Evaluating Variations of Bladder Volume Using an Ultrasound Scanner in Rectal Cancer Patients during Chemoradiation: Is Protocol-Based Full Bladder Maintenance Using a Bladder Scanner Useful to Maintain the Bladder Volume?

Hong In Yoon1, Yoonsun Chung2, Jee Suk Chang1, Joo Yong Lee3, Soo Jung Park4, Woong Sub Koom1,5*

1 Department of Radiation Oncology, Yonsei University College of Medicine, Seoul, Korea, 2 Department of Radiation Oncology, Samsung medical center, Seoul, Korea, 3 Department of Urology, Yonsei University College of Medicine, Seoul, Korea, 4 Department of Internal Medicine and Institute of Gastroenterology, Yonsei University College of Medicine, Seoul, Korea, 5 Yonsei Song-Dang Institute for Cancer Research, Yonsei University College of Medicine, Seoul, Korea

* mdgold@yuhs.ac (WSK); SJPARK@yuhs.ac (SJP)

Abstract

Purpose

The maintenance of full bladder is important to reduce radiation-induced toxicities and maintain the therapeutic consistency in locally advanced rectal cancer patients who underwent radiotherapy (RT). So, the aim of this study was to evaluate the effectiveness of protocol-based full bladder maintenance by assessing bladder volume variation using an ultrasound bladder scanner to maintain bladder volume.

Materials and Methods

From March 2011 to May 2011, twenty consecutive rectal cancer patients receiving external beam RT participated in this prospective study. Protocol-based full bladder maintenance consisted of education, training and continuous biofeedback by measuring bladder volume. Bladder volume was measured by bladder scan immediately before simulation CT scan and before each treatment three times weekly during the RT period. The relative bladder volume change was calculated. Intra-patient bladder volume variations were quantified using inter-quartile range (IQR) of relative bladder volume change in each patient. We compared intra-patient bladder volume variations obtained (n=20) with data from our previous study patients (n=20) performing self-controlled maintenance without protocol.
Results
Bladder volumes measured by bladder scan highly correlated with those on simulation CT scan (R=0.87, p<0.001). Patients from this study showed lower median IQR of relative bladder volume change compared to patients of self-controlled maintenance from our previous study, although it was not statistically significant (median 32.56% vs. 42.19%, p=0.058). Upon logistic regression, the IQR of relative bladder volume change was significantly related to protocol-based maintenance [relative risk 1.045, 95% confidence intervals (CI) 1.004-1.087, p=0.033]. Protocol-based maintenance included significantly more patients with an IQR of relative bladder volume change less than 37% than self-controlled maintenance (p=0.025).

Conclusion
Our findings show that bladder volume could be maintained more consistently during RT by protocol-based management using a bladder scan.

Introduction
Preoperative chemoradiation (CRT) is a standard treatment for locally advanced rectal cancer, which increases local control and sphincter preservation rates compared with adjuvant treatments [1–4]. Additionally, for rectal cancer patients with adverse pathologic factors such as T3-4 or positive lymph nodes, postoperative CRT is required to improve treatment outcome [5]. Generally, pelvic irradiation for rectal cancer patients is safe and tolerable.

Small bowel damage is a common treatment-related complication for rectal cancer patients receiving CRT. It is known that the risk of gastrointestinal (GI) toxicities is related with irradiated small bowel volume and the dose within the radiation field [6]. In some cases, patients develop acute toxicities such as nausea, vomiting, abdominal pain, and abnormal absorption, as well as chronic toxicities including bowel obstruction, perforation, and intestinal stricture. In preoperative CRT for rectal cancer, grade 3 or 4 acute toxicity and diarrhea is reported to develop in 12~36% of patients, whereas long-term GI toxicity occurs in 9% of patients [2,7,8]. Similarly, 18~35% of patients receiving postoperative CRT are reported to have severe acute toxicity and 5~15% of patients have late toxicity related to small bowel damage [2,7,9,10]. Therefore, many investigators have attempted to reduce the irradiated dose of small bowel using non-surgical methods such as the prone position, as well as small bowel displacement devices like the belly board and bladder distension [11,12]. Since 2009, our institution has treated all rectal cancer patients by distending the bladder fully with a belly board in the prone position preoperatively or postoperatively.

However, after CT simulation with a full bladder volume, patients find it very difficult to maintain the bladder volume similar to that of the simulation CT during the entire duration (5 to 6 weeks) of radiotherapy (RT). Some studies using ultrasound bladder scanner for cervical or prostate cancer reported large variations of bladder volume but noted the bladder scanner was useful and accurate to assess the inter-fractional variation of bladder volume [13–16]. Our institution has assessed bladder volume every other day using a bladder scanner since February 2011 and in our previous pilot study, we reported that there were bladder volume variations and reductions in bladder volume in rectal cancer patients receiving CRT [17]. It was previously reported that biofeedback could improve the consistency of bladder volume despite a lack of statistical significance [16]. Thus, the purpose of this prospective study was to evaluate the
usefulness of protocol-based full bladder maintenance by assessing variations of bladder volume using an ultrasound bladder scanner to maintain the bladder volume during radiation for locally advanced rectal cancer.

