Therapeutic Benefit of Intracavitary-interstitial Brachytherapy in Cervical Cancer Patients with Small and Large High-risk Clinical Target Volume

Makbule TAMBAŞ

1Department of Radiation Oncology, University of Groningen, University Medical Center Groningen, Groningen-Netherlands

OBJECTIVE
We aimed to investigate the added value of interstitial brachytherapy (IS-BT) over classical intracavitary BT (IC-BT) in terms of target coverage and organ at risk (OAR) sparing among patients for whom an optimal dose distribution could not be provided without IS-ICBT and also to determine if the magnitude advantage provided by IS-BT is similar in patients smaller (<30 cm³) and larger (≥30 cm³) high-risk clinical target volume (CTV_RH).

METHODS
24 patients treated with IS-ICBT were included in this study. IS-BT was performed 76 of 93 BT fractions. For each patient, two additional IC-BT planning were created: (1) ICBT_Target-focused plan: The priority was adequate coverage of CTV_RH. Then, the OARs were spared as much as possible. (2) ICBT_OARs-focused plan: The priority was given to the OAR sparing. Then, highest CTV_RH coverage was tried to achieve within the allowed OAR dose limits. The IS-ICBT plans were compared with these two plans in terms of target coverage and OAR doses.

RESULTS
13 patients had large and 11 patients had small CTV_RH. In IS-ICBT plans, EQD2_10 CTV_RH D90 doses were significantly higher compared with ICBT_OARs-focused plans (Δdose: 10.5±6.2 Gy, p<0.001), whereas EQD2, OAR D2cc doses were significantly lower compared with ICBT_Target-focused plans (Average Δdose, bladder: 24.5±25.9 Gy [p<0.001], rectum: 7.6±9.7 Gy [p=0.001], sigmoid: 18.3±15.3 Gy [p<0.001]). There was no significant difference between patients with small and large CTV_RH in terms of ∆doses of both target and OARs.

CONCLUSION
IS-BT provides significant therapeutic advantage over IC-BT for patients both with small and large CTV_RH.

Keywords: Cervical cancer; CTVHR volume; interstitial brachytherapy; intracavitary brachytherapy.
MRIpostEBRT imaging was performed in the last week of EBRT to evaluate the patient's response and suitability for BT, which was planned within the 1st week after EBRT completion. The patient preparation, clinical workflow, and CT-guided needle insertion were described in detail in our previous study.[5] Briefly, BT was performed under sedoanalgesia. A CT scan was performed after applicator insertion, that is, CTpreneedle. Together with the MRIpostEBRT scan, this scan was evaluated by the radiation oncologist to decide needle indication, channels, and insertion lengths. If the IS-BT was indicated based on the tumor extension, applicator position and OAR location on the CT, the needles were inserted on the CT table and a second CT scan was performed after needle insertion (CTpostneedle). CTpreneedle and CTpostneedle were used to create BT plans for patients treated with IC-BT and IC-ISBT, respectively.

CT scanning was performed with a 1.25-cm slice thickness using the GEHC Discovery CT750 HD (Waukesha Wisconsin, USA). Three-dimensional BT planning was performed using the Oncentra Brachytherapy Planning System v4.5.3 (Elekta, Veenendaal, The Netherlands) after contouring of the residual gross tumor volume (GTVres), CTVHR, and OARs, including the bladder, rectum, and sigmoid on the CTpostneedle.[6-9] The plan was initiated by activating all source positions and was continued by manual optimization of the dwell times in the channels of the intrauterine tandem, ovoids, and needles.

All the treatment procedures reported in this study were a part of the routine clinical practice in the institution and were conducted after obtaining consent as relevant. The ethics committee deemed that additional informed consent for this study was not required, based on the Liv Hospital-Ulus Department of Radiation Oncology Medical Research Involving Human Subjects Act. However, all patients were informed that their data could be used for research purposes and that they could refuse consent for such use.

Dosimetry Goals
The summed biologically equivalent doses in 2-Gy fractions (EQD2) of EBRT and BT were calculated with α/β of 10 (EQD210) and 3 (EQD23) for CTVHR and OARs, respectively. The aims and limits of planning in the EMBRACE II protocol were used during plan optimization.[10]

Planning Without Needles
To determine the advantage provided by ISBT, plans without needles were created in total for the 76 IC-ISBT patients a larger CTVHR. Moreover, an increase by 10% in 3-year local control has been reported in patients with a larger high-risk clinical target volume (CTVHR ≥30 cm³) using IS-ICBT, with no increased toxicity.[1]

