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# **Defecation into clothing without forewarning and mean radiation dose to bowel and anal-sphincter among gynecological cancer survivors**

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**ABSTRACT**

*Background:* To analyze the relationship between mean radiation dose to the bowels and the anal sphincter and occurrence of “defecation into clothing without forewarning”, a specific and serious fecal incontinence symptom after gynecological radiotherapy. Additional potential risk factors associated with the symptom are explored.

*Material and Methods:* Data were collected for 519 eligible gynecological cancer survivors, treated with pelvic radiotherapy, with a median follow-up of 5.8 years, using a study-specific questionnaire and medical records. Correlations between defecation into clothing without forewarning and mean dose to organs at risk; the anal-sphincter region, the rectum, the sigmoid and the small intestines were investigated, also taking other risk factors into account.

*Results:* Twelve per cent reported having had the symptom at least once in the preceding six months. Mean doses  $> 50$  Gy to the anal-sphincter region, the rectum, the sigmoid and the small intestines were related to the occurrence of the symptom. Significantly associated risk factors were deliveries with high birth weight, heart failure and lactose and/or gluten intolerance. After adjusting for these factors, mean doses  $> 50$  Gy to the anal-sphincter region, the sigmoid and the small intestines remained related to the occurrence of the symptom.

*Conclusion:* Mean doses to the bowels and anal-sphincter region are related to the risk of defecation into clothing without forewarning in long-term gynecological cancer survivors treated with pelvic radiotherapy. Further radiobiological modeling may distinguish which organ(s) contribute most to development of the symptom.

## INTRODUCTION

Involuntary defecation into clothing is a devastating event remembered for many years. Among gynecological cancer survivors, we have described a symptom in which the survivor suddenly and unexpectedly defecates into the clothes (1), a defecation using the reflexes seen during a regular toilet visit. This symptom, here cited as “defecation into clothing without forewarning”, is thus a completely different symptom than having fecal leakage without the reflexes of defecation. The underlying conditions include a combination of decreased sensitivity for rectal filling as well as an increased activity of the bowel as seen in frequent defecation urgency. This endpoint is associated with a decreased sensitivity that entails not being able to sense the need to go to the toilet and defecate and it also includes an irritative component that is responsible for the sudden emptying of a large volume of stools.

Having the symptom “defecation into clothing without forewarning” may result in adverse psychological, sexual and social consequences (1-3). At present we lack dose-volume data that allow us to examine possible associations of dose-volume with this devastating symptom.

Fecal continence is maintained by the pelvic floor muscles, the anal sphincter and the rectum (4). Dysfunction leads to fecal incontinence and the anorectal function has been extensively evaluated after pelvic radiation therapy (5, 6). Weakness of the anal sphincter, reduction in rectal compliance, change in rectal sensitivity, and altered stools consistency have been reported among gynecological cancer survivors treated with pelvic radiation therapy (7, 8). It is evident that the mechanism leading to fecal incontinence after radiation therapy is complex (9) perhaps involving many organs. However, most attention thus far has been given to

the anal sphincter and the rectum. Other organs at risk (OARs) have not been studied as carefully as they should be in order to understand the mechanism. Furthermore, the concept of “fecal incontinence” as commonly used encompasses a variety of symptoms and thus may be too general. We have found that by *atomizing* the term “incontinence” into detailed patient-reported symptoms, it may be possible to reveal specific radiation pathophysiologies that otherwise would remain hidden. Information on dose to normal tissue and outcome data will help to predict the risk of normal-tissue injury and thereby guide the radiotherapist in the choice of competing treatment plans (10).

In Sweden unique personal identity numbers and official population-based registers offer excellent conditions to follow cancer survivors without selection-induced problems. In 2006 a population-based study was performed among 616 gynecological cancer survivors on late symptoms after pelvic radiation therapy. A matched control population of 344 non-irradiated women was included. Information was provided through a study-specific validated questionnaire comprising questions on demographics, physical and psychological symptoms, sexuality and social functioning. We found a higher prevalence of the symptom defecation into clothing without forewarning at least occasionally during the preceding six months among the survivors (12 %) than among the controls (less than 1 %) (2). In the present study we report on the relationship between mean dose to bowel organs at risk and the occurrence of this specific fecal incontinence symptom in 519 survivors for whom we had electronically stored external beam radiation therapy (EBRT) dose plans.