Materials and Methods

Patients
This prospective observational study received approval from the Severance internal review boards of the Severance hospital (IRB No. 4-2010-0832). From March 2011 to May 2011, twenty consecutive rectal cancer patients receiving external beam RT participated in this study and provided written informed consent. The inclusion criteria of this study were as follows: age ≥20 years with planned preoperative or postoperative RT for locally advanced rectal cancer. We excluded patients who had urinary frequency of more than once per hour, nocturia more than 4 times per day, urinary incontinence, median laparotomy scar or lymphocele interfering with bladder scanning, aberrant bladder form, or poor compliance. Patient characteristics are listed in Table 1.

Chemoradiotherapy
CT simulation and treatment were performed with the patient with a full bladder in a prone position with a belly board [18]. All patients received CRT consisting of 3-dimensional conformal RT (3D-CRT). Treatment planning was accomplished with Pinnacle3 (Philips Medical Systems, Andover, MA). Target volumes and critical adjacent organs, including the bladder and small bowel, were delineated by the attending radiation oncologists. The clinical target volume included the primary tumor mass or postoperative tumor bed, mesorectum, presacral space, as well as the pelvic lymph nodes and/or external iliac lymph nodes if indicated. For the whole pelvic field, the superior border was defined as the L5-S1 interspace, the inferior border as 3–4 cm below the primary tumor or inferior border of obturator foramen, and the lateral border as 1.5 cm behind the true bony pelvis. For the lateral fields, we defined the anterior border as the posterior margin of the symphysis pubis and the posterior border as the anterior bony sacral margin. In case of a tumor extending directly to a pelvic organ with external drainage, we modified the anterior border to the anterior border of the symphysis pubis to include the external iliac lymphatics. The 3D-CRT consisted of 41.4–45 Gy of whole pelvic RT and 4.5–9.0 Gy of local boost RT to the postoperative tumor bed or primary tumor. The median total dose was 50.4 Gy (range, 45–54). Concurrent intravenous chemotherapy was administered with a 5-fluorouracil (425 mg/m2) and leucovorin (20 mg/m2) bolus on weeks 1 and 5 of 3D-CRT.

Protocol-based full bladder maintenance by education, training, and continuous biofeedback
Protocol-based full bladder maintenance consisted of education, training and continuous biofeedback by measuring bladder volume. The specific bladder-filling instructions regarding comfortably full bladder were given as follows. First, immediately after patients void the bladder, they drink 500 ml water to have a comfortably full bladder. Second, patients wait until they have a definite but easily tolerable micturition urge (grade 3) or intolerable micturition urge (grade 4). Third, patients record each duration of grade 3 or grade 4 micturition urge in a micturition diary. Fourth, patients visit the hospital at an intermediate status (between grade 3 and 4 of micturition urge) to have a comfortably but maximally full bladder. Until the date of CT simulation, patients were asked to train at least twice according to these specific instructions. Next, all patients were additionally trained three times a week (Monday, Wednesday,
and Friday) during the 5- or 6-week RT course according to the biofeedback protocol. The aim of the biofeedback protocol was to improve the constancy of bladder volume measured on bladder scan. We attempted to improve the ability of patients to detect bladder-filling sensations, similar to bladder fullness on simulation CT scan. The feedback consisted of advising patients to drink more water or suppress urination longer according to their daily bladder volume. When the bladder volume of patients ranged from 80% to 120% of that of the simulation CT scan, patients were instructed to keep the same pattern of bladder filling the following day. When the bladder volume was less than 80% of that of the simulation CT scan, patients

Table 1. Patient characteristics.

| Characteristics                        | n   | (%)  |
|----------------------------------------|-----|------|
| Age (years)                            | Median 56 | (22–73) |
| Gender                                 | Female 6 | (30.0) |
|                                        | Male 14 | (70.0) |
| Performance status                     | ECOG PS 0 8 | (40.0) |
|                                        | ECOG PS 1 12 | (60.0) |
| Distance from AV (cm)                  | Median 7 | (3–12) |
|                                        | Range 6 | (30.0) |
| Tumor location                         | Lower rectum 8 | (40.0) |
|                                        | Mid rectum 6 | (30.0) |
| Pathology                              | Adenocarcinoma 20 | (100.0) |
| Tumor grade                            | WD 4 | (20.0) |
|                                        | MD 15 | (75.0) |
|                                        | PD 0 | (0.0) |
|                                        | Unknown 1 | (5.0) |
| Aim of radiotherapy                    | Preoperative 13 | (65.0) |
|                                        | Postoperative 7 | (35.0) |
| Clinical T stage for preoperative RT   | T2 2 | (15.4) |
|                                        | T3 10 | (76.9) |
|                                        | T4 1 | (7.7) |
| Clinical N stage for preoperative RT   | N0 3 | (23.1) |
|                                        | N1 1 | (7.7) |
|                                        | N2 9 | (69.2) |
| Pathologic T stage for postoperative RT| T2 2 | (28.6) |
|                                        | T3 5 | (71.4) |
|                                        | T4 0 | (0.0) |
| Pathologic N stage for postoperative RT| N0 2 | (28.6) |
|                                        | N1 3 | (42.9) |
|                                        | N2 2 | (28.6) |
| RT dose (Gy)                           | Median 50.4 | (45–54) |
| Fractional dose (Gy)                   | Median 1.8 | (1.8–2.0) |

Patient characteristics are listed.

