Assessing cumulative dose distributions in combined external beam radiotherapy and intracavitary brachytherapy for cervical cancer by treatment planning based on deformable image registration

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Background: This study aimed to validate the feasibility of deformable image registration (DIR) in assessing the cumulative dose distributions in combined external beam radiotherapy (EBRT) and intracavitary brachytherapy (ICBT) for cervical cancer.

Methods: This retrospective study included 23 patients with stage IIB disease treated with combined EBRT to the whole pelvis (50.4 Gy in 28 fractions) using an intensity-modulated radiotherapy technique with 6-MV X-ray, followed by three-dimensional (3D) ICBT (28 Gy in 4 fractions). Tumor gross target volume at diagnosis (GTV-Tinit), tumor gross target volume before brachytherapy, high-risk clinical target volume (HR-CTV), intermediate-risk clinical target volume (IR-CTV), and parametrium and organs at risk were recontoured on computed tomography images of EBRT and ICBT, respectively. The dose-volume parameters were also determined. The DIR results were reviewed using MIM Maestro (Reg Review) and modified by function (Reg Refine). To evaluate the accuracy of DIR, DIR-based cumulative dose-volume histogram (DVH) parameters and simple DVH parameter addition were compared using Wilcoxon rank-sum tests.

Results: The cumulative dose distributions of EBRT and four ICBT sessions were successfully illustrated using DIR. The mean tumor diameters were 68.35 cm³ at diagnosis and 29.63 cm³ at ICBT initiation. The mean tumor regression was 56.6%. The median minimum dose covering 90% (D90) of HR-CTV, GTV-Tinit, IR-CTV, and parametrium were 69.58±4.94, 68.81±7.98, 59.28±3.78, and 60.97±1.1 Gy_{α/β}=10, respectively, for DIR and 69.11±5.68, 68.49±8.62, 58.89±3.59, and 61±1.49 Gy_{α/β}=10, respectively, with conventional simple DVH parameter addition. No statistically significant differences in dosimetric parameters were observed between the two methods.

Conclusions: Although there were limitations in the DIR accuracy, DIR-based dose accumulation was significantly beneficial in visually showing the cumulative dose distribution in the target area to clinicians in combined radiotherapy for cervical cancer in routine clinical practice.

Keywords: Cervical cancer; radiotherapy; deformable image registration (DIR); brachytherapy

Submitted Feb 25, 2020. Accepted for publication Aug 21, 2020.
doi: 10.21037/tcr-20-1196

View this article at: http://dx.doi.org/10.21037/tcr-20-1196
Introduction

The 2019 National Comprehensive Cancer Network (1) guidelines for Clinical Practice of Cervical Cancer 2019 (1st edition) recommend parametrial boosting in select cases with bulky parametrial/pelvic side wall disease after initial whole pelvic radiation, which has been added in the principle of radiotherapy. In such cases, the cumulative dose distributions of clinical targets should be determined and the dose-volume histograms (DVHs) calculated to identify the cumulative dose-volume associations for the target volumes and organs at risk (OARs) in combined external beam radiotherapy (EBRT) and intrauterine brachytherapy (ICBT). However, the accurate evaluation of cumulative dose distributions at the targets in the combined EBRT and ICBT is still a challenge for clinicians, specifically for patients with uncontrolled or recurrent tumors who require secondary radiotherapy.

Brachytherapy is an irreplaceable radiation treatment for cervical cancer. Recently, the field has transitioned from point- to volume-based planning (2), allowing for improved target volume and OAR visualization and enhanced dose conformity (3). When patients receive EBRT, brachytherapy is usually initiated toward the latter part of treatment. Sufficient primary tumor regression at this time permits satisfactory brachytherapy apparatus geometry. However, because of the different dose profiles and fractionations of ICBT, organ movement, and changes in target volume, the calculation of DVH parameters for the target volume and OARs may be difficult when assessing cumulative dose distributions with EBRT in conventional treatment planning systems. Several studies have determined the effect of deformable image registration (DIR) on cumulative OARs such as bladder and rectum DVH parameter summation for brachytherapy fractions (4-6). They demonstrated that several factors affect the systematic underestimation of dose. Specifically, the complete overlap of the DVH parameter hot spots may not always be accurate, and DIR in cervical cancer brachytherapy leads to lower average OAR DVH parameters (5,6). To date, the clinical effect of DIR on cumulative OAR DVH parameter summation between brachytherapy fractions is also controversial. In combined EBRT and BT treatment, van Heerden et al. used structure-based DIR to evaluate cumulative DVH parameters for cervical cancer in the bladder and rectum (7). Xu et al. applied the full parameter addition methods with respect to the image registration-based methods for evaluating combined EBRT and ICBT (8).

