The association of rectal equivalent dose in 2 Gy fractions (EQD2) to late rectal toxicity in locally advanced cervical cancer patients who were evaluated by rectosigmoidoscopy in Faculty of Medicine, Chiang Mai University

Ekkasit Tharavichtikul, MD1, Pooriwat Meungwong, MD2, Taned Chitapanarux, MD3, Somvilai Chakrabhandhu, MD1, Pitchayaponne Klunklin, MD1, Wimrak Onchan, MD1, Somsak Wanwilairat, PhD1, Patinee Traisathit, PhD4, Razvan Galalae, MD, PhD5, Imjai Chitapanarux, MD1

1Division of Therapeutic Radiology and Oncology, Department of Radiology, Faculty of Medicine, Chiang Mai University, Chiang Mai, Thailand; 2Lampang Cancer Hospital, Lampang, Thailand; 3Department of Medicine, Faculty of Medicine, Chiang Mai University, Chiang Mai, Thailand; 4Biostatistics and Applied Statistics Laboratory, Department of Statistics, Faculty of Science, Chiang Mai University, Chiang Mai, Thailand; 5Faculty of Medicine, Christian-Albrechts University at Kiel, Kiel, Germany

Purpose: To evaluate association between equivalent dose in 2 Gy (EQD2) to rectal point dose and gastrointestinal toxicity from whole pelvic radiotherapy (WPRT) and intracavitary brachytherapy (ICBT) in cervical cancer patients who were evaluated by rectosigmoidoscopy in Faculty of Medicine, Chiang Mai University.

Materials and Methods: Retrospective study was designed for the patients with locally advanced cervical cancer, treated by radical radiotherapy from 2004 to 2009 and were evaluated by rectosigmoidoscopy. The cumulative doses of WPRT and ICBT to the maximally rectal point were calculated to the EQD2 and evaluated the association of toxicities.

Results: Thirty-nine patients were evaluated for late rectal toxicity. The mean cumulative dose in term of EQD2 to rectum was 64.2 Gy. Grade 1 toxicities were the most common findings. According to endoscopic exam, the most common toxicities were congested mucosa (36 patients) and telangiectasia (32 patients). In evaluation between rectal dose in EQD2 and toxicities, no association of cumulative rectal dose to rectal toxicity, except the association of cumulative rectal dose in EQD2 >65 Gy to late effects of normal tissue (LENT-SOMA) scale ≥ grade 2 (p = 0.022; odds ratio, 5.312; 95% confidence interval, 1.269–22.244).

Conclusion: The cumulative rectal dose in EQD2 >65 Gy have association with ≥ grade 2 LENT-SOMA scale.

Keywords: Cervical cancer, Radiotherapy, Cumulative dose, Rectum, Toxicity, Association

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Correspondence: Ekkasit Tharavichitkul, MD, Division of Therapeutic Radiology and Oncology, Department of Radiology, Faculty of Medicine, Chiang Mai University, Chiang Mai, 239 Huay Kaew Road, Muang District, Chiang Mai 50200, Thailand. Tel: +66-53945456, Fax: +66-53945491, E-mail: paan_31@hotmail.com

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Introduction

Cervical cancer is the second most common cancer in incidence and mortality in Thai women. In Thailand, the age-standardized rates were 24.5 and 12.8 in incidence and mortality, respectively [1]. Concurrent chemotherapy and radiotherapy is the treatment of choice for locally advanced cancers because combination of these two modalities improves tumor control, overall survival, and progression-free survival [2-6]. In radiotherapy part, combination of whole pelvic radiotherapy (WPRT) and intracavitary brachytherapy (ICBT) has become a standard treatment modality for cervical cancer. Initial WPRT is performed to obtain tumor shrinkage to permit optical ICBT placement [7]. According to the American Brachytherapy Society recommendations for high-dose-rate (HDR) ICBT for carcinoma of the cervix, the whole pelvic external beam radiotherapy (EBRT) dose varies from institution to institution. The HDR fraction size and number depends on the EBRT dose [8]. HDR brachytherapy has been used for more than 30 years in Japan and Europe, and its applications have been increasing in the United States [9,10]. ICBT has been used to increase the dose to cervix to ≥80 Gy. However organs in the proximity of the cervix, such as urinary bladder, small bowel and rectum, can receive rather high doses and adverse side effects can occur. The risk of late effects (i.e., obstruction, fibrosis/necrosis, or fistula) is related to the volume, total dose, dose per fraction, and specific intrinsic radiosensitivity of the normal tissue irradiated. Biologically effective dose (BED) had been used to evaluate the dose to target and organs at risk [11,12]. Because numerical values of BED (in Gy$_{eq}$ or Gy$_{B}$) are not familiar to many medical professionals, it is a common practice to convert these BEDs to the biologic equivalent of a total dose in 2 Gy fractions (BED of 2 Gy). These simplified biologically equivalent doses are called either EQD2, denoting equivalent total doses in 2-Gy fractions [13]. Rectal bleeding is a common late toxicity of pelvic radiation. Late toxicity to the rectum includes radiation proctitis, ulceration, stricture, and fistula. Some symptoms may resolve in a few months, whereas others will have a longer course that may involve 1 to 2 years of rectal bleeding. For severe cases, surgical intervention may be needed [14,15].