Abbreviations: ECOG PS = Eastern Cooperative Oncology Group Performance Status Scale; WD = well differentiated; MD = moderately differentiated; PD = poorly differentiated; RT = radiotherapy

doi:10.1371/journal.pone.0128791.t001
were instructed to drink 100 or 200 ml more fluid or to suppress urination longer. When the bladder volume was less than 50% of the simulation CT scan volume, patient was advised to suppress urination for 30 minutes to 1 hour. When the measurement was greater than 120% of the simulation CT scan, patients were advised to drink less.

**Bladder volume measurements**

Using a portable automated ultrasonic bladder scanner (Biocon-700, Mcube Technology, Korea), bladder volume was measured by two experienced physicians immediately before the simulation CT scan as well as prior to treatment during 5 or 6 weeks of RT. Patients were comfortably positioned in a supine position. Next, the operator placed the scanner probe on two fingers over the symphysis pubis at the midline of abdomen and angled the probe towards the bladder. Operators checked the three-dimensional real time image before scanning to locate the bladder position for more accurate measurement. Each bladder scan time and bladder volume (ml) measured by the bladder scanner was documented. Before simulation CT, the bladder was scanned consecutively 5 times, and the median value was recorded as the baseline bladder volume. The bladder volume was also calculated using the contour of bladder inner wall based on the simulation CT images. During the entire treatment period, each patient’s bladder was scanned 3 days per week (Monday, Wednesday, and Friday) just prior to RT, and 5 times daily to report the median value. Each ultrasound scan required approximately 1–2 min and the addition of an ultrasound scan did not affect overall treatment time.

**Toxicity evaluation and follow-up**

Patients were prospectively followed up at 1 and 3 months following the completion of RT, then every 3 months during the first 2 years after RT, and every 6 months from the third year. Treatment-related adverse events were evaluated according to the National Cancer Institute Common Toxicity Criteria (version 3.0).

**Statistical analyses**

When bladder volume measurement data was missing for any reason, we replaced the missing values for bladder volume measured using the last observation carried forward (LOCF) method. Eighteen patients (90%) completed planned measurement of bladder volume according to schedule. In two patients, bladder volume measurement data was missing, (one data in one patient and five data in the other). In order to assess the accuracy of bladder ultrasound scan, the correlation between the bladder volume measured by bladder scanner and the bladder volume calculated using simulation CT images was analyzed using Pearson’s correlation tests. The bladder volumes during the RT and the baseline bladder volumes measured by bladder ultrasound scan before CT simulation were compared using Wilcoxon signed-rank tests. To evaluate the protocol-based full bladder maintenance, we compared 20 patients from this study with 20 patients performing self-controlled maintenance without protocol-based education, training and continuous biofeedback from our previous study [17]. The relative bladder volume change of all patients was calculated using the following equation:

\[
\text{Relative bladder volume change (\%)} = \frac{\text{Bladder volume during RT (ml)} - \text{Baseline bladder volume (ml)}}{\text{Baseline bladder volume (ml)}} \times 100
\]

We calculated the interquartile range (IQR) of relative bladder volume change for each patient in this study and each patient in the previous study. Intra-patient bladder volume
variations were quantified by IQR of relative bladder volume change in each patient. Next, we compared relative bladder volume change and IQR of relative bladder volume change of each patient between two studies using Mann-Whitney U tests. For further evaluation of the effect of protocol-based full bladder maintenance on improvement of the intra-patient bladder volume variations, we analyzed the correlation between IQR of relative bladder volume change and protocol-based full bladder maintenance using binary logistic regression for a total of 40 patients from this and previous studies. We determined the cut-off value of IQR of relative bladder volume change for protocol-based full bladder maintenance using receiver operating characteristic (ROC) curves. The 40 patients were divided into two groups according to the cut-off value. We performed Chi-square tests to investigate the impact of protocol-based full bladder maintenance on the two groups according to the cut-off value. Statistical significance was defined as \( p < 0.05 \), and SPSS 20.0.0 (IBM Corporation, Armonk, New York, USA) was used for all analyses.

