Analysis of clinical utilization of ring applicator for combined intracavitary/interstitial image-guided brachytherapy treatment in Chinese patients with locally advanced cervical cancer

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Abstract

Purpose: To analyze the results of clinical utilization of ring applicator for combined intracavitary (IC)/interstitial (IS) image-guided brachytherapy (BT) technique in locally advanced cervical cancer (LACC), and to explore the application conditions of IC/IS BT approach with this applicator for Chinese patients.

Material and methods: IC/IS BT was performed in total of 95 LACC patients, who already received external beam radiotherapy. In forty-three of these patients, utilization of ring applicator in brachytherapy was done, including IC/IS BT approach with ring applicator for 65 fractions. Clinically optimized IC/IS BT plan with ring applicator (IC/IS treatment) and additionally generated optimized plan without needle use (IC research) were designed, respectively. Dosage differences in target regions and organs at risk (OARs) were analyzed between both the plans. Four planes were selected in the longitudinal direction of uterine cervix and divided the 360° area around the tandem into eight equal sectors. Tumor target features and prescription dose distributions on different planes were described.

Results: The usage rate of ring applicator with most suitable ring diameter of 26 mm was 45.3% in Chinese cervical cancer patients. The high-risk clinical target volume (HR-CTV) D_{90}/D_{98} and intermediate-risk CTV (IR-CTV) D_{90}/D_{98} in IC/IS treatment were significantly higher than IC research. As compared with IC research plan, the D_{2.0 \text{ cm}^3} of bladder and rectum in IC/IS treatment were significantly lower (p < 0.05). There was no significant difference in D_{2.0 \text{ cm}^3} of sigmoid and small bowel between two plans. When the target volume was larger than 36 cm³, IC alone had an inadequate coverage of target.

Conclusions: Nearly half of Chinese patients with LACC could be treated with the ring applicator. The application conditions of IC/IS BT approach with ring applicator for Chinese patients were achieved. Combined IC/IS image-guided BT based on magnetic resonance imaging (MRI) planning is clinically feasible in Chinese patients.

Key words: cervical cancer, intracavitary, interstitial, brachytherapy, ring applicator.

Purpose

The standard treatment for locally advanced cervical cancer includes cisplatin-based concurrent chemo-radiotherapy, followed by brachytherapy. For brachytherapy treatment, different techniques such as intracavitary brachytherapy (ICBT), interstitial brachytherapy (ISBT), or hybrid intracavitary/interstitial brachytherapy (IC/IS BT) are being increasingly applied for cervical cancer. ICBT has been previously described in the literature, and with this technique, excellent tumor control can be expected for small size tumors [1]. The American Brachytherapy Society recommendations for high-dose-rate (HDR) brachytherapy for cervical carcinoma revealed that the eligibility criteria for undergoing ISBT were bulky lesions, narrow vagina, inability to enter cervical os, more than proximal third of parametrium extension, and lower vaginal extension [2]. In order to produce better target coverage while optimally sparing organs at risk (OARs), the combined IC/IS compatible computed tomography/magnetic resonance imaging (CT/MRI) applicator has a remarkable contribution. An analysis from the RetroEMBRACE study found that us-

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ing combined IC/IS brachytherapy improves therapeutic ratio and local control in large tumors without adding treatment-related late morbidity [3]. Although the ring applicator is one of the most widely used applicators for IC/IS technique in the world, there are only few departments to implement IC/IS BT for cervical cancer using ring applicator, with less experience in China. However, the clinical feasibility of this applicator on a larger Chinese group of patients with cervical cancer have yet to be demonstrated. The aim of this present study was to investigate the preliminary results of clinical use and to analyze the indication of ring applicator in the treatment of Chinese patients with locally advanced cervical cancer.

Material and methods

Patients

A total of 95 patients with cervical squamous cell cancer (aged, 49.80 ±9.65 years; Karnofsky Performance Score [KPS] ≥ 70) were enrolled into the study. These subjects were all Chinese of Han origin, inhabiting the Northeast area of China. The patients were admitted to our hospital between November 2014 and October 2016. Initial locoregional staging included a clinical evaluation with stage IB2 (9 patients), stage IIA2 (23 patients), stage IIIA (30 patients), stage IIIB (22 patients), and stage IVA (11 patients) was performed by a well-trained gynecologic surgeon and radiation oncologist according to the 2009 Federation International of Gynecology and Obstetrics (FIGO) classification.