IS-BT not only facilitates coverage of the parametrial extension of the tumor with an adequate dose, but also accomplishes unacceptable OAR doses due to topography.[4] In our previous study, we have demonstrated the feasibility of the CT-guided needle insertion during combined IS-ICBT in patients with locally advanced cervical cancer using tandem-ovoid Utrecht applicator (Elekta, Veenendaal, The Netherlands). In addition to information provided by magnetic resonance imaging (MRI) in the last week of external-beam radiotherapy (MRIpostEBRT), OAR location and positioning of the tandem on the day of BT evaluated in the CT imaging taken after tandem insertion (CTpreneedle) was used to determine IS-BT indication, needle channels, and insertion lengths. This method is especially beneficial for patients with smaller CTVHR as unexpected indication may emerge based on CTpreneedle in these patients, whereas IS-BT indication is already determined based on MRIpostEBRT in patients with larger CTVHR. However, it remains unknown if advantage provided by the IS-BT in patients with smaller CTVHR is as great as for those with larger CTVHR.

Therefore, we aimed to investigate the added value of IS-BT over classical IC-BT in terms of target coverage and OAR sparing among patients for whom an optimal dose distribution could not be provided without IS-ICBT and also determine if the magnitude advantage provided by IS-BT is similar in patients smaller and larger CTVHR.

Materials and Methods

Patients and Treatment
Between May 2018 and January 2020, 74 patients with inoperable cervical cancer were evaluated for BT at Liv Hospital-Ulus Department of Radiation Oncology following EBRT, scheduled at 45-50.4 Gy/25-28 fr, and concomitant weekly cisplatinum (40 mg/m²). Among these, 32% (n=24) of the patients were treated with IS-BT using the Utrecht applicator, which enabled the study center to be recognized as an IC/IS center based on the definition used in the retroEMBRACE study. [1] The BT was scheduled as 6.5-7.5 Gy in 3-4 fractions based on the EBRT dose. For those 24 patients included in this study, a total of 266 interstitial needles were inserted under CT guidance during 76 of 93 BT fractions. In 17 of these 93 fractions, needle insertion was not indicated.
fractions, in which dose optimized was in two different ways (in total 76 fractions * 2 plans = 156 plans):

1. ICBT\textsubscript{Target-focused} plan: The priority was that CTV\textsubscript{HR} D90 > 85 Gy was achieved. Then, the OARs were spared as much as possible. The aim was to determine the IS-BT advantage in terms of OAR sparing when optimal target coverage was maintained.

2. ICBT\textsubscript{OARs-focused} plan: The priority was given to the OAR sparing. The highest CTV\textsubscript{HR} D90 was tried to achieve within the allowed OAR dose limits. The aim was to determine the IS-BT advantage in terms of target coverage when OARs sparing was maintained.

The EQD\textsubscript{210} dose for CTV\textsubscript{HR} D90 and EQD\textsubscript{23} doses for the 2 cc of the OARs (D2cc) including bladder, rectum, and sigmoid were calculated, and summed with the EBRT EQD2 dose. Eventually, for each of the 24 patients, three different plans were obtained: (1) Plan with needle (IS-ICBT), (2) ICBT\textsubscript{Target-focused}, and (3) ICBT\textsubscript{OARs-focused}.

To determine the benefit of needle use, IS-ICBT plans were compared with these two plans (for target coverage: IS-ICBT vs. ICBT\textsubscript{OARs-focused} for OAR sparing: IS-ICBT vs. ICBT\textsubscript{OARs-focused}). The dose difference (Δdose) between the plans was compared further between patients with small and large CTV\textsubscript{HR} to establish if the advantage of IS-BT was similar between these two patient groups.

**Statistical Analysis**

The normality of the continuous variables was determined using the Kolmogorov-Smirnov test, and Q-Q plots were checked. Between-group comparisons of continuous variables were performed using the independent t-test and Mann-Whitney U-test for normally and non-normally distributed variables, respectively. For comparisons between more than two groups, one-way ANOVA and the Kruskal-Wallis test were used for normally and non-normally distributed continuous variables, respectively. Categorical variables were compared using the chi-square test. A two-sided p ≤ 0.05 was considered statistically significant. All analyses were performed using the Statistical Package for the Social Sciences (SPSS) for Windows, version 21.0 (SPSS Inc., Chicago, IL, USA).

**Results**

For the 24 patients included in this study, a total of 266 interstitial needles were inserted under CT guidance during 76 of 93 BT fractions (three fractions in three patients and four in 21 patients).

The FIGO staging of the patients was as follows: Stages IIA (n = 1), IIB (n = 4), IIIB (n = 4), IIIC1 (n = 9; [T2bN1 (n = 7), T3bN1 (n = 2)]), IIIC2 (n = 2; [T1b2N1 (n = 1), T2bN1 (n = 1)]), IVA (n = 3; [T4N0 (n = 1), T4N1 (n = 2)]), and IVB (n = 1; T3aN1M1). Eleven patients had a small CTV\textsubscript{HR} and 13 had a large CTV\textsubscript{HR}.