**Results**

**Treatment-related acute and chronic toxicity**

Most acute and chronic toxicities in this prospective study were mild (grade 0–2). Grade 4 or 5 acute toxicity was not observed in any patient. Eleven of the patients (55%) experienced grade 3 diarrhea, five patients (25%) had acutely grade 3 tenesmus, and one patient (5%) had a grade 3 skin rash (Table 2). Grade 4 or 5 chronic toxicity was also not observed in any patient. No patients experienced grade 3 chronic toxicity, excepting 3 patients (15%) with grade 3 diarrhea (Table 3).

**Bladder volume measurement validation**

All patients received bladder ultrasound scans a median of 16 times (range: 15–18; 3 days a week according to the RT schedule). All patients received prearranged scans a median of 16 times (range: 11–18). The median bladder volume on simulation CT image was 450 ml (IQR 195, range: 264–860). The median volume measured by bladder ultrasound scanner before simulation CT scan on the same day was 398 ml (IQR 240; range: 264–821). We observed that

| Grade 0 | Grade 1 | Grade 2 | Grade 3 | Grade 4 |
|---------|---------|---------|---------|---------|
| Nausea  | 15      | 75%     | 4       | 20%     |
| Vomiting| 0       | 0%      | 0       | 0%      |
| Fatigue | 3       | 15%     | 13      | 65%     |
| Weight loss | 18  | 90%     | 2       | 10%     |
| Anorexia | 6     | 30%     | 12      | 60%     |
| Diarrhea | 3     | 15%     | 1       | 5%      |
| Tenesmus | 9     | 45%     | 3       | 15%     |
| Incontinence, anal | 17  | 85%     | 3       | 15%     |
| Skin rash | 15   | 75%     | 4       | 20%     |
| Cystitis | 17    | 85%     | 3       | 15%     |
| Incontinence, urinary | 0   | 0%      | 0       | 0%      |
| Abdominal pain | 16  | 80%     | 3       | 15%     |

The data of acute toxicity are listed.

doi:10.1371/journal.pone.0128791.t002
Table 3. Prospective evaluation of late toxicity (n = 20).

| Grade 0 | Grade 1 | Grade 2 | Grade 3 | Grade 4 |
|---------|---------|---------|---------|---------|
| Nausea  | 20      | 100%    | 0       | 0%      | 0       | 0%      | 0       | 0%      |
| Vomiting| 0       | 0%      | 0       | 0%      | 0       | 0%      | 0       | 0%      |
| Fatigue | 14      | 70%     | 6       | 30%     | 0       | 0%      | 0       | 0%      |
| Weight loss | 20   | 100%    | 0       | 0%      | 0       | 0%      | 0       | 0%      |
| Anorexia | 15     | 75%     | 5       | 25%     | 0       | 0%      | 0       | 0%      |
| Diarrhea | 10     | 50%     | 3       | 15%     | 4       | 20%     | 3       | 15%     |
| Tenesmus | 15     | 75%     | 2       | 10%     | 3       | 15%     | 0       | 0%      |
| Incontinence, anal | 16 | 80%     | 4       | 20%     | 0       | 0%      | 0       | 0%      |
| Skin rash | 20    | 100%    | 0       | 0%      | 0       | 0%      | 0       | 0%      |
| Cystitis | 20     | 100%    | 0       | 0%      | 0       | 0%      | 0       | 0%      |
| Incontinence, urinary | 0   | 0%      | 0       | 0%      | 0       | 0%      | 0       | 0%      |
| Abdominal pain | 20   | 100%    | 0       | 0%      | 0       | 0%      | 0       | 0%      |

The data of chronic toxicity are listed.

doi:10.1371/journal.pone.0128791.t003

Bladder volumes measured by bladder scan highly correlated with those on simulation CT scan (R = 0.87, p < 0.001, Fig 1).

Time trends of bladder volume measured using bladder ultrasound scan

The bladder volume measurements and the number of patients adjusted by LOCF according to time sequence are described in Table 4. Due to missing data, we used LOCF from V4-2 to V5-3.

![Fig 1. Correlation of bladder volume measured using a bladder scan and simulation CT scan. This figure depicts that bladder volumes measured by bladder scan highly correlated with those on simulation CT scan (R = 0.87, p < 0.001).](doi:10.1371/journal.pone.0128791.g001)
Compared to baseline bladder volume by bladder scan, all median values of bladder volumes measured during RT were significantly lower ($p < 0.05$), except the first scan after the start of treatment ($p = 0.087$, Table 4).