## **MATERIALS AND METHODS**

We have followed the hierarchical step-model (11) in the design of the data collection and the interpretation of the results. A detailed description of the study methodology has previously been published (2). In previous papers from our group on “defecation of stools into clothing without forewarning” we used the phrasing directly reported by the survivors i.e. “emptying of all stools into clothing without forewarning”.

The Regional Ethics Committee at the Karolinska Institute approved the study.

### *Survivors' characteristics*

Clinical data and three-dimensional (3D) EBRT plans including dose-volume histograms (DVHs) for the anal-sphincter region, the rectum, the sigmoid and the small intestines were collected for the 519 eligible gynecological cancer survivors. The treatments were given with curative intent at Radiumhemmet, Stockholm or at Jubileumskliniken, Gothenburg in 1991 to 2003. Endometrial cancer and cervical cancer were the most common diagnoses, 62 and 22 % respectively. In addition there were survivors of ovarian, fallopian tube, vaginal and vulvar cancers and uterine sarcomas. In addition to EBRT, approximately 90 % also had had surgery, 82 % brachytherapy and 27 % chemotherapy. The chemotherapy received was single cisplatinum, combination platinum-taxane, combination platinum-antracycline or other. Median time since completing EBRT was 5.8 years (range 2 -14 years).

### *Treatment planning and delivery*

EBRT 3D treatment planning was based on Computed Tomography (CT) scans performed prior to therapy. Scans were made in the supine position on a flat table top, using laser markers and conversion factors to electron density. Slice thickness was

usually 5-10 mm. The EBRT dose was prescribed either at the isocenter or as the mean dose to the target covering at least 95% of the planning target volume (12). The treatment was administered with linear accelerators or a racetrack accelerator with 6-50 MV photons using two opposing fields or a four-field box technique with prescribed daily fractions of 1.6, 1.8 or 2.0 Gy. EBRT treatment position was verified by portal image films and check-and-confirm systems.

Prescribed doses for endometrial cancer was 40-46 Gy and to uterine sarcomas 50 Gy. For cervical cancer similar techniques were used in an initial treatment phase and in a second phase a boost covering a smaller volume with a prescribed total dose of 55-70 Gy, depending if brachytherapy was added or not. Ovarian and Fallopian tube cancers had a prescribed dose of 20 Gy to the abdomen and an additional 20 Gy to a volume with lowered cranial margin.

Brachytherapy (BT) was applied using standardized techniques and applicator templates. The BT dose was prescribed according to local practice. Orthogonal x-ray images verified the position of the BT applicators. High-dose rate BT for endometrial cancer was prescribed at 5 Gy per fraction in two fractions or 3.75 Gy per fraction in three fractions. For cervical cancer low-dose rate BT was prescribed at 10.0-24.0 Gy per fraction in one to three fractions depending on tumor size and EBRT dose or as high-dose rate BT at 4.0 Gy per fraction in three fractions.

#### *Organs at risk and Dose-Volume Histograms*

In the present study we contoured four intestinal OARs; anal-sphincter region, rectum, sigmoid and small intestines. The “anal-sphincter region” was represented by the inner muscle layer of the sphincter up to the anal verge. The “rectum” was

depicted by its outer contour with filling extending from the anal verge to the recto-sigmoid junction. The “sigmoid colon” was contoured from where the rectum deviates from its mid-position to where it turns cranially in the left part of the abdomen connecting to the colon descendens. The “small intestines” were all visible small bowels in the small pelvic cavity to the caudal part of the sacroiliac joints. Continuous CT slices resulted in three-dimensional volumes (Figure 1) where the absorbed doses were calculated. The contouring was performed by two persons at each clinic under the supervision of senior oncologists (H.L. and A.-C.W.) during 2006 and 2007. Guidance was provided by a Contouring Manual with illustrations (on-line Appendix) and written instructions. The dose-volume histograms (DVHs) were exported for the four OARs for each patient using the TMS (Nucletron, Veenendaal, the Netherlands), Cadplan or Eclipse (Varian Medical Systems, Palo Alto, the USA) treatment planning systems.