**Needle Dwell Intensities**

The average dwell intensity of an individual needle was 11±8% (range, 0-42), and the dwell intensity was >15% in 63 out of 266 needles. The total contribution of the needles inserted in a fraction was 37.2±19.2% (range, 1.3-84.1). The average contribution of the needles to the complete BT treatment was 30.3±18%.

**The Comparison of the IS-ICBT and IC-BT Plans**

The IS-ICBT plan resulted in a significant increase in EQD\textsubscript{210} CTV\textsubscript{HR} D90 compared with ICBT\textsubscript{Target-focused} plans, with an average Δdose 10.5±6.2 Gy, which translated into a relative dose increase by 11.9±7.9% (Table 1, Figs. 1, 2).

The OAR 2cc EQD\textsubscript{2} 3 doses were significantly decreased with the IS-ICBT plans compared to ICBT\textsubscript{Target-focused} plans, with an average Δdose of 24.5±25.9 Gy, 7.6±9.7 Gy and 18.3±15.3 Gy for bladder, rectum, and sigmoid, respectively (Table 1, Figs. 1, 3).

**The Comparison of Patients with Small and Large CTV\textsubscript{HR}**

There was no significant difference between patients with small and large CTV\textsubscript{HR} in terms of CTV\textsubscript{HR} D90.
Discussion

A large CTV$_{HR}$ is regarded as a standard indication for IS-ICBT. However, 46% of our patients who received IS-ICBT consisted of patients with a small CTV$_{HR}$, which was in line with the previous series that reported approximately a frequency of 40%.[4,11,12] In these patients, asymmetrical extensions of the CTV$_{HR}$ in relation to uterus, an OAR located close to the high dose region, larger unilateral target extensions (>3.5 and >2.5 cm at vaginal applicator and point A level, respectively) make IS-ICBT necessary.[4,13]

True benefit of the IS-ICBT can be best demonstrated by the comparison of the plans with and without IS-CT in the same patient, rather than comparison of different patient groups or historical controls. Using such an in-patient pairwise comparison, the addition of IS-BT has been shown to increase CTV$_{HR}$ D90 EQD2 doses by 4-8 Gy on average without a significant increase in the OARs doses.[2,3,11,14] Similar therapeutic advantage was also demonstrated in studies comparing different patient groups.[4,13,15,16] The average Δdose between IS-ICBT and ICBT OAR-focused plan was 10.5±6.2 Gy, which was consistent between patients with small and large CTV$_{HR}$.

A notable characteristic of the current study was that IS-BT effect on both target dose and OARs sparing was evaluated separately, to the best of our knowledge, which was not investigated in the previous studies. The

![Fig. 1. The dose difference between IS-ICBT and ICBT plans in patients with small and large CTV$_{HR}$. IS-ICBT plans were used as reference plan and compared with ICBT$_{OARs-focused}$ for CTV$_{HR}$ D90 and ICBT$_{OARs-focused}$ for D2cc of bladder, rectum and sigmoid. IS-ICBT: Interstitial intracavitary brachytherapy; CTV$_{HR}$: High-risk clinical target volume; Gy: Gray.](image1)

![Fig. 2. CTV$_{HR}$ D90 EQD2$_{Gy}$ doses of the 24 patients included in the study, based on their CTV$_{HR}$ volume. Red area indicates the doses which is lower than the minimum criteria for CTVHR D90, that is, 85 Gy. IS-ICBT: Interstitial intracavitary brachytherapy; CTV$_{HR}$: High-risk clinical target volume; OAR: Organ at risk; Gy: Gray.](image2)
The benefit of intracavitary-interstitial brachytherapy (IS-BT) in cervical cancer can be explained by more effective use of IS-BT using CT-guidance on the day of treatment after applicator insertion, rather than MRI post-external beam radiation therapy (EBRT). A 5-10 Gy difference in dose could have a clinical effect on tumor control and side effects, based on the evidence from image-guided studies. Not only was there remarkable reduction in the prescribed dose between patients with and without IS-BT plans, with an average Δdose of 24 and 18 Gy, respectively. Although non-significant, the sigmoid sparing was even higher in patients with small clinical target volume high-risk (CTV_{HR}) compared with those with large CTV_{HR} (21.8±19.6 Gy vs. 15.2±10.1 Gy). These extreme dose differences between patients with and without IS-BT plans can be explained by more effective use of IS-BT using CT-guidance on the day of treatment after applicator insertion, rather than MRI post-EBRT. Considering that 5-10 Gy difference had a clinical effect on tumor control and side effects, based on the evidence from image-guided studies, not only pa-

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**Fig. 3.** OAR D2cc EQD23 doses of the 24 patients included in the study, based on their CTV_{HR} volume. Red area indicates the doses which is higher than the maximal allowable dose criteria for that OAR, that is, 90 Gy for bladder D2cc and 75 Gy for rectum and sigmoid D2cc.