Assessment of intra-patient variation according to protocol-based maintenance

The median values and IQRs of relative bladder volume change of each patient from this and our previous study patients were calculated (Table 5). Comparing median intra-patient relative bladder volume change between current and previous study, protocol-based full bladder maintenance group showed less bladder volume reduction than self-controlled maintenance group although a difference is not statistically significant (median 27.4% reduction in current study vs. 32.9% reduction in previous study, $p = 0.55$). To evaluate intra-patient variation, we investigated the difference between median values of each patient’s IQR of relative bladder volume change between patients receiving protocol-based full bladder maintenance ($n = 20$) and patients from the previous study who performed self-controlled full bladder maintenance ($n = 20$). Patients from this study receiving protocol-based maintenance showed lower median IQR of relative bladder volume change although a difference is not statistically significant [median 32.56% (range 18.03–61.09) vs. 42.19% (range 17.79–96.6), $p = 0.058$]. In Fig 2, we observed that patients receiving no protocol-based maintenance had wider IQR ranges of relative bladder volume change. Therefore, to further evaluate the correlation between IQR of relative

Table 4. Time trends of bladder volume from CT scans and bladder scans.

| No. of patients | Median volume (ml) | IQR | Range (ml) | $p$ value\textsuperscript{a} |
|-----------------|-------------------|-----|------------|-----------------------------|
| $V_{CT}$        | 20                | 450 | 195        | 264–860                    | - |
| $V_{B-SCAN}$    | 20                | 398 | 240        | 264–821                    | - |
| $V_{1-1\text{ week}}$ | 20     | 319 | 234        | 147–929                    | 0.087 |
| $V_{1-2\text{ week}}$ | 20     | 306 | 146        | 60–734                     | 0.001 |
| $V_{1-3\text{ week}}$ | 20     | 270 | 135        | 50–561                     | 0.001 |
| $V_{2-1\text{ week}}$ | 20     | 371 | 161        | 136–642                    | 0.017 |
| $V_{2-2\text{ week}}$ | 20     | 300 | 154        | 219–536                    | 0.002 |
| $V_{2-3\text{ week}}$ | 20     | 358 | 196        | 173–488                    | 0.01 |
| $V_{3-1\text{ week}}$ | 20     | 348 | 166        | 124–634                    | 0.03 |
| $V_{3-2\text{ week}}$ | 20     | 268 | 62         | 147–606                    | <0.001 |
| $V_{3-3\text{ week}}$ | 20     | 317 | 179        | 126–557                    | 0.014 |
| $V_{4-1\text{ week}}$ | 20     | 333 | 144        | 117–528                    | 0.006 |
| $V_{4-2\text{ week}}$ | 20     | 252 | 182        | 134–532                    | 0.001 |
| $V_{4-3\text{ week}}$ | 20     | 272 | 122        | 84–482                     | 0.001 |
| $V_{5-1\text{ week}}$ | 20     | 336 | 140        | 147–485                    | 0.003 |
| $V_{5-2\text{ week}}$ | 20     | 277 | 182        | 98–528                     | <0.001 |
| $V_{5-3\text{ week}}$ | 20     | 282 | 130        | 137–511                    | <0.001 |
| $V_{6-1\text{ week}}$ | 17      | 284 | 65         | 130–436                    | 0.002 |
| $V_{6-2\text{ week}}$ | 8       | 335 | 130        | 216–560                    | 0.025 |
| $V_{6-3\text{ week}}$ | 7       | 258 | 266        | 149–448                    | 0.018 |

The bladder volume measurements and the number of patients adjusted by LOCF according to time sequence are described.

\textsuperscript{a} $p$ value between VB-SCAN and $V_{x-x\text{ week}}$

Abbreviations: IQR = interquartile range; VCT = bladder volume measured from simulation CT scans; VB-SCAN = bladder volume scanned by bladder scan prior to simulation CT scans; $V_{x-x\text{ week}}$ = bladder volume scanned by bladder scan at post-RT $x-x\text{ week}$.
### Table 5. Relative bladder volume change for each patient.

| Patient | Current study (protocol-based maintenance) | Previous study (self-controlled maintenance) |
|---------|---------------------------------------------|-----------------------------------------------|
|         | Median (%) | IQR (%) | Median (%) | IQR (%) |
| Case 1  | -19.66     | 52.52   | -35.10     | 17.79   |
| Case 2  | -40.72     | 26.17   | -56.15     | 19.49   |
| Case 3  | 1.71       | 30.82   | -67.60     | 21.81   |
| Case 4  | 4.55       | 61.09   | -66.95     | 24.36   |
| Case 5  | 5.30       | 27.46   | -68.24     | 31.42   |
| Case 6  | -27.45     | 42.84   | -55.50     | 31.92   |
| Case 7  | -53.96     | 21.16   | -41.99     | 33.12   |
| Case 8  | -31.15     | 20.49   | -9.17      | 34.86   |
| Case 9  | -27.42     | 18.02   | -26.67     | 39.63   |
| Case 10 | -38.17     | 36.98   | -65.69     | 41.42   |
| Case 11 | -18.35     | 24.73   | -26.90     | 42.95   |
| Case 12 | -51.83     | 26.19   | -76.04     | 45.64   |
| Case 13 | -24.04     | 35.91   | 23.03      | 51.69   |
| Case 14 | -41.90     | 34.29   | -33.12     | 57.25   |
| Case 15 | -35.75     | 49.74   | 0.00       | 66.41   |
| Case 16 | -22.91     | 34.97   | 41.27      | 76.05   |
| Case 17 | -15.52     | 34.07   | 77.55      | 96.60   |

The median values and interquartile ranges of relative bladder volume change of each patient from this and our previous study patients are described.