### *Statistical analyses*

Cancer survivors were dichotomized into having had or not having had defecation into clothing without forewarning at least once during the preceding six months. We used risk ratios (RRs) and odds ratios (ORs) with 95% CI to compare symptom prevalence between the two groups. RRs and p-values were estimated through the log-binomial model. For multivariable modeling we used ORs estimated through logistic regression.

Characteristics of the study population and univariate RRs are given in Table 1 and Table 2.

In order to identify potential co-variables for the symptom, we performed an exploratory variable selection (Table 1) using logistic regression with forward selection. We used complete cases and  $\alpha=0.05$  as inclusion criteria (Table 3). Survivors were sorted into five mean EBRT dose intervals and symptom prevalence was calculated within each dose level for the OARs, using the previously identified factors for adjustments (Table 4). Exclusion of survivors who had received BT with Iridium  $> 11.25$  Gy or Radium +/- Cesium was made to test the impact of BT. Correlations between mean doses in the four OARs were calculated using the Spearman correlation coefficient.

Differences between mean DVHs of those with and without the symptom were assessed with pointwise t-tests for each dosebin. All tests were performed two-sided and at the 5 % significance level; individuals with missing data were excluded in each calculation. Calculations were performed using SAS software (version 9.2, SAS Institute Inc., Cary, NC, USA). The EBRT doses have been corrected to 2 Gy per fraction using the Linear-Quadratic model with an  $\alpha/\beta$ -ratio of 3 Gy (13).

## **RESULTS**

Demographic and clinical characteristics of the 63 gynecological cancer survivors with the symptom defecation into clothing without forewarning and the 456 survivors without the symptom occurring at least once the past six months are presented in Table 1 and in Table 2. Deliveries with high birth weight, cardiovascular disease, lactose intolerance, rheumatism and thrombosis were more frequent among survivors with the symptom.

Cervical cancer and uterine sarcomas were overrepresented compared to endometrial cancer, which was the reference level. Affected survivors had more often been treated with radiation as only treatment and usually to higher doses compared to survivors treated following surgery (Table 2).

Mean total EBRT dose was higher among survivors with the symptom than among those without the symptom and 52 % of the affected survivors had had a total dose > 45 Gy compared to 35 % of non-affected survivors. There was no difference regarding field technique or target area. BT was less common among survivors with the symptom. There was no statistically significant increased risk of developing the symptom with time after EBRT, OR = 1.03 (95% CI 0.94-1.12) per year.

The multivariable analyses identified three risk factors for the symptom “defecation into clothing without forewarning”, in addition to EBRT. Delivery of at least two children with birth weight exceeding four kg (RR=2.2, 95% CI 1.2-4.1), heart failure (RR=3.4, 95% CI 2.0-6.0), and lactose intolerance and/or gluten intolerance (RR=2.6, 95% CI 1.4-4.7) were significantly associated with a risk of having the symptom (Table 3).

Mean doses to the four OARs were too closely correlated to be included in the same regression analysis, with Spearman correlation coefficients ranging from 0.412 to 0.724. The prevalence of “defecation into clothing without forewarning” was higher among survivors with mean doses > 50 Gy for at least one of the OARs than among those with lower mean doses (Table 4). The corresponding unadjusted RRs and ORs for mean doses > 50 Gy were significantly increased. Adjustment for the risk factors resulted in losing the significantly increased OR for mean dose > 50 Gy to the rectum.

The risk of the outcome was highest among patients with 4 OARs with mean dose > 50Gy.