All patients received pelvic external beam radiotherapy (EBRT) of 45 Gy, with three-dimensional conformal radiotherapy (3D CRT) technique (18 patients) or intensity-modulated radiotherapy (IMRT) technique (77 patients) and 4 fractions of 3D IGBT (once a week) with 7 Gy of prescribed dose (PD). Brachytherapy was initiated at the 5th week into EBRT (nearly 36-40 Gy). Then, 4 fractions of 3D IGBT were performed once a week. In addition, we applied 2 fractions of brachytherapy at the 8th week in certain cases in order to control the total treatment time within 8 weeks. Chemotherapy was given during EBRT, with intravenous cisplatin 40 mg/m² once a week, for 5-6 weeks in all patients.

Equipment and applicator

Micro-Selectron HDR Iridium-192 (192Ir) brachytherapy (Nucletron, an Elekta company, Elekta AB, Stockholm, Sweden), Oncentra v.4.3, Nucletron (Veenendaal, The Netherlands), and 3.0T magnetic resonance imaging (Siemens, Erlangen, Germany) were used in the study. Interstitial ring CT/MRI applicator set (Nucletron, an Elekta company, Elekta AB, Stockholm, Sweden) is a gynecological applicator for brachytherapy procedures. This applicator is available in different diameters (26, 30, and 34 mm). Guide holes in the ring tube allow placement of MRI-compatible IS ProGuide needles using the ring tube as a needle template. A maximum of 9 ProGuide needles can be placed.

Imaging, contouring, treatment planning, and dose reporting

In each patient, an assessment of tumor position by MRI examination and clinical gynecological examination was implemented before first brachytherapy session. BT procedure with ultrasound-guided insertion of ring applicator under general anesthesia was performed. During the operation, the implant depths was adjusted according to the results of ultrasound guidance. After insertion, the needles were locked, and the rectal dose monitor was placed in the rectum. In the subsequent treatments, appropriate applicator should be chosen according to tumor shrinking condition.

Magnetic resonance images of every brachytherapy fraction were obtained by 3.0T MRI scanner. Images were acquired according to our institutional protocol, which includes axial, sagittal, and coronal T2-weighted turbo spin-echo images, with 4.5 mm slice thickness and a balanced steady-state free precession scan with a slice thickness of 1.5 mm for applicator reconstruction purposes.

Based on the recommendations of the Gynecologic Groupe European de Curietherapie and the European Society for Therapeutic Radiology and Oncology (GEC-ESTRO), the target regions and OARs in patients were contoured according to MRI images.

The target regions included high-risk clinical target volume (HR-CTV) and intermediate-risk clinical target volume (IR-CTV), and OARs involved bladder, rectum, sigmoid, and small bowel. Target volume delineation was based on the MRI image before and after external beam radiation.

On the basis of MRI images, the 3D IC/IS BT plan was optimized for HR-CTV D90/D90 (minimum doses delivered to 90% and 98% of the target volume) and OAR D2.0 cm³ (minimum doses to 2 cm³ volumes of the OAR that receive the maximum dose). Dose calculation and reporting were based on the total (EBRT + BT) biologically equivalent dose in 2 Gy fractions (EQD2). Linear quadratic (LQ) model for radiation damage repair was used with α/β = 10 Gy for the tumor target and α/β = 3 Gy for the OARs. The EQD2 of EBRT and BT was then added to evaluate the optimized plan regarding the dose-volume-histogram (DVH) constraints. The goal of combined EBRT and BT was set to achieve HR-CTV D90 ≥ 85 Gy, IR-CTV D90 ≥ 60 Gy, bladder D2.0 cm³ ≤ 80 Gy, and rectum/sigmoid/small bowel D2.0 cm³ ≤ 70 Gy. The DVH and the reference doses were used to design the treatment plan.