Their research proved that DIR could serve as an acceptable plan evaluation tool for clinical purposes in combined EBRT and BT treatment. In this study, we focused on the evaluation of cumulative dose distributions on targets and DVHs in combined radiotherapy for cervical cancer using a commercially available DIR algorithm (MIM Maestro®) (9) in clinical practice. We present the following article in accordance with the STROBE reporting checklist (available at http://dx.doi.org/10.21037/tcr-20-1196).

Methods

Patients and treatment

Twenty-three patients having anteflexed or upright uterus with stage IIB disease previously treated with combined ICBT and EBRT were randomly selected for this retrospective study. All patients received EBRT to the whole pelvis including the parametrium (50.4 Gy in 28 fractions) using the intensity-modulated radiotherapy (IMRT) (Varian, 1X, Pinnacle 9.8 plan system, USA) technique with 6-MV X-ray followed by three-dimensional (3D) ICBT (28 Gy in 4 fractions). Computed tomography (CT) [Philips, Brilliance BigBore (slice thickness, 5 mm)]-based 3D image-guided brachytherapy (Elekta, Nucletron microSelectron V3, Oncentra version 4.3 plan system, Sweden) was performed for the first fraction of ICBT. The other three fractions used the same initial CT-based brachytherapy planning. Since the ICBT and EBRT doses were delivered continuously in each fraction, their physical doses were summed directly, assuming the same biological effectiveness. The total equieffective prescription for a total point A dose was 90 Gy/α/β=10 (assuming an α/β ratio of 10 Gy). ICBT was performed weekly for 4 consecutive weeks using high-dose rate brachytherapy. In all cases, tandem and ovoid with sufficient vaginal packing were used to ensure appropriate placement and planning. At least 6 Gy was prescribed to the high-risk clinical target volume (HR-CTV) minimum dose covering 90% of the tumor volume (D90) in each ICBT session. The dose constraint was 75 Gy/α/β=10 in the minimum doses to the highest irradiated 2 cc area of the rectum (D2cc). The dose prescriptions and target coverages were modified based on the dose constraints for D2cc of 90 Gy/α/β=10 to the bladder and 70 Gy/α/β=3 to the rectum according to Groupe Européen de Curiethérapie and the European Society for Radiotherapy & Oncology (GEC-ESTRO) recommendations (10).

The study was conducted in accordance with the
Declaration of Helsinki (as revised in 2013). The study was approved by the Human Investigation Committee of Tianjin Medical University Cancer Institute and Hospital, Tianjin, China (No. bc2019029). Written informed consent for the scientific use of the clinical data was obtained from all patients.

**Computed tomography simulation**

This study requested that patients collect urine 1 hour before undergoing CT imaging for EBRT treatment planning to maintain the relative horizontal position of the uterus. We used the initial planning CT image with a 5-mm slice thickness per patient rather than per fraction.

**Target and organ-at-risk definitions**

The CTV of EBRT comprised the cervical tumor gross target volume at diagnosis (GTV-Tinit), the entire uterus, the parametrium, at least the upper half of the vagina, and the pelvic lymph nodes (including the common, external, and internal iliac and presacral lymph nodes). The planned target volume of EBRT included the CTV plus a 7-mm safety margin. For ICBT, CT contour of the HR-CTV, intermediate-risk CTV (IR-CTV), cervical tumor gross target volume before brachytherapy, parametrium, rectum, and bladder were adapted based on the CT-standardized contour guidelines (11). Based on the findings of gynecologic examinations at diagnosis and brachytherapy, the HR-CTV was delineated. Next, the IR-CTVs were generated by adding a 1-cm expansion around the HR-CTV, modifying the disease extent at diagnosis with a comparison to the initial extension in either the clinical diagram or initial magnetic resonance imaging (MRI) scan, and modifying the volume by deleting contours extending into the bladder, sigmoid, or rectum. Table 1 shows the details of the parametrical borders.

