging is used for image-guided brachytherapy (IGBT), magnetic resonance imaging (MRI) can be used to delineate the high-risk clinical tumor volume for volumetric planning [4,5,6]. When MRI scans of implanted HDR applicators are not readily available for target delineation, computed tomography (CT) is typically used to define the implant geometry, dose prescription points (point A), and organs-at-risk [3,7,8,9]. For dose calculation, the ABS recommends use of point A in their reports [1,3], located 2 cm perpendicularly to the tandem plane, and 2 cm plus the radius of the ovoids superiorly along the tandem from a line connecting mid-dwell positions of the ovoid pair. Even for volume-based planning, ABS [3], the Groupe Européen de Curiethérapie and the European Society for Radiotherapy and Oncology (GEC-ESTRO) [7] require reporting of dose to point A: “The dose delivered to point A should be reported for all brachytherapy applications regardless of treatment-planning technique” [3]. Defining
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A consistent dose point relative to the applicator would facilitate correlation of dose with clinical outcomes as clinics transition from 2D planning to volumetric 3D planning, according to GEC-ESTRO’s recommendation “to strictly follow the rules of a certain system with an enduring clinical tradition (for dose prescription)” [7]. With more advanced imaging technologies and brachytherapy modalities, volumetric prescription to the clinical target volumes is becoming prevalent for IGBT [9]. However, as of 2015, nearly 50% of centers continue to prescribe to point A despite the predominate use of CT-based planning [8]. Automated point coordinate calculations reduces dependence on either the user or the treatment planning system (TPS), thereby alleviating potential human errors in dose calculation point placement. Such errors were identified as one of the high-risk failure modes in HDR planning procedures [10,11,12,13]. Automated calculation of point A could also increase efficiency in HDR brachytherapy treatments of cervical cancer [11,14]. We have developed an automated point coordinate calculation algorithm for Fletcher-Suit-Delclos (FSD) tandem and ovoid (T&O) implants (Varian Medical Systems, Palo Alto, CA, USA) in order to expedite the process while simultaneously improving the precision of point localization.

Material and methods

**High-dose-rate brachytherapy treatment planning data**

This study utilized 34 CT scans with 2 mm slice thickness from 7 cervical cancer patients implanted with titanium FSD T&O for HDR treatment at our institution in 2014. High-dose-rate treatment plans were performed in BrachyVision v.11 (Varian Medical Systems, Palo Alto, CA, USA). To reduce applicator positioning uncertainties due to imaging artifacts, we utilize the solid applicators with pre-defined geometries provided by the manufacturer. The solid applicators are inserted into the CT scans, and the user translates and rotates them to match the implanted FSD T&O. All plans are generated with the 0.5 cm length 192Ir source (VariSource, Varian Medical Systems, Palo Alto, CA, USA) using a contiguous 0.5 cm step size and a 0.3 cm retraction from the distal tip to account for wire over-travel. The total source dwell length in the tandem is determined depending upon the cervical sleeve length inserted into the patient and the location of the cervical os. A standard source dwell length of 2 cm (i.e., 4 active dwell positions) is utilized in each ovoid. After insertion of the solid applicators, the user manually places points A by translating and rotating to multiple 3D views intersecting the applicators.

**Automated calculation of point A coordinates**

A stand-alone Matlab (Mathworks, USA) executable application calculates point A locations described in the previous section. The GUI-based application reads in a text file specifying the geometry of the implanted applicators, including the dwell positions in the applicators and the diameter of the ovoids, which is generated from the plan. The application first calculates the “tandem plane” in the sagittal plane, in which all tandem source dwell positions reside. The “tandem plane” is defined from the tip, middle, and most inferior source dwell positions in the tandem. The point A coordinate calculation begins by defining the ovoid central plane (Figure 1 – A), in which both ovoid dwell lines, assumed to be coplanar, are contained. The ovoid central plane is determined from any combination of three points in the ovoid dwell.
positions, one from one ovoid and two from the other ovoid. For our calculation, the 2nd dwell position in the left ovoid ($LO_2$), and the 1st ($RO_1$) and 3rd ($RO_3$) dwell positions in the right ovoid are selected. The vector normal to the ovoid central plane ($\hat{n}$, Figure 1 – C) is calculated as in Eq. 1.

$$\text{MIDOV}_1 = (RO_1 - LO_2), \text{MIDOV}_2 = (RO_3 - LO_2),$$

$$\hat{n} = \frac{\text{MIDOV}_1 \times \text{MIDOV}_2}{|\text{MIDOV}_1||\text{MIDOV}_2|}$$

(Eq. 1)

where $\text{MIDOV}_1$ and $\text{MIDOV}_2$ are the vectors connecting two dwell points between the ovoids. The “plane at ovoids’ top surface” (Figure 1 – B) is determined from the three points, $P_0$, $P_1$, and $P_2$ that are calculated from Eq. 2, where $R_{\text{coul}}$ is the radius of the ovoids.

$$P_0 = R_{\text{coul}} \hat{n} + LO_2, P_1 = R_{\text{coul}} \hat{n} + RO_1, P_2 = R_{\text{coul}} \hat{n} + RO_3$$

(Eq. 2)

The “point at the ovoids’ top surface”, $Ovtop$ (marked as $\times$ in Figure 1) is the point on the tandem intersecting with the plane at ovoids’ top surface. The point A is located 2 cm superiorly to $Ovtop$ and 2 cm laterally in the normal direction to the tandem plane.