Study analyses and experimental design

Patients enrolled in this study who were treated with combined IC/IS BT with the ring applicator were defined as IC/IS treatment group. Each fractional treatment plan of IC/IS treatment group generated a simulation plan by changing the source dwell times of the tandem only, leaving the interstitial part aside, and it was defined as IC research group. This optimized IC research plan was not used clinically. The aim dose of target and the limit dose of OAR in IC research plan were the same as the criterion of IC/IS treatment group.
Each needle application was evaluated, and the number of needles, positions through the ring, and insertion depths were registered. These needle choices were compared with the location of HR-CTV. This location was determined in para-transverse planes (perpendicularly to the tandem). A total of 4 parallel planes were observed: plane I at the level of ring tube surface, plane II at the parallel layer of 1 cm above ring tube surface, plane III at the parallel layer of 2 cm above ring tube surface (ICRU point A), and plane IV at the parallel layer of 1 cm above plane III (3 cm above ring tube surface). We divided the 360° area around the tandem into eight equal sectors. The dividing line from 1 to 8 in clockwise direction was numbered and the first line at 0 o’clock direction was marked (Figure 1).

We evaluated 65 fractions of IC/IS treatment group plans and the same fractions of IC research plans with the following criteria: for each direction of every plane analyzed, if the area of the target volume was covered by the prescription isodose curve, it was defined as qualified; if not, it was interpreted as unqualified. Therefore, we obtained the “qualified” ratio in each direction of every plane and in this way, the “prescription dose coverage rate” in each plane and direction was analyzed.

The HR-CTV D90/D98 and IR-CTV D90/D98 of target were calculated in the IC/IS treatment and IC research plans, respectively. Moreover, D2.0 cm³ of OARs (bladder, rectum, sigmoid, and small bowel) were compared between the two treatment regimens. Tumor target features and prescription dose distributions on 4 planes were described.

Data processing and statistical analysis

Data were presented as mean ± standard deviation. The comparison of dosimetric differences between two groups was done using pair t-test. Statistical significance was set at $p < 0.05$ at two-sided level. The linear regression model was used for the HR-CTV D90 and target volume analysis. All statistical analysis was performed using SPSS version 17.0 (SPSS, Chicago, IL, USA).

Results

Treatment outcome

In forty-three of 95 patients with locally advanced cervical cancer, the application of ring applicator was performed. The ring applicator usage rate was nearly 45.26%. Forty-three cases were treated with 3D IGBT in 172 fractions, including implementation of IC/IS IGBT approach in 65 fractions.

The mean volume of HR-CTV was 48.2 ±27.2 cm³, and the mean volume of IR-CTV was 97.4 ±59.3 cm³.

Dosimetric comparison of target exposure and OARs between the two plans

Table 1 presents the dose parameters of target and OARs with ring applicator in the two investigated plans. The HR-CTV D90/D98 and IR-CTV D90/D98 in IC/IS treatment group were significantly higher than those in IC research group ($p < 0.05$). As compared with IC research plans, the D2.0 cm³ of bladder and rectum in IC/IS treatment plans were significantly lower ($p < 0.05$). There was no significant difference in the D2.0 cm³ of sigmoid and small bowel in the two plans. The total EQD2 dose of 43 patients treated with combined IC/IS BT was shown in Table 2.

Needles implantation

The total of 43 patients were treated by combined intracavitary/interstitial brachytherapy in 65 fractions. The total number of needles was 277, and the average number of needles used was 4.3 for each patient, with the average depth of 4.1 cm. Hole distributions in the ring applicator in our research are shown in Figure 2. The situation of each hole is shown in Table 3.
Regression fitting of power function for target volume and HR-CTV D_{90}

Regression fitting of power function with linear regression model as well as the relationship between target volume and HR-CTV D_{90} under the two plans (Figure 3) was analyzed. According to the EQD_{2} value of LQ model, when the EQD_{2} of HR-CTV D_{90} was 85 Gy in terms of four equivalent calculations, the physical dose of HR-CTV D_{90} was 7.13 Gy. When HR-CTV D_{90} dose value was 7.13 Gy, the target volume was 36 cm³ in ICBT curve. The result of regression curve analysis showed that it might be difficult for IC BT alone to meet the dose requirements of tumor target when its volume was equal to or greater than 36 cm³.