IS-ICBT: Interstitial intracavitary brachytherapy; CTV_{HR}: High-risk clinical target volume; OAR: Organ at risk; Gy: Gray.
tients with a large CTV_{HR} but also with small CTV_{HR} can take a great advantage of IS-BT, when indicated.[17-21]

Limitations of the Study
Our study has some limitations. First, it is a single center study, which may not precisely reflect the practice in other institution. Second, IS-ICBT plans were the clinical plans used for patient treatment, whereas IC-BT plans were retrospectively created without time pressure. Therefore, even better dose distribution could have been achieved with IS-ICBT plans if they were also created only for research purposes without haste. Third, CT planning was used in this study, whereas results with MRI planning can deviate from our results.

Conclusion
A considerable proportion of the patient treated with IS-ICBT consists of patients small CTV_{HR} (40-45%). The therapeutic advantage provided by IS-BT was similar between patients with small and large CTV_{HR} with regard to better target coverage (10 Gy increase on average) and OARs sparing. Dramatic dose reduction was obtained in sigmoid and bladder using IS-ICBT in both patient groups.

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Conflict of Interest: All authors declared no conflict of interest.

Ethics Committee Approval: All the treatment procedures reported in this study were a part of the routine clinical practice in the institution and were conducted after obtaining consent as relevant. The ethics committee deemed that additional informed consent for this study was not required, based on the Liv Hospital-ULus Department of Radiation Oncology Medical Research Involving Human Subjects Act. However, all patients were informed that their data could be used for research purposes and that they could refuse consent for such use.

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References
4. Kirisits C, Lang S, Dimopoulos J, Berger D, Georg D, Potter R. The Vienna applicator for combined intracav-

Table 2  Comparison of the average IS-ICBT doses, Δdose (the difference between IS-ICBT and ICBT) in CTV_{HR} D90 and OAR D2cc values between patients with small and large CTV_{HR}

<table>
<thead>
<tr>
<th>CTV_{HR}</th>
<th>Small CTV_{HR} (n=11)</th>
<th>Large CTV_{HR} (n=13)</th>
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<tr>
<td>EBRT dose</td>
<td>22.0±5.9 cm³</td>
<td>54.4±22.8 cm³</td>
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<tr>
<td>BT dose per fr</td>
<td>49.5±4.2 Gy</td>
<td>50±4.2 Gy</td>
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<td>CTV_{HR} D90</td>
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<td>6.8±0.6 Gy</td>
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<tr>
<td>Bladder D2cc</td>
<td>88±4.8 Gy</td>
<td>88±4.3 Gy</td>
<td>0.858</td>
</tr>
<tr>
<td>Rectum D2cc</td>
<td>82.2±7.1 Gy</td>
<td>85.1±7.4 Gy</td>
<td>0.370</td>
</tr>
<tr>
<td>Sigmoid D2cc</td>
<td>66.7±8.3 Gy</td>
<td>68.4±6.0 Gy</td>
<td>0.565</td>
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<tr>
<td>Δdose CTV_{HR} D90</td>
<td>75.1±9.1 Gy</td>
<td>77.3±6.9 Gy</td>
<td>0.507</td>
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<tr>
<td>Δdose Bladder D2cc</td>
<td>-9.6 (5.8-14.8) Gy</td>
<td>-12.1 (4.8-16.3) Gy</td>
<td>0.765</td>
</tr>
<tr>
<td>Δdose Rectum D2cc</td>
<td>12.5 (2.9-36.0) Gy</td>
<td>21.8 (7.5-50.6) Gy</td>
<td>0.543</td>
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<tr>
<td>Δdose Sigmoid D2cc</td>
<td>2.7 (1.3-12.6) Gy</td>
<td>3.1 (0.0-15.2) Gy</td>
<td>0.931</td>
</tr>
</tbody>
</table>
| Δdose Mean±SD values are given. Median (Q25-Q75) are given for the Δdose. *EQD2 10 values: ΔDose=ICBT OARs-focused - IC-ISBT. ‘EQD2 3 values: ΔDose=ICBT Target-focused - IC-ISBT. IS-ICBT: Interstitial intracavitary brachytherapy; CTV_{HR}: High-risk clinical target volume; EBRT: external-beam RT; OAR: Organ at risk; Gy: Gray.
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