Further analyses were restricted to survivors treated with Iridium BT dose  $\leq 11.25$  Gy. All OARs with the exception of the rectum showed significantly increased ORs for mean doses > 50 Gy (anal sphincter: 27.5 (5.0-150.1), rectum: 9.4 (0.9-95.9), sigmoid: 8.8 (20-39.3), small intestines: 7.0 (1.5-32.5)). After adjusting for risk factors the significantly increased ORs for mean doses > 50 Gy remained for all OARs except for the rectum Gy (anal sphincter: 36.3 (6.0-221.7), rectum: 11.1 (0.9-136.6), sigmoid: 9.9 (2.0-48.2), small intestines: 9.3 (1.7-50.7)). The differences in average dose DVHs with p-values for EBRT for survivors with and without defecation into clothing without forewarning for the four OARs are presented in Figure 2. The DVHs for the anal-sphincter region were significantly separated ( $p < 0.05$ ) for doses in the interval of 34.5-66.5 Gy, for the rectum in 39.0-41.5 Gy and 45.0-68.0 Gy respectively, for the sigmoid in 38.0-70.0 Gy and for the small intestines in the interval of 45.5-50.5 and 53.0-69.5 Gy respectively.

## **DISCUSSION**

Our results show a dose-effect relationship between mean doses > 50 Gy to the anal-sphincter region, the rectum, the sigmoid, and the small intestines and the occurrence of defecation into clothing without forewarning among long-term gynecological cancer survivors treated with pelvic radiation therapy. The dose distributions for patients with and without the symptom were significantly separated for the studied OARs for intermediate and high doses indicating that the doses to these OARs is an important factor for the development of the symptom “defecation into clothing without forewarning”. In a recent publication we reported on a study of a subgroup of

the survivors treated without BT. The DVHs for these OARs were also significantly separated for intermediate and high doses (14).

To the best of our knowledge, these are the first studies investigating the relationship between dose-distribution data of EBRT to the bowels and the anal-sphincter region and the occurrence of involuntary defecation among gynecological cancer survivors treated with pelvic radiation therapy. In contrast, a vast number of studies have reported on the relationship between anorectal dose parameters and the risk of late fecal incontinence in prostate and other pelvic cancers (15). In a study by Fiorino *et al.*, dosimetric rectal data from 506 prostate cancer patients were analyzed, where rectal volume receiving  $\geq 40$  Gy ( $V_{40}$ ) and surgery were the strongest predictors of fecal incontinence defined as “use of pads” (16). Based on 641 prospectively scored (RTOG/EORTC scale) prostate cancer survivors Peeters *et al.* found fecal incontinence requiring pads to be associated with anal wall parameters (17). Similar results have previously been reported by our own group for prostate cancer, where a significant correlation between mean dose in the interval of 45-55 Gy to the anal-sphincter region and the risk of fecal incontinence was found (18).

Some researchers favor the hypothesis that symptoms may originate from specific anatomic regions. Smeenk *et al.* reported on urgency and incontinence, that originated from both the anal wall and rectal wall, while frequency seemed mostly associated with rectal wall dysfunction (19). In addition they found that dose-effect relations differed between the described symptoms. The importance of discriminating between different symptoms and their origin in order to increase specificity is supported by Heemsbergen *et al.*, who performed an anorectal dose-surface map analysis and found a dose-effect relation for fecal incontinence in the anal region and lower rectum (20).

In the study by Fonteyne *et al.*, the sigmoid colon was suggested as being co-responsible for the development of lower intestinal toxicity beside the anal sphincter and the rectum. They also found that the volume of the small bowel receiving doses in the range of 50-60 Gy is predictive for the development of late side-effects, which is in line with our results (21). In our recent paper we found steep dose-response relationships for the anal-sphincter, rectum, sigmoid and the small intestines and the development of “defecation into clothing without forewarning”. The mean doses to the OARs were however highly correlated with each other, and it is difficult to say if only one or if multiple organs are involved in the development of the symptom (14). On the other hand Andreyev *et al.*, have recently questioned the anatomically based approach arguing that symptoms originating from the pelvic area have multiple causes (22). We consider “defecation into clothing without forewarning” neither to be fecal incontinence nor a pure urgency symptom. This implies the engagement of all the investigated OARs and is in line with our result that the risk of the symptom “defecation into clothing without forewarning” is highest among patients with mean dose >50 Gy to all four of the investigated organs at risk. We strongly support the importance of proper diagnostic procedures and that increased knowledge of radiotherapy related atomized symptoms may lead to refinement of treatment and development of less radiotherapy induced long-term side-effects. We would like to note that the treatment technique used for the survivors included in the study was either opposing fields or four-field box. Nowadays, in order to decrease toxicity, it is recommended to use IMRT and VMAT for the EBRT treatment.