To define $Ovtop$, the algorithm first finds the tandem segment vector, which crosses the plane at the ovoids’ top surface. The tandem segment vectors are defined between consecutive tandem dwell positions, $\vec{v}_i = t_{i+1} - t_i$. Here $t_i$ and $t_{i+1}$ are the $i$th and $(i+1)$th dwell positions from the most inferior source position in the tandem. Each tandem segment line is determined by the general parametric equation for a line, $t_i + \vec{v}_i \times s$, where $\vec{v}_i \times s$ is the difference vector between $t_i$ and $t_{i+1}$, and represents the distance from $t_i$ to a point on the line along $\vec{v}_i$. $Ovtop$ is the point where a tandem segment line intersects with the plane at the ovoids’ top surface. This surface is defined using the general equation for a plane, $P_0 + (P_1 - P_0)u + (P_2 - P_0)v$, where $P_0$, $P_1$, and $P_2$ are the three points as described above, and $u$, $v$ are parameters of a point in the plane formed by $P_1 - P_0$ and $P_2 - P_0$. To search for this intersecting point Eq. 3 is solved to determine $s$ for each tandem segment line.

$$t_i + (t_{i+1} - t_i)s = P_0 + (P_1 - P_0)u + (P_2 - P_0)v$$

(Eq. 3)

The tandem segment line, which crosses the plane is identified by $s \in [0, 1]$. The resulting crossing point, calculated from $t_i + \vec{v}_i \times s$, is assigned as $Ovtop$.

Once $Ovtop$ is determined, the algorithm locates the point on the tandem 2 cm superior from $Ovtop$. The “vector along the tandem”, $T_{\text{along}}$, is defined between $Ovtop$ and a tandem dwell position typically 2-2.5 cm above $Ovtop$, traces the superior-inferior direction of the tandem as shown in Figure 1 – D as a dotted line vector.

The algorithm finds the point 2 cm away from $Ovtop$ along $T_{\text{along}}$ and projects it back onto the actual tandem. To find this point, first the tandem segment vectors $\vec{v}_i$ are each projected onto $T_{\text{along}}$. The projection of $\vec{v}_i$ on $T_{\text{along}}$ is $\vec{v}_i \cos \theta_i$, where $\cos \theta_i$ is the cosine of the angle between $\vec{v}_i$ and $T_{\text{along}}$. The projection is computed and accumulated one segment vector at a time until the final ($i^\text{th}$) segment is reached. The final segment and its projection are computed as follows. The projections of individual segment vectors are accumulated as $d_{\text{sum}} = \sum_{i=1}^{3} |\vec{v}_i|$. until the next $\vec{v}_{i+1}$ projection to be added would cause the sum $d_{\text{sum}}$ to exceed 2 cm. For the final ($i+1)^\text{th}$ segment projection, the remaining distance $d_{\text{remain}} = 2 \text{ cm} - d_{\text{sum}}$ on $T_{\text{along}}$ is projected back onto the actual tandem to locate the corresponding point by solving $d_{\text{remain}} = |\vec{v}_{i+1}| \cos \theta_{i+1} + t_{i+1}$.

Finally, the points A are located 2 cm away from this point on the tandem described above in the direction normal to the tandem plane.

Data comparison and statistical analysis

To evaluate the accuracy and precision of the algorithm, a single experienced user (HK) retrospectively and manually placed points A for each plan to compare with the automatically calculated point coordinates. Because retrospective manual point placement by a single user avoids introducing inter-user variability and alleviates time constraints/pressure, points were considered the “gold standard”. To evaluate intra-user variability, the same user (HK) repeated point A placement in a second but independent session. To characterize inter-user variability within a time-pressured clinical setting, points A defined by various users for these clinical plans are also compared. Doses were calculated to check the dosimetric effects of point coordinate differences. The time span required to place points A manually was tabulated when the “gold standard” was placed, and used to estimate the increased efficiency that could potentially be gained by automatic point coordinate calculation.