**Image registration**

Image registration was performed using MIM Maestro version 6.7.10 (MIM Software Inc., Cleveland, OH, USA) (12). CT image datasets from ICBT were rigidly fused by matching structures on CT images from IMRT. The rigid registration was determined to be “as good as possible” based on bone alignment. DIR was performed to accumulate doses of four fractions of ICBT with the same dose of the initial planning CT image per fraction. The results of the DIR were carefully reviewed using the MIM Maestro function in Reg Review and modified using the Reg Refine function. Figure 1 shows the schematics and workflow. Patients were required to defecate before being treated to avoid gas artifacts. Rectal dilatation was confirmed by treatment planning CT of EBRT and ICBT. If gas is found in the rectum during ICBT, gas drainage is performed to reduce rectal expansion. Subsequently, CT is performed again.

**Dose-volume evaluations**

The radiotherapy doses were converted into biologically equivalent doses in GyEQD2 using the linear quadratic model with α/β =10 Gy for tumor tissues and 3 Gy for normal tissues (13). To evaluate the combination of IMRT and ICBT for each plan generated, we calculated the median value of all plans based on the dose values received by 90% of the target volume (D90), the dose received by 100% of the target volume (D100), and the volume receiving 100% of the prescription dose (V100) compared to those for conventional simple DVH parameter addition, according to the EMBRACE II study (14) (Table 2). “Simple straight addition” was performed with the rigid image registration based on straight addition where there will be one final D90.

**Statistical analyses**

Wilcoxon rank-sum tests were used to investigate the differences between DIR-based dose accumulation and DVH parameter addition with regard to D90, D100, and D100.
Image of 1st fraction of ICBT

Whole uterus as region of interest

EBRT image

Lock local registrations

Rerun the deformable registration

Figure 1 Reg refine-based correction of deformation.

Table 2 Median D100, D90, and V100 values of the GTV-Tinit, HR-CTV, IR-CTV, and parametrium (α/β=10 Gy)

| Parameter      | DIR            | Simple addition | P value |
|----------------|----------------|-----------------|---------|
| GTV-Tinit      |                |                 |         |
| D100 (Gy)      | 61.89±3.53     | 61.55±4.15      | 0.330   |
| D90 (Gy)       | 68.81±7.98     | 68.49±8.62      | 0.301   |
| V100 (%)       | 30.27±4.44     | 30.03±4.61      | 0.057   |
| HR-CTV         |                |                 |         |
| D100 (Gy)      | 60.45±3.79     | 60.25±3.71      | 0.144   |
| D90 (Gy)       | 69.58±4.94     | 69.11±5.68      | 0.52    |
| V100 (%)       | 35.56±4.96     | 35.28±4.91      | 0.117   |
| IR-CTV         |                |                 |         |
| D100 (Gy)      | 54.16±4.01     | 53.99±3.93      | 0.456   |
| D90 (Gy)       | 59.28±3.78     | 58.89±3.59      | 0.073   |
| V100 (%)       | 28.92±6.13     | 28.92±6.13      | 0.616   |
| Parametrium    |                |                 |         |
| D100 (Gy)      | 55.71±2.73     | 55.51±2.27      | 0.692   |
| D90 (Gy)       | 60.97±1.1      | 61±1.49         | 0.648   |

GTV-Tinit, GTV at diagnosis; HR-CTV, high-risk clinical target volume; IR-CTV, intermediate-risk clinical target volume; V100, volume receiving 100% of the prescription dose; D90, dose received by 90% of the target volume; D100, dose received by 100% of the target volume.
V100 deviations for GTV-Tinit, HR-CTV, IR-CTV, and parametrium, respectively. Wilcoxon signed-rank tests were performed for each of the parameters mentioned above to disclose systematic differences between DIR-based accumulation and DVH parameter addition. Analyses were performed using the International Business Machines Corporation Statistical Package for the Social Sciences Statistics for Windows version 19.0.