Normal tissue injury induced by ionizing radiation is thought to be a progressive process. However, there are reports showing both an increase and a decrease of rectal symptoms in prostate cancer survivors with time (23). In the present study we did not

find any statistically significant increase in risk for developing defecation into clothing without forewarning during follow up from 2-14 years after pelvic radiation therapy.

One of the strengths of this study is the large population-based survivor cohort. The use of unique personal identity numbers, public registers and the fact that all gynecological cancer patients in Sweden belong to two of six geographical catchment areas minimizes the risk of selection-induced problems. Interviewer-induced bias was avoided by the use of a numbered postal questionnaire, a practice that mimics the technique of blinding. Access to all medical records has ensured correct information regarding clinical characteristics. The large cohort in combination with the long follow-up time has enabled us to investigate the symptom prevalence over time. Our data were based on women under the age of 80 years, and the results may not be possible to generalize to older populations.

The major limitation of this study comprises the difficulties in estimating the contribution from BT. A variety of techniques were used regarding isotopes, applicators, anatomical arrangements and doses. In addition treatments were performed without the aid of 3D BT planning systems. Based on our clinical experience, we made the assumption that Iridium BT in the adjuvant setting to treat endometrial cancer patients postoperatively with a total prescribed dose of 10-11.25 Gy did not substantially affect the dose to the studied OARs. The survivors that were treated with this kind of BT had in general a low prescribed EBRT dose. We therefore excluded survivors receiving Iridium > 11.75 Gy, or Radium with or without Cesium, in an attempt to investigate the impact of BT given with “high” doses. The resulting

prevalence ratios were even higher indicating that EBRT to the four OARs is related to the occurrence of defecation into clothing without forewarning.

To improve the specificity the contouring of the OARs was made with a zero margin in order not to unintentionally include parts of other organs. Reviewing of the contouring was applied to decrease interobserver variety. A limitation of our study is the potential effect of organ motions and variations in set-up of patients since the contouring is based on pretreatment CT scans, which represents a static picture. We have previously reported that the position of the sigmoid may vary and has the largest deviation anteriorly but also that overlapping dose is most pronounced in the anterior rectal wall (24).

This study shows that mean dose to bowel organs and anal-sphincter region is related to the occurrence of “defecation into clothing without forewarning”. Our results suggest that not only the rectum and anal sphincter should be acknowledged in radiation therapy planning but also the sigmoid and the small intestines. The results should be taken into consideration when comparing competing dose plans. In addition there is a need of more advanced radiobiological modeling to further explore the contribution from each OAR for predicting the risk of “defecation into clothing without forewarning”.

## **REFERENCES**

1. Bergmark K, Åvall-Lundqvist E, Dickman PW, Henningsohn L, Steineck G. Patient-rating of distressful symptoms after treatment for early cervical cancer. *Acta Obstet Gynecol Scand.* 2002 May;81(5):443-50.