Abbreviations: IQR = interquartile range

doi:10.1371/journal.pone.0128791.t005

---

**Fig 2.** The distribution of interquartile range of relative bladder volume change according to protocol-based maintenance. The interquartile range of relative bladder volume change for patients receiving protocol-based full bladder maintenance (black squares) had smaller variation than that of patients performing self-controlled maintenance (open triangles).

doi:10.1371/journal.pone.0128791.g002
bladder volume change and protocol-based maintenance, we performed binary logistic regression. Upon logistic regression, the IQR of relative bladder volume change was significantly related to protocol-based maintenance [relative risk 1.045, 95% confidence intervals (CI) 1.004–1.087, \( p = 0.033 \)]. We calculated cut-off values of IQR of relative bladder volume change for protocol-based maintenance using ROC curve analysis (AUC 0.68, 95% CI 0.51–0.85) and determined 37% as a cut-off value (Table 6). Using Chi-square tests, the current study included significantly more patients with an IQR of relative bladder volume change less than 37% than our previous study (\( p = 0.025 \), Table 6).

### Discussion

The findings from this prospective study showed that although absolute values of bladder volume measured by bladder scans differed from bladder volume on simulation CT scan, there was significant positive correlation between bladder volumes of simulation CT and bladder ultrasound scans. Bladder scan is a useful and accurate device to measure bladder volume easily. During the entire RT course for rectal cancer, there was significant reduction and a wide range of variation in bladder volume compared to baseline bladder volume, despite protocol-based education, bladder filling training and continuous biofeedback. However, this study demonstrated that compared to our previous study, protocol-based full bladder maintenance could reduce the intra-patient variation of bladder volume during the RT course and improve the consistency of bladder volume.

This study has some special considerations. First, this study is an observation study. In addition, we used patients in the preliminary study as a control group, which may lower the quality of data in the control group due to the lack of experience using the bladder scanner in the initial stages of the preliminary study. However, bladder scans and CT bladder volumes were highly correlated in the control group, and patients in the control group fit the inclusion criteria of this study. Therefore, we consider our results are sufficient to show the importance of protocol-based full bladder maintenance. For a more accurate study, further randomized trials are necessary to study the effect of protocol-based maintenance. Second, basic bladder function tests were not performed on patients enrolled in this study. However, we excluded patients with micturition problems and no patient showed micturition problems during the study period. Any patient with mild or severe urinary frequency or nocturia was not included in this study. Thus, the absence of basic bladder function tests in this study likely did not significantly impact the results of this study. Third, there was no significant difference of median relative bladder volume change of each patient between two groups due to small number of enrolled patients. But, we consider that despite small number of patients, our findings showed less bladder volume reduction compared to initial planning and day-to-day variation through protocol-based bladder volume maintenance. Forth, in the era of intensity-modulated RT (IMRT), the protocol-based full bladder maintenance might be less applicable and important since IMRT could reduce radiation-induced toxicity for rectal cancer patients. Nevertheless, we consider that the addition of full

| Variables          | Current study (protocol-based maintenance) | Previous study (self-controlled maintenance) | \( p \) value |
|--------------------|---------------------------------------------|-----------------------------------------------|--------------|
| IQR of RBVC <37    | 15 (75.0)                                   | 8 (40.0)                                      | 0.025        |
| IQR of RBVC ≥37    | 5 (25.0)                                    | 12 (60.0)                                     |              |

The current study included significantly more patients with an interquartile range of relative bladder volume change less than 37% than our previous study. Abbreviations: IQR = interquartile range; RBVC = Relative bladder volume change

\[ \text{doi:10.1371/journal.pone.0128791.t006} \]
Bladder maintenance to IMRT could help minimize irradiated dose to organs at risk and radiation-induced toxicity in IMRT for rectal cancer. In this context, further study about the applicability of bladder scanner in IMRT for rectal cancer would be needed. Considering these limitations, properly designed randomized trial with more patients should be needed.

In a previous study from another institute, the usefulness of a bladder scan to achieve better reproducibility of bladder filling during pelvic RT was evaluated [16]. These authors showed that daily bladder volume variation was not significantly improved by biofeedback protocols by comparing the control and feedback group (47.2% and 40.1%, respectively, \( p = 0.2 \)). However, although not statistically significant, bladder volume during RT was better, and daily bladder volume variations were smaller, in the feedback group. This was perhaps due to reasons such as baseline micturition problems of prostate cancer patients. Similarly, another study of prostate patients also reported large bladder volume variance [19]. However, although a small number of patients were evaluated, we enrolled rectal cancer patients without baseline micturition problems according to strict inclusion and exclusion criteria. In addition, we used a more definitive and objective biofeedback protocol. Thus, in rectal cancer patients without baseline micturition problems, we consider that definitive protocol-based continuous biofeedback training of bladder filling using bladder ultrasound scan can improve bladder volume consistency during the RT course, thereby maintaining effective and safe treatment.