**Prescription dose coverage on different planes**

The prescription dose coverage rate in IC/IS treatment plans was 93.2%, which was significantly higher than that in IC research (69.7%). In IC research plans, the prescription dose could cover the target volume at 8 directions on plane I; the location of less than 60% of the prescription dose coverage were centered in the direction 3 (48.1%), 4 (44.4%), 6 (51.8%), and 7 (44.4%) on plane I, the maximum distance between prescription dose and tandem was 20 mm at these four directions; the location of less than 60% of prescription dose coverage were centered in the direction 3 (44.4%), 4 (48.1%), 6 (44.4%), and 7 (51.8%).

**Table 1. Comparison of dose parameters of target and OARs with ring applicator in two plans**

| Parameters | IC/IS treatment | IC research | t  | P-value |
|------------|-----------------|-------------|----|---------|
| HR-CTV D_{90} | 7.94 ±0.65 | 6.98 ±0.75 | 8.48 | < 0.001 |
| HR-CTV D_{98} | 6.66 ±0.75 | 5.85 ±0.66 | 7.27 | < 0.001 |
| IR-CTV D_{90} | 4.86 ±0.53 | 4.37 ±0.63 | 7.07 | < 0.001 |
| IR-CTV D_{98} | 3.75 ±0.58 | 3.34 ±0.69 | 3.99 | < 0.001 |
| Bladder D_{2.0cm³} | 5.18 ±0.66 | 5.61 ±0.50 | –6.71 | < 0.001 |
| Rectum D_{2.0cm³} | 3.83 ±0.60 | 4.05 ±0.49 | –3.05 | 0.005 |
| Sigmoid D_{2.0cm³} | 3.56 ±0.90 | 3.48 ±0.93 | 1.76 | 0.091 |
| Small bowel D_{2.0cm³} | 2.68 ±1.32 | 2.72 ±1.36 | –1.32 | 0.199 |

**Table 2. Dose-volume histogram parameters in 43 patients treated with combined IC/IS BT**

| Parameters | EQD_{2} dose (EBRT + BT) Mean ± SD (Gy) |
|------------|----------------------------------------|
| HR-CTV D_{90} | 90.33 ±9.20 |
| IR-CTV D_{90} | 67.89 ±3.59 |
| HR-CTV D_{98} | 80.17 ±7.45 |
| IR-CTV D_{98} | 62.15 ±4.11 |
| Bladder D_{2.0cm³} | 76.53 ±6.49 |
| Rectum D_{2.0cm³} | 64.25 ±6.76 |
| Sigmoid D_{2.0cm³} | 61.79 ±8.01 |
| Small bowel D_{2.0cm³} | 60.15 ±6.80 |

**Table 3. Total number of implanted needles and average insertion depth in 65 cases treated with ring applicator**

| Needle channel | Number of needles | Average depth (cm) |
|----------------|-------------------|--------------------|
| 4              | 24                | 4.0                |
| 5              | 58                | 4.2                |
| 6              | 55                | 4.3                |
| 7              | 17                | 2.9                |
| 8              | 55                | 4.2                |
| 9              | 51                | 4.3                |
| 10             | 17                | 4.0                |
on plane III, the maximum distance between prescription dose and tandem was 17.8 mm at these four directions; the location of less than 60% of prescription dose coverage were in the direction 6 (44.4%) and 7 (51.8%) on plane IV, the maximum distance between prescription dose and tandem was 17 mm at these two directions (Figure 4).

**Discussion**

The combined IC/IS BT can consider high-dose radiation and conformal dose distribution to target in brachytherapy procedure for locally advanced cervical cancer. The standardized applicators for IC/IS BT includes Utrecht applicator (tandem + ovoid + needles) and ring applicator (tandem + ring + needles). Because needles of ring applicator are parallel to tandem, it is necessary to have a model. When the model was converted to an ovoid for dosimetric comparison of this ring applicator in order to provide a reliable reference for its clinical practice. We analyzed the dosimetric comparison of this ring applicator in order to provide a reliable reference for its clinical practice.

Only 43 patients could tolerate the ring applicator, which was limited to the minimum size (diameter with 26 mm), and the usage rate of this applicator for combined IC/IS BT treatments was 45.26%. In our study, we found that the ring applicator was applied to patients with anteversion of uterus or higher vaginal stretch conditions.