2. Lind H, Waldenström AC, Dunberger G, al-Abany M, Alevronta E, Johansson KA, et al. Late symptoms in long-term gynaecological cancer survivors after radiation therapy: a population-based cohort study. *Br J Cancer*. 2011 Sep 6;105(6):737-45.
3. Dunberger G, Lind H, Steineck G, Waldenström AC, Nyberg T, al-Abany M, et al. Fecal incontinence affecting quality of life and social functioning among long-term gynecological cancer survivors. *Int J Gynecol Cancer*. 2010 Apr;20(3):449-60.
4. Bajwa A, Emmanuel A. The physiology of continence and evacuation. *Best Pract Res Clin Gastroenterol*. 2009;23(4):477-85.
5. Maeda Y, Hoyer M, Lundby L, Norton C. Faecal incontinence following radiotherapy for prostate cancer: a systematic review. *Radiother Oncol*. 2011 Feb;98(2):145-53.
6. Putta S, Andreyev HJ. Faecal incontinence: A late side-effect of pelvic radiotherapy. *Clin Oncol (R Coll Radiol)*. 2005 Sep;17(6):469-77.
7. Yeoh E, Sun WM, Russo A, Ibanez L, Horowitz M. (4, 5). *Int J Radiat Oncol Biol Phys*. 1996 Jul 15;35(5):1003-10.
8. Dunberger G, Lind H, Steineck G, Waldenström AC, Onelöv E, Åvall-Lundqvist E. Loose stools lead to fecal incontinence among gynecological cancer survivors. *Acta Oncol*. 2011 Feb;50(2):233-42.
9. O'Brien PC. Radiation injury of the rectum. *Radiother Oncol*. 2001 Jul;60(1):1-14.
10. Bentzen SM, Constine LS, Deasy JO, Eisbruch A, Jackson A, Marks LB, et al. Quantitative Analyses of Normal Tissue Effects in the Clinic (QUANTEC): an introduction to the scientific issues. *Int J Radiat Oncol Biol Phys*. 2010 Mar 1;76(3 Suppl):S3-9.

11. Steineck G, Hunt H, Adolfsson J. A hierarchical step-model for causation of bias-evaluating cancer treatment with epidemiological methods. *Acta Oncol.* 2006;45(4):421-9.
12. ICRU. Prescribing, recording, and reporting photon beam therapy 1993.
13. Emami B, Lyman J, Brown A, Coia L, Goitein M, Munzenrider JE, et al. Tolerance of normal tissue to therapeutic irradiation. *Int J Radiat Oncol Biol Phys.* 1991 May 15;21(1):109-22.
14. Alevronta E, Lind H, Al-Abany M, Waldenström AC, Olsson C, Dunberger G, et al. Dose-response relationships for an atomized symptom of fecal incontinence after gynecological radiotherapy. *Acta Oncol.* 2012 Nov 1.
15. Fiorino C, Rancati T, Valdagni R. Predictive models of toxicity in external radiotherapy: dosimetric issues. *Cancer.* 2009 Jun 19;115(S13):3135-40.
16. Fiorino C, Fellin G, Rancati T, Vavassori V, Bianchi C, Borca VC, et al. Clinical and dosimetric predictors of late rectal syndrome after 3D-CRT for localized prostate cancer: preliminary results of a multicenter prospective study. *Int J Radiat Oncol Biol Phys.* 2008 Mar 15;70(4):1130-7.
17. Peeters ST, Lebesque JV, Heemsbergen WD, van Putten WL, Slot A, Dielwart MF, et al. Localized volume effects for late rectal and anal toxicity after radiotherapy for prostate cancer. *Int J Radiat Oncol Biol Phys.* 2006 Mar 15;64(4):1151-61.
18. al-Abany M, Helgason AR, Cronqvist AK, Lind B, Mavroidis P, Wersäll P, et al. Toward a definition of a threshold for harmless doses to the anal-sphincter region and the rectum. *Int J Radiat Oncol Biol Phys.* 2005 Mar 15;61(4):1035-44.
19. Smeenk RJ, Hopman WP, Hoffmann AL, van Lin EN, Kaanders JH. Differences in Radiation Dosimetry and Anorectal Function Testing Imply That

Anorectal Symptoms May Arise from Different Anatomic Substrates. *Int J Radiat Oncol Biol Phys.* 2010 Oct 13:1-8.

20. Heemsbergen WD, Hoogeman MS, Hart GA, Lebesque JV, Koper PC. Gastrointestinal toxicity and its relation to dose distributions in the anorectal region of prostate cancer patients treated with radiotherapy. *Int J Radiat Oncol Biol Phys.* 2005 Mar 15;61(4):1011-8.