Differences from the gold standard coordinates were calculated for automatically placed points, the repeated placements by a single user, the clinically used coordinates placed by multiple users, and used to assess the algorithm accuracy, intra-user variability, and inter-user variability, respectively. Coordinate differences in each dimension, in 3D magnitude, and associated dose ratios were compared for the three groups using a non-parametric, paired Friedman test, which does not require data to be normally distributed. Differences in a group were found, a paired Wilcoxon test was used to identify the pair of groups that differed.

To compare the variances of these coordinate differences among the same three groups, the nonparametric counterpart to the F-test, the Ansari-Bradley test, was used. All statistical tests were assessed at a significance level of 0.05.

**Results**

Table 1 summarizes the signed coordinate differences ($\Delta x$, $\Delta y$, $\Delta z$) and their 3D magnitudes ($\Delta 3D$) as well as dose ratios averaged over left and right points A from 34 HDR CT scans. All differences are referenced to the
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We developed a stand-alone application to automate point A coordinate calculation based solely on T&O implant geometry for HDR brachytherapy. This tool expedites planning and reduces user-dependence from the dose point localization process. The point placement time was reduced to less than 3-5 minutes, including generating the text input file and typing the point coordinates back into the TPS. The calculation time of the algorithm is less than one second. Because the initial planner and the second physicist checking the plan must both localize points A independently, the mean time added to the overall planning process is on the order of 10 minutes.

An advantage of automating point A coordinate calculations is to minimize the potential for human error and reduce the variability among various planners. Automation can improve efficiency and result in consistent point A localization even for planners who are not familiar with a TPS. In this study, the automatically calculated point A coordinates show submillimeter localization agreement to 0.35 mm on average, and dosimetric agreement to 1% on average compared to retrospective point insertion by a single experienced user (i.e., the gold standard).

While a single user localizing point A in a retrospective and non-rushed setting could mimic the high accuracy and precision of the algorithm, this was not true when multiple planners inserted point A into clinical plans in a time-pressured setting. Our study demonstrated that for multiple users, the 3D magnitude differences were significantly larger as was the variance of the coordinates in x, y, z, and 3D. The localization accuracy achieved by a single user in a retrospective setting could have been achieved at the expense of time, potentially leading to an overestimation of the time efficiency afforded by the point placement algorithm. Although the coordinate localization accuracy was decreased among multiple planners, the doses were not significantly impacted to within 1-2% on average. While not significant, differences in dose were up to 7% as shown in Table 1. In addition, we simulated the effects of 1 mm-shifts in L-R and S-I on dose and found that dose differences were larger, up to 5%, when point A was shifted laterally to the tandem compared to when it was shifted in S-I, for which differences were within 1%. The accuracy of both the coordinates and dosimetric agreement to the automatically placed points A demonstrates that the algorithm is able to perform robustly, thus reducing potential human errors arising from clinical time pressure. Studies on failure mode in high-dose-rate dosimetry identifies incorrect dose point placement as a mode with high severity [10]. Other studies have shown that failure modes associated with the highest risks originate most frequently from the treatment planning process [12] and from human failures [13].

Table 1. Summary of the signed coordinate differences (Δx, Δy, Δz) and their 3D magnitudes (Δ3D) as well as dose ratios averaged over left and right points A from 34 high-dose-rate computed tomography scans

| Groups compared to gold standard | Coordinate difference (mm); mean ± STD | Dose ratio (%); mean ± STD (range) |
|----------------------------------|----------------------------------------|-----------------------------------|
| Automatic calculation (algorithm accuracy) | L-R (Δx) A-P (Δy) S-I (Δz) 3D (Δ3D) | 99.9 ± 0.8 (97.7, 102.2) |
| Second manual placement (intra-user variability) | 0.02 ± 0.25 0.02 ± 0.21 0.02 ± 0.29 0.38 ± 0.21 | 100.1 ± 1.2 (96.8, 103.6) |
| Clinical plan (inter-user variability) | 0.00 ± 0.37* 0.03 ± 0.53* 0.06 ± 0.54* 0.71* ± 0.44* | 100.2 ± 2.0 (93.0, 104.7) |

The asterisk (*) indicates statistical significance (p < 0.05)
Automated point A calculation assumes that all ovoid central dwell positions are contained on the ovoid central plane (Figure 1 - A), and thus, any combination of three dwell positions should describe the same plane. This assumption could be violated in a clinical implant, particularly when the cervical anatomy of patient is atypical leading to asymmetric ovoid geometry. Non-coplanar ovoids can result in different central ovoid planes depending on which three dwell positions are selected to define the plane. To estimate the effects of non-coplanar ovoids and errors in localizing the ovoid tip on our methodology, point A was re-calculated with the dwell positions shifted by 3 mm along the ovoid lines, and also by swapping the combination of dwell positions, two dwells from the left and one from the right ovoids, instead of one from the left and two from the right. The uncertainties of the point A coordinates from these potential variations in ovoid localization was typically less than 0.1 mm. Because the algorithm compares the dwell positions within a curved tandem to the straight line drawn from the top of the ovoids plane, and a dwell position 2-2.5 cm superiorly in the tandem (i.e., dashed line shown in Figure 1), the algorithm did fail in 3-4 instances when the curved tandem approximated the straight dashed line (e.g., 15°).