To evaluate the rectal toxicity, many scores in objective and subjective approaches were used to evaluate. Radiation Therapy Oncology Group/European Organization for Research and Treatment of Cancer (RTOG/EORTC) toxicity scale were used in objective aspect only while Vienna Rectoscopy Score (VRS) was used in subjective aspect [16,17]. Moreover, there were the combinations of subjective and objective evaluations as late effects of normal tissues (LENT-SOMA) scales [18]. For this study, we retrospectively evaluated the endoscopic findings in cervical cancer patients who were treated by radical radiotherapy and also accumulative dose using EQD2 to rectum is associated with gastrointestinal toxicity ≥grade 2.

Materials and Methods

This study designed as retrospective study in 336 patients who were treated by radical radio-chemotherapy for locally advanced cervical cancer in the Division of Therapeutic Radiology and Oncology, Faculty of Medicine, Chiang Mai University during 2004–2009. We enrolled 39 patients of them who were evaluated the rectal toxicity by rectosigmoidoscopy during the follow-up period according to symptomatic indications or patient preferences. Treatment schedules for these patients were shown as the followings.

1. External beam radiotherapy
WHRT was performed using conventional radiotherapy with conventional fractionation (2 Gy/fraction) by 6- or 10-MV X-ray with two- or four-field techniques according to anteroposterior diameter. After 40 to 44 Gy of WPRT, central shielding was inserted using 3-or 4-cm width rectangular block at midline. The field was coned down after the initial 50 Gy in an effort to boost parametrium and exclude small bowel to 56 Gy. For concurrent chemotherapy, mitomycin-C 10 mg/m$^2$, 1st and 6th week or weekly cisplatin 40 mg/m$^2$ was given intravenously during WPRT [18].

2. High-dose-rate brachytherapy
First HDR-ICBT was started after 4th week of the treatment. Patients were treated with HDR-ICBT using Fletcher-type (Fletcher-Williamson Asian-Pacific) applicators. Anterior and posterior vaginal packing with gauze was used to maximize the distance from the source to the bladder wall and the rectal wall. The dose was prescribed to Point A as the dose of 6 to 7.2 Gy in 3 to 4 fractions. To evaluate rectal point dose, a rectal probe was used to identify the anterior rectal wall. The International Commission on Radiation Units and Measurements (ICRU) rectal points were identified using a laterally orthogonal radiographs taken during each individual application and defined as the points that an line drawn from the tip of ovoid applicator to the anterior border of the rectal
The affect of rectal dose to late toxicity

probe, the 4 another rectal points were digitized in 1- and 2-cm superior and inferior directions according to the first rectal point (Fig. 1). Calculation of the dose profiles was also based on these radiographs, and the maximum point dose was recorded.

3. Calculation of rectal dose
To determine the rectal dose from the WPRT plus ICBT, the total dose to rectal point was calculated as EQD2 using the linear quadratic model [13,19].

4. Follow-up schedule
Late complications after RT are defined as side effects developing 3 months after RT completion. For each endpoint, the grade of late morbidity will be the maximal grade ever recorded, even if the side effect later subsided. The data will be retrospectively collected from treatment data and patient record of each patient. Patients that were sent to perform endoscopy were retrospectively collected to evaluate. The endoscopy report consists of a written description and a video photographic documentation. Rectal toxicity scoring is according to systematic sigmoidoscopy score for radiation-induced proctitis proposed by Medical University of Vienna using standard endoscopic terminology that had been previously used to evaluate the radiation-induced changes in the rectal mucosa of patients with prostate cancer treated with radical radiotherapy [17]. The grading system determines from the endoscopic terminology of the World Organization for Digestive Endoscopy [20].