Our findings showed that bladder scans were fairly accurate in measuring bladder volume compared to bladder volume on simulation CT scan. Other previous studies have also reported that there was a strong correlation between bladder ultrasound scan and simulation CT scan [14–16]. Because CT scan cannot be performed every day or weekly during the entire treatment period, bladder scans can be utilized as an easy, convenient, and useful tool to check bladder volume and allow steady biofeedback training progress. However, we managed the full bladder maintenance protocol using bladder scans and improved bladder volume consistency during RT, and bladder volume during RT period was still significantly lower than baseline bladder volume. Ahmed et al. also reported that bladder volumes were dramatically reduced during treatment despite of full bladder training [14]. Thus, it is necessary to develop and evaluate additional methods or materials to reduce the differences in absolute bladder volume between baseline and the RT period.

**Conclusions**

In conclusion, we report that it would be possible to maintain bladder volume steadily during the entire RT course through protocol-based education, training, and continuous biofeedback and bladder scan utilization. Further study will be required to critically evaluate the influence of protocol-based full bladder maintenance, and additional methods need to be studied for improved consistency of bladder volume.

**Author Contributions**

Conceived and designed the experiments: HIY WSK. Performed the experiments: JSC WSK. Analyzed the data: HIY YC WSK. Contributed reagents/materials/analysis tools: HIY YC WSK. Wrote the paper: HIY YC WSK. Contributed to this work for prospective assessment and management of acute and late GI toxicities: SJP. Supervision of bladder scanner and management of acute and late urologic toxicities: JYL.

**References**

1. Navarro M, Dotor E, Rivera F, Sanchez-Rovira P, Vega-Villegas ME, Cervantes A, et al. A Phase II study of preoperative radiotherapy and concomitant weekly irinotecan in combination with protracted
Gastrointestinal Tumor Study Group. Prolongation of the disease-free interval in surgically treated rectal carcinoma. J Clin Oncol. 2006; 24(28):4620–5. Epub 2006/06/20. pii: JCO.2009.22.0467. doi: 10.1200/JCO.2009.22.0467. PMID: 19770376; PubMed Central PMCID: PMC2773471.

2. Sau R, Becker H, Hohenberger W, Rodel C, Wittekind C, Fietkau R, et al. Preoperative versus postoperative chemoradiotherapy for rectal cancer. The New England journal of medicine. 2004; 351(17):1731–40. Epub 2004/10/22. pii: S0140-6736(01)06409-1 PMID: 15496622.

3. Sau R, Liesch T, Merkel S, Fietkau R, Hohenberger W, Hess C, et al. Preoperative versus postoperative chemoradiotherapy for locally advanced rectal cancer: results of the German CAO/ARO/AIO-94 randomized phase III trial after a median follow-up of 11 years. J Clin Oncol. 2012; 30(16):1926–33. Epub 2012/04/25. pii: JCO.2011.40.1836. doi: 10.1200/JCO.2011.40.1836 PMID: 22529255.

4. Sebag-Montefiore D, Stephens RJ, Steele R, Monson J, Grieve R, Khanna S, et al. Preoperative radiotherapy versus selective postoperative chemoradiotherapy in patients with rectal cancer (MRC CR07 and NCIC-CTG C016): a multicentre, randomised trial. Lancet. 2009; 373(9666):811–20. Epub 2009/03/10. pii: S0140-6736(09)60484-0. doi: 10.1016/S0140-6736(09)60484-0 PMID: 19269519; PubMed Central PMCID: PMC2688947.

5. Colorectal Cancer Collaborative Group. Adjuvant radiotherapy for rectal cancer: a systematic overview of 8,507 patients from 22 randomised trials. Lancet. 2001; 358(9290):1291–304. Epub 2001/10/31. pii: S0140-6736(01)06409-1 PMID: 11694209.

6. Kavanagh BD, Pan CC, Dawson LA, Ressmeyer TK, Denekas D, Henrys AJ, Bedford J, Norman A, Tait DM. The effect of treatment position, proctitis, and NCIC-CTG C016: a randomised phase III trial after a median follow-up of 11 years. J Clin Oncol. 2010; 28(34):4610–6. Epub 2010/07/16. pii: JCO.2010.28.34.4610. doi: 10.1200/JCO.2010.28.34.4610. PMID: 20620417.

7. Drzymala M, Hawkins MA, Henrys AJ, Bedford J, Norman A, Tait DM. The effect of treatment position, proctitis, and NCIC-CTG C016: a randomised phase III trial after a median follow-up of 11 years. J Clin Oncol. 2010; 28(34):4610–6. Epub 2010/07/16. pii: JCO.2010.28.34.4610. doi: 10.1200/JCO.2010.28.34.4610. PMID: 20620417.