The calculation algorithm could be easily applied to other TPS as long as a text file specifying the applicator geometry is available and can easily account for either more or fewer dwells in either the tandem or ovoids. The algorithm could also be extended to tandem and ring applicators due to the similarities in the definition of point A [3]. We recommend that TPS manufacturers incorporate such algorithms into their software system automating both calculation of point A coordinates and insertion into the 3D plan. While this would ensure that plans use correctly defined dose points without potential human errors in the data transfer between TPS and standalone point placement applications, verification by the end user would always be necessary. Such implementation would be the first step to standardization of dose point definition for brachytherapy.

Conclusions

By utilizing the automated dose point calculation algorithm, points are defined consistently between treatment plans without inter-user variability that might affect dosimetric evaluation and reporting. Not only does this provide compliance with the ABS recommendation of consistent point A placement and dose reporting [1,3] but it also addresses one of the failure modes identified in HDR treatment planning [10,11,12]. Automated calculation has the potential to reduce human errors in radiotherapy planning, particularly in clinics with a single planner for HDR brachytherapy [11].

Disclosure

Authors report no conflict of interest.

References

1. Nag S, Erickson B, Thomadsen B et al. The American Brachytherapy Society recommendations for high-dose-rate brachytherapy for carcinoma of the cervix. Int J Radiat Oncol Biol Phys 2001; 48: 201-211.
2. Yin G, Wang P, Lang J et al. Dosimetric study for cervix carcinoma treatment using intensity modulated radiation therapy (IMRT) compensation based on 3D intracavitary brachytherapy technique. J Contemp Brachytherapy 2016; 8: 221-232.
3. Viswanathan AN, Thomadsen B, American Brachytherapy Society Cervical Cancer Recommendations Committee, American Brachytherapy Society. American Brachytherapy Society consensus guidelines for locally advanced carcinoma of the cervix. Part I: general principles. Brachytherapy 2012; 11: 33-46.
4. Pottér R, Fidarova E, Kirisits C et al. Image-guided adaptive brachytherapy for cervix carcinoma. Clin Oncol (R Coll Radiol) 2008; 20: 426-432.
5. Viswanathan AN, Erickson BA. Three-dimensional imaging in gynecologic brachytherapy: a survey of the American Brachytherapy Society. Int J Radiat Oncol Biol Phys 2010; 76: 104-109.
6. Pavamani S, D’Souza DP, Portelance L et al. Image-guided brachytherapy for cervical cancer: a Canadian Brachytherapy Group survey. Brachytherapy 2011; 10: 345-351.
7. Pottér R, Haie-Meder C, Van Limbergen E et al. Recommendations from gynaecological (GYN) GEC ESTRO working group (II): concepts and terms in 3D image-based treatment planning in cervix cancer brachytherapy-3D dose volume parameters and aspects of 3D image-based anatomy, radiation physics, radiobiology. Radiother Oncol J Eur Soc Ther Radiol Oncol 2006; 78: 67-77.
8. Grover S, Harkenrider MM, Cho LP et al. Image Guided Cervical Brachytherapy: 2014 Survey of the American Brachytherapy Society. Int J Radiat Oncol Biol Phys 2016; 94: 598-604.
9. Otter S, Franklin A, Ajaz M et al. Improving the efficiency of image guided brachytherapy in cervical cancer. J Contemp Brachytherapy 2016; 8: 557-565.
10. Wilkinson DA, Kolar MD. Failure modes and effects analysis applied to high-dose-rate brachytherapy treatment planning. Brachytherapy 2013; 12: 382-386.
11. Damato AL, Devlin PM, Bhagwat MS et al. Independent brachytherapy plan verification software: improving efficacy and efficiency. Radiother Oncol 2014; 113: 420-424.
12. Mayadev J, Dieterich S, Harse R et al. A failure modes and effects analysis study for gynecologic high-dose-rate brachytherapy. Brachytherapy 2015; 14: 866-875.
13. Wadi-Ramahi S, Alnajjar W, Mahmood R et al. Failure modes and effects analysis in image-guided high-dose-rate brachytherapy: Quality control optimization to reduce errors in treatment volume. Brachytherapy 2016; 15: 669-678.
14. Damato AL, Lee LJ, Bhagwat MS et al. Redesign of process map to increase efficiency: Reducing procedure time in cervical cancer brachytherapy. Brachytherapy 2015; 14: 471-480.