According to dosimetric comparison of the two plans, the target dose in IC/IS treatment plans was significantly higher than that in IC research plans. Moreover, IC/IS treatment plans could reduce the D2.0 cm³ of rectum and bladder significantly, as compared with IC research plans. There was no significant difference in the dose of the sigmoid and small bowel in the two plans. This result was consistent with Fokdal et al. [3] that combined IC/IS BT to deliver significantly higher dose to the target area without increasing the dose to OARs.

In this study, the highest number of needles were implanted through the ring applicator at lateral and latero-posterior hole (Table 3). Similar results were found in previous studies [4,5] using Utrecht applicator, and another study performed with tandem/ring applicator (GammaMed, Varian).

The dose constraint of rectum was more of rigorous standard in this study, since the higher increase of D2.0 cm³ of rectum, the higher incidence rate of side effects. Dose effect relationships have been demonstrated for rectal morbidity [6,7]. Mazeron et al. found that D2.0 cm³ of rectum should be lower than 75 Gy [7], and when rectum D2.0 cm³ dose further descended to 65 Gy, the incidence of side effects was noted at below 10% [8]. When the rectum received an appropriate dose (60-70 Gy), only mild side effects occurred and included mainly fibrosis, telangiectasia, or inflammation [9]. The incidence of G2-G4 side effects was 8.6% in 3 years follow-up, when the D2.0 cm³ of rectum was between 60 to 65 Gy, which was the 1/3 of incidence when the rectum D2.0 cm³ was up to 75 Gy [7]. If the D2.0 cm³ of rectum was more than 75 Gy, the risk of rectal fistula was nearly 12.5%. Therefore, the appropriate rectal D2.0 cm³ dose limit for this study was 70 Gy.

Mazeron et al. found that the incidence of G2-G4 side effects in rectum and bladder was related to the D2.0 cm³. When D2.0 cm³ was less than 81.4 Gy, the incidence of adverse reaction of bladder (30%) was higher than that of the rectum [10]. As the radiation dose increased, the sensitivity of the bladder to the dose response decreased. In this study, the restriction dose of bladder was set as 80 Gy. If the HR-CTV D90 exceeded 87 Gy (EQD2), the 3-year local control results could be better than 95% [11]. In order to improve the local control, this study required the HR-CTV D90 to reach 85 Gy or more, with the HR-CTV D90 requirement appropriately increased, when there are high-risk factors to consider. For example, in case of stage III-IV cervical cancer patient, target volume > 30 cm³, treatment time > 55 days, and tumor diameter > 5 cm at diagnosis, etc., the HR-CTV D90 needs to be increased to 97, 92, 105, and 92 Gy, respectively [12].

In the present study, the physical dose of HR-CTV D90 obtained by the two plans was fitted to the regression curve of the target volume. According to EQD2 value in LQ model, when the EQD2 of HR-CTV D90 was 85 Gy in terms of four equivalent calculations, the physical dose of HR-CTV D90 was equal to 7.13 Gy. The result of regression curve analysis showed that it might be difficult for IC BT alone to meet the dose requirements of tumor target when its volume was equal to or greater than 36 cm³ in Chinese patients with locally advanced cervical cancer. Yoshida et al. used a cube as a model to investigate the HR-CTV volume threshold of IC BT alone [13]. It was considered that HR-CTV less than 4 × 3 × 3 cm³ (equivalent to 36 cm³) required image-guided IC BT and recommended IC/IS BT or IS BT for HR-CTV larger than this volume. This result was calculated using a cube as a model. When the model was converted to an ovoid for
volume calculation, this critical volume was only 18.8 cm$^3$, which was smaller than the results of this study. The reasons for this difference may be as follows. First, the HR-CTV in the Yoshida et al.’s study was obtained by the volume calculation formula and might have some differences from the direct measurement of clinical target volume in this study. Second, the shape of HR-CTV was different in the two studies. The HR-CTV studied by Yoshida et al. was a fully symmetrically distributed cube. However, the HR-CTV in this study was the actual target region, and the full symmetry of the shape was difficult to achieve. Third, the Yoshida et al.’s study selected fewer models with a total of 8 different HR-CTV volumes (8, 12, 27, 36, 64, 80, 96, and 112 cm$^3$). However, the number of patients enrolled in this research was large, and the difference in target volume was insignificant. The estimated value was fitted by the regression curve relationship between target volume and HR-CTV. This is also the main reason for the large difference between the two data. In the study of Yoshida et al., the tandem just passed the HR-CTV center, which was rare in actual clinical situation. Anderson et al. suggested that the conventional IC BT alone could meet the clinical requirements of HR-CTV D$_{90}$ for a < 40 cm$^3$ HR-CTV. When HR-CTV > 40 cm$^3$, IC BT combined with image guidance was required. When the volume of HR-CTV was further increased (median, 74 cm$^3$), the target coverage evaluated by HR-CTV D$_{90}$ in image-guided IC BT could reach only 80 ±16% (PD%), and IC/IS BT was recommended [14]. The result of volume threshold reported by Anderson et al. was greater than that estimated in our study, which might be related to higher doses of OAR in their research.