**Results**

**Cumulative doses**

The cumulative dose distributions, comprising EBRT and four sessions of ICBT, were successfully illustrated using DIR (Figure 2). The HR-CTV was covered with 70–80 Gy\(^{\alpha/\beta=10}\). The parametria were covered with 50–60 Gy\(^{\alpha/\beta=10}\).

**Gross target volume and DVH parameters**

Among the 23 patients included in the present study, the mean tumor diameters were 68.35 cm\(^3\) at diagnosis and 29.63 cm\(^3\) at ICBT initiation. The mean tumor regression was 56.6%. The mean ± standard deviation (SD) HR-CTV D90 and D100 of the GTV-Tinit, IR-CTV, and parametrium were 69.58±4.94, 61.89±3.53, 54.16±4.01, and 55.71±2.73 Gy\(^{\alpha/\beta=10}\), respectively, for DIR and 69.11±5.68, 61.55±4.15, 53.99±3.93, and 55.51±2.27 Gy\(^{\alpha/\beta=10}\), respectively, for the conventional simple DVH parameter addition (Figures 3-5). No statistically significant differences were observed in the dosimetric parameters between the
two calculation methods (Table 2). No systematic differences were observed between DIR-based accumulation and DVH parameter addition (P=0.11).

Discussion

DIR is now accepted as a plan evaluation tool for clinical purposes at several radiation oncology centers not only for automatic contouring but also for diagnostic image registration and dose accumulation. The quality of DIR is dependent on image quality, user skills, and software algorithm. Several centers also use an Excel sheet-based EQD2 dose composite recommended by the American Brachytherapy Society. In these applications, these registrations must be characterized in phantom studies and also checked on a per-patient basis around regions in which deformable registration guided the clinical decisions (12).

DIR accuracy is of utmost importance in evaluating the DIR-based cumulative dose distributions and DVHs of the tumor and OARs. All selected cases in the present study had anteflexed or upright uterus. The uterus became horizontal after the insertion of the uterine tube in the ICBT. Considering the comfort of patients without general anesthesia in every ICBT session, the patients were only required to collect urine during ICBT to stabilize bladder size and position. With these methods, the cumulative images by rigid fusion were based on the bony structure (Figure 2).

However, in brachytherapy, the target volume and parts of the OARs could move together with the applicator after the applicator's implant (15). As the applicator changed its position relative to the bony structures, it is essential to match the region of interest, including the target volume and the parts of the OARs moving together with the applicator rather than only matching the brachytherapy applicator or bony structures (16). Therefore, the use of a proper deformation registration method based on a rigid registration improved fusion accuracy with automatic tools in planning systems (17).

Contouring and reconstruction performed in the same image series allowed the dose to targets and organs to be directly calculated without any image fusion. However, the fusion uncertainties could translate into absorbed dose calculation when several image series were combined. For example, contouring was performed in one image series and reconstruction in another. Tanderup et al. reported that uncertainties resulted in a miscalculation of DVH dose parameters by typically 4% to 6% mm$^{-1}$ of fusion error between CT and MRI (18). van Heerden et al. (19) recently used DIR to determine the absorbed dose distribution of daily image-guided adaptive EBRT and brachytherapy. The cone-beam CT of the first EBRT fraction and the brachytherapy planning MRI were registered by DIR. The cumulative dose to 2 cm$^3$ with the highest dose (D 2 cm$^3$) from EBRT and brachytherapy to the bladder and rectum were calculated and compared to those for direct addition, assuming a uniform EBRT dose (UD). The differences between UD and DIR were less than 3.9 Gy$_{EQD2}$. Our study only used CT images with the same field of view and slice thickness for fusion. The DIR-based DVH parameter values did not differ significantly from those derived from the conventional simple DVH parameter addition (Figure 3). According to the Groupe Européen de Curiethérapie and the GEC-ESTRO recommendations (10), the conventional simple DVH parameters for the target volumes and OARs in EBRT and each ICBT session were simply added to estimate the cumulative DVH parameters.