5. Statistical analysis
Observations will be censored at date of last visit, at 3 months of follow-up, or at date of failure. Logistic regression analysis was used to evaluate the association of cumulative rectal dose in EQD2 to toxicity. Descriptive statistics were used to evaluated the characteristic data of patients (age, treatment time, and etc.) All statistical analyses will be performed using IBM SPSS statistical software ver. 19.0 (IBM, Armonk, NY, USA).

Results
The mean cumulative dose in term of EQD2 to Point As and rectum were 76.4 and 64.2 Gy, respectively. All Characteristic data of these patients were shown in Table 1.

1. Toxicity profiles
Grade 0–1 toxicity was observed in 34 patients for RTOG/EORTC toxicity criteria and 25 patients for LENT-SOMA scale. In endoscopic findings, the first and second most common toxicities were congested mucosa (36 patients) and telangiectasia (32 patients). No necrosis was observed in this group. According to VRS proposed by Medical University of Vienna, grade 1 toxicity (51%) was the most common finding. When we evaluated in VRS, there were 6, 20, 7, 3, and 3 patients for score 0, 1, 2, 3, and more than 3, respectively. The data of toxicities were showed in Table 2.

When we considered the cumulative dose to rectum
and toxicity scale, we found the associations between the cumulative dose to rectum more than 65 Gy and LENT-SOMA scale at least grade 2 (p = 0.022; odds ratio, 5.312; 95% confidence interval, 1.269–22.244). When we evaluated the association of the same dose level (>65 Gy) and other toxicity scale, no association was found in RTOG/EORTC (p = 0.137) and VRS (p = 0.497) (Table 3).

Discussion and Conclusion

During the past decade, BED was used to evaluate the cumulative dose to tumor and organs-at-risk. For cervical cancer, the cumulative dose to rectum and bladder in terms of BED correlated to rectal toxicity [21,22].

Rectal toxicity is the most common toxicity after radical radiotherapy for cervical cancer. The combined dose of EBRT and brachytherapy to rectum predicted the toxicity in terms of BED. Toita et al. [23] suggested that cumulative BED at the rectal point should be kept below 100–120 Gy3 to prevent late rectal complication. However, Ogino et al. [26] also showed that the time-dose factor and the BED at rectal point values were significant correlated with the incidence of late rectal complication.

In terms of EQD2, Sakata et al. [27] found that rectal radiation tolerance dose using EQD2, 5% and 50% complication probability, was about 64 and 79 Gy, respectively, and the probability of rectal complications increased drastically after the maximal rectal dose was >60 Gy.

From our study, only association between cumulative rectal dose in EQD2 >65 Gy and LENT-SOMA ≥2 were observed (p = 0.022). No statistical significance was found between rectal dose and toxicity in terms of RTOG/EORTC scale (p = 0.137) and VRS (p = 0.497). The possible explanation is that LENT-SOMA scale covered both objective and subjective evaluations while RTOG/EORTC and VRS were purely objective and subjective approaches, respectively. So LENT-SOMA scale should be more appropriate to evaluate the rectal toxicity. Moreover, in this study, the association of cumulative rectal toxicity in EQD2 to toxicity ≥1 was not found statistical significance in all toxicity score. This correlates to the clinical aspects that the decision to treat toxicity starts practically when patients developed ≥grade 2 toxicity [28].

However, there are some limitations in this study. The accuracy of the rectal dose calculation is affected by several factors. First, to calculate the rectal dose from EBRT, it assumed that the rectum was completely shielded after central shielding. However, the rectum might be irradiated with some doses according to anatomic variations. Secondly, using orthogonal films for calculation might cause problem to evaluate the really maximal dose to rectum due to anatomy. Volume-based
planning brachytherapy could be more accurate on delineating organs-at-risk including rectum and assessing doses through dose volume histograms [29]. The study from Kato et al. [30] reported the dose evaluation of 48 patients who were treated by combination of WPRT plus HDR-ICBT. Computed tomography-based three dimensional (3D) dose-volume parameters of the rectum may be effective for predicting late rectal complications while no correlation between rectal dose in ICRU point and toxicity was found [30]. The using of 3D-based approach will be reported in the future. Thirdly, the number of patients who performed endoscopic exam in our study was small because rectosigmoidoscopy was not routine evaluation after treatment in our country.

Our findings showed correlation between cumulative EQD2 to rectal point >65 Gy and rectal toxicity in term of LENT-SOMA scale at least grade 2. The volume-based planning in both WPRT and ICBT improve the accuracy in dosimetry and toxicity evaluation.

Conflict of Interest

No potential conflict of interest relevant to this article was reported.

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