In this study, the target area coverage of IC/IS BT was much higher than that of IC BT alone (93.2% vs. 69.7%), which was related to better conformation of PD curve to the coverage of target in IC/ISBT insertion needles. It was
found that the location of coverage rate ratio less than 60% in IC research plans were concentrated in direction 3, 4, 6, and 7 on the plane II and III; direction 6 and 7 on the plane IV. In addition, on the III and IV planes, the PD curve formed by IC BT created approximately a regular circle. Plane I in this study was the parallel layer for upper surface of the ring tube. In clinical applications, larger diameter of ring applicator should be selected, so that the shape of ring tube and the cervix front-end could be as consistent as possible. In this case, both IC BT and IC/IS BT could produce good dose coverage of tumor target on plane I. At planes II and III, IC BT alone could not fully cover the dorso-lateral part of HR-CTV, which was related to the distribution of target area. Coia et al. reported that the irregular shape of target volume and paracervical excessive invasion were the main reasons to unsatisfactory dose coverage and insufficient dose distribution [15]. In our study, plane IV was located at the parallel layer of 3 cm above ring tube surface, and the dose coverage of IC BT was better than that in planes II and III. The possible reason might be that the height of some HR-CTV was less than 3 cm. It could be seen that there was a minimum dose contribution of the ring tube at the parallel layer of ICRU point A or above, and the radiation dose was likely to be insufficient in the direction of dorsal part of tumor target.

In the present study, when arbitrary directions in the lower quadrant (directions 3, 4, 6, and 7) of the plane II exceeded 20 mm, and the arbitrary directions of planes III and IV exceeded 17.8 and 17 mm, respectively, it was difficult to satisfy the coverage requirements of the target area by IC BT alone. However, Dimopoulos et al. evaluated their experience with tandem/ring/needles applicator for cervical cancer [16]. The IC/IS BT was selected on the condition that the unilateral tumor expanded more than 3.5 cm at the ring level, and/or more than 2.5 cm at ICRU point A level, and/or more than 2.2 cm at a distance of 3 to 4 cm from the upper ring surface. The differences between the anatomy and the ring applicator diameter applied might be the actual reasons for discrepancies in the above studies. In addition, the relatively strict dose limitation in our research was also an important factor.

Conclusions

The IC/IS BT technique with the ring applicator was chosen if IC BT alone was unable to produce sufficient coverage of target volume with the PD. This occurred in two clinical situations: 1. if HR-CTV was more than 36 cm³, and 2. if the cervical dorsal tumor extension exceeded 20 mm at a distance 1 cm cranial to the upper ring surface, or unilateral tumor extension exceeded 17.8 mm at the level of ICRU point A and 17 mm at a distance 3 cm cranial to the upper ring surface. Combined IC/IS image-guided BT based on MRI treatment planning with the ring applicator in situ is clinically feasible in Chinese patients with locally advanced cervical cancer, which need a reasonable selection of indications. Our experience with 95 patients has helped us to create institutional guidelines. In the near future, we expect that IC/IS image-guided BT in combination with the ring applicator will make an important contribution to the treatment of cervical cancer in China.

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Disclosure

The authors report no conflict of interest.

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