Mayr et al. reported that the tumor could shrink from 60% to 80% of the pre-therapeutic tumor volume during the first 2–3 weeks of EBRT (20). The patient’s anatomy at the time of brachytherapy with the applicator in situ differed significantly from that at EBRT. In this study, the mean tumor regression was 56.6% when brachytherapy
was initiated after 3–4 weeks of EBRT. Although pelvic organs, such as the bladder, rectum, and intestine, differed significantly in movement and filling, the processed image fusion met the requirements for clinical target dose assessment by planning systems.

The previous analyses proved that ICBT, with the new target concept based on HR-CTV and IR-CTV and combined with the pear concept used in the past for ICBT, had a significantly high local control rate (21-23). Thus, it was appropriate and feasible to evaluate the extension of the dose into the parametria with cervical cancer (24,25). Nomden et al. compared the treatment planning of four cervical cancer patients with different sizes of HR-CTV between four centers (23). In their study, the median HR-CTV D90 was 68.9 Gy, whereas the mean ± SD HR-CTV D90 of our result was 69.58±4.94 Gy in the treatment plan with equieffective prescription for a total point A dose of 90 Gy\(\alpha/\beta\)=10. The results did not meet the demand for HR-CTV \(\geq 80\) Gy\(\alpha/\beta\)=10 for large or poorly responding tumors (1), for a total point A prescription dose (as recommended in the guidelines) of 80 Gy\(\alpha/\beta\)=10. The pear-shaped isodose without optimization implied a large non-conformality region compared to the HR-CTV definition. Although the doses to the HR-CTVs and OARs could be regulated through optimization, large residual or asymmetrically located tumors could not be managed by ICBT alone (26). A satisfactory D90 was not achieved for extensive tumors with an optimized ICBT approach alone because the main role of optimization was to reduce the D2cc to the OARs (27). Thus, clinicians must always consider the risk of local recurrence due to insufficient doses caused by these techniques. Furthermore, adequate evaluation of ICBT with EBRT is required for further parametrial-boosting radiotherapy.

According to the GEC-ESTRO recommendations (10), IR-CTV included the GTVres and residual pathological tissue. Margins were added for suspected adjacent residual microscopic disease in the direction of potential residual microscopic spread considering the natural anatomical borders. The selection and contouring of the IR-CTV were based on macroscopic tumor extension at diagnosis (GTV-Tinit), which had a high risk of local recurrence due to insufficient doses caused by these techniques. Furthermore, adequate evaluation of ICBT with EBRT is required for further parametrial-boosting radiotherapy.

Conclusions

Although it showed no significance in all DVH parameters between DIR and straight addition, DIR-based dose accumulation was significantly helpful to visually show the cumulative dose distribution in the target area to clinicians in combined radiotherapy for cervical cancer in routine clinical settings.

Acknowledgments

Funding: None.

Footnote

Reporting Checklist: The authors have completed the STROBE reporting checklist. Available at http://dx.doi.org/10.21037/tcr-20-1196

Data Sharing Statement: Available at http://dx.doi.org/10.21037/tcr-20-1196
Conflicts of Interest: All authors have completed the ICMJE uniform disclosure form (available at http://dx.doi.org/10.21037/tcr-20-1196). The authors have no conflicts of interest to declare.

Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013). The study was approved by institutional Ethics Review Board of Tianjin Medical University Cancer Institute and Hospital (No. bc2019029) and informed consent was taken from all the patients.

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Cite this article as: Zeng J, Chen J, Zhang D, Meng M, Zhang B, Qu P, Pang Q, Wang P. Assessing cumulative dose distributions in combined external beam radiotherapy and intracavitary brachytherapy for cervical cancer by treatment planning based on deformable image registration. Transl Cancer Res 2020;9(10):6107-6115. doi: 10.21037/tcr-20-1196