8. Kavanagh BD, Pan CC, Dawson LA, Das SK, Li XA, Ten Haken RK, et al. Radiation dose-volume effects in the stomach and small bowel. Int J Radiat Oncol Biol Phys. 2010; 76(3 Suppl):S101–7. Epub 2010/03/05. pii: S0140-6736(09)03286-6. doi: 10.1016/j.ijrobp.2009.06.071 PMID: 20171953.

9. Roh MS, Colangelo LH, O'Connell MJ, Yothers G, Deutsch M, Allegre CJ, et al. Preoperative multimodality therapy improves disease-free survival in patients with carcinoma of the rectum: NSABP R-03. J Clin Oncol. 2009; 27(31):5124–30. Epub 2009/09/23. pii: JCO.2009.22.0467. doi: 10.1200/JCO.2009.22.0467. PMID: 19770376; PubMed Central PMCID: PMC2773471.

10. Gerard JP, Conroy T, Bonnetain F, Bouche O, Chapelot O, Closon-Dejardin MT, et al. Preoperative radiotherapy with or without concurrent fluorouracil and leucovorin in T3-4 rectal cancers: results of FFCD 9203. J Clin Oncol. 2006; 24(28):4620–5. Epub 2006/06/03. pii: 24/28/4620. doi: 10.1200/JCO.2006.06.7629 PMID: 17008704.

11. Crook JE, Moertel CG, Gunderson LL, Wieand HS, Collins RT, Beart RW, et al. Effective surgical adjuvant therapy for high-risk rectal carcinoma. The New England journal of medicine. 1991; 324(11):709–15. Epub 1991/01/14. doi: 10.1056/NEJM199103131241101 PMID: 1997835.

12. Nijman MP, Boonstra H, Buenger J, Tans M,比較のための分離体検査の標準化。放射線治療と放射線物質の効果性を評価する放射線治療。Br J Radiol. 2009; 82(976):321–7. Epub 2009/02/04. pii: 57848689. doi: 10.1259/bjr/57848689 PMID: 19188240.

13. Hynds S, McGarry CK, Mitchell DM, Early S, Shum L, Stewart DP, et al. Assessing the daily consistency of bladder filling ultrasound Bladderscan device in men receiving radical conformal radiotherapy for prostate cancer. Br J Radiol. 2011; 84(1005):813–8. Epub 2010/12/17. pii: 10.1259/bjr/50048151 PMID: 21159811; PubMed Central PMCID: PMC3473792.

14. Ahmad R, Hoogeman MS, Quint S, Mens JW, de Pree I, Heijmen BJ. Interm-fraction bladder filling variations and time trends for cervical cancer patients assessed with a portable 3-dimensional ultrasound bladder scanner. Radiat Oncol. 2008; 39(2):172–9. Epub 2008/08/16. pii: S0167-8140(08)00367-8. doi: 10.1016/j.radonc.2008.07.005 PMID: 18703248.

15. O’Doherty UM, McNair HA, Norman AR, Miles E, Hooper S, Davies M, et al. Variety of bladder filling in patients receiving radical radiotherapy to the prostate. Radiother Oncol. 2006; 79(3):335–40. Epub 2006/06/20. pii: S0167-8140(06)00206-4. doi: 10.1016/j.radonc.2006.05.007 PMID: 16781790.

16. Stamm MR, van Lin EN, van der Vught LP, Kaanders JH, Visser AG. Bladder filling variation during radiation treatment of prostate cancer: can the use of a bladder ultrasound scanner and biofeedback optimize bladder filling? Int J Radiat Oncol Biol Phys. 2006; 65(2):371–7. Epub 2005/12/39. doi: 10.1016/j.ijrobp.2005.12.039 PMID: 16542790.

17. Chang JS, Yoon HI, Cha HJ, Chung Y, Cho Y, Keum KC, et al. Bladder filling variations during concurrent chemoradiotherapy and pelvic radiotherapy in rectal cancer patients: early experience of bladder...
volume assessment using ultrasound scanner. Radiat Oncol J. 2013; 31(1):41–7. Epub 2013/04/27. doi: 10.3857/roj.2013.31.1.41 PMID: 23620868; PubMed Central PMCID: PMC3633230.

18. Chung Y, Yoon HI, Keum KC, Kim JH, Choi WH, Nam KC, et al. Effect of belly board with bladder compression device on small bowel displacement from the radiotherapy field for rectal cancer. Onkologie. 2013; 36(5):241–6. doi: 10.1159/000350299 PMID: 23689217.

19. Fiorino C, Foppiano F, Franzone P, Broggi S, Castellone P, Marcenaro M, et al. Rectal and bladder motion during conformal radiotherapy after radical prostatectomy. Radiother Oncol. 2005; 74(2):187–95. Epub 2005/03/01. pii: S0167-8140(04)00470-0. doi: 10.1016/j.radonc.2004.10.002 PMID: 15734207.