21. Fonteyne V, De Neve W, Villeirs G, De Wagter C, De Meerleer G. Late radiotherapy-induced lower intestinal toxicity (RILIT) of intensity-modulated radiotherapy for prostate cancer: the need for adapting toxicity scales and the appearance of the sigmoid colon as co-responsible organ for lower intestinal toxicity. *Radiother Oncol.* 2007 Aug;84(2):156-63.

22. Andreyev HJ, Wotherspoon A, Denham JW, Hauer-Jensen M. "Pelvic radiation disease": new understanding and new solutions for a new disease in the era of cancer survivorship. *Scand J Gastroenterol.* 2011 Apr;46(4):389-97.

23. Odrazka K, Dolezel M, Vanasek J, Vaculikova M, Zouhar M, Sefrova J, et al. Time course of late rectal toxicity after radiation therapy for prostate cancer. *Prostate Cancer Prostatic Dis.* 2009 Jun;13(2):138-43.

24. Waldenström AC, Alsadius D, Pettersson N, Johansson KA, Holmberg E, Steineck G, et al. Variation in position and volume of organs at risk in the small pelvis. *Acta Oncol.* 2010 May;49(4):491-9.

**Table 1. Demographic and Clinical Characteristics for 519 Gynecological Cancer Survivors with and without Defecation into Clothing without Forewarning after Pelvic Radiation Therapy**

	Survivors with Defecation into Clothing without Forewarning	Survivors without Defecation into Clothing without Forewarning	Unadjusted Relative Risks	p-value*
	N=63 n (%)	N=456 n (%)	RR (95 % CI)	
Age				0.108
28-49	8 (15)	47 (85)	1.0	
50-59	4 (5)	78 (95)	0.3 (0.1-1.1)	
60-69	25 (13)	173 (87)	0.9 (0.4-1.8)	
70-79	26 (14)	158 (86)	1.0 (0.5-2.0)	
Smoking				0.461
Never smoker	28 (12)	215 (88)	1.0	
Former smoker	17 (11)	139 (89)	0.9 (0.5-1.7)	
Current smoker	18 (16)	97 (84)	1.4 (0.8-2.4)	
BMI				0.37
< 18.5 (underweight)	2 (14)	12 (86)	1.0 (0.3-3.7)	
18.5-25.0 (normal weight)	31 (14)	185 (86)	1.0	
25.0-30.0 (overweight)	15 (9)	157 (91)	0.6 (0.3-1.1)	
≥ 30.0 (obese)	12 (14)	72 (86)	1.0 (0.5-1.8)	
No. of births				0.826
0	15 (11)	118 (89)	1.0	
1-3	44 (13)	302 (87)	1.1 (0.7-2.0)	
> 3	4 (10)	36 (90)	0.9 (0.3-2.5)	
Delivery				
Fast < 5 hours, no	35 (11)	275 (89)	1.0	
Fast < 5 hours, yes	27 (13)	174 (87)	1.2 (0.7-1.9)	0.471
Slow > 24 hours, no	45 (11)	348 (89)	1.0	
Slow > 24 hours, yes	17 (14)	101 (86)	1.3 (0.7-2.1)	0.397
Vacuum, no	56 (12)	419 (88)	1.0	
Vacuum, yes	6 (17)	30 (83)	1.4 (0.7-3.1)	0.408
Forceps, no	61 (12)	440 (88)	1.0	
Forceps, yes	1 (10)	9 (90)	0.8 (0.1-5.4)	0.830
Episiotomy, no	48 (12)	354 (88)	1.0	
Episiotomy, yes	14 (13)	95 (87)	1.1 (0.6-1.9)	0.799
Caesarian, no	60 (12)	428 (88)	1.0	
Caesarian, yes	2 (9)	21 (91)	0.7 (0.2-2.7)	0.589
Breech birth, no	60 (12)	433 (88)	1.0	
Breech birth, yes	2 (11)	16 (89)	0.9 (0.2-3.4)	0.891
Delivery with birth weight > 4 kg				0.045
0	49 (12)	367 (88)	1.0	
1	5 (8)	60 (92)	0.7 (0.3-1.6)	
≥ 2	9 (26)	26 (74)	<b>2.2 (1.2-4.1)</b>	
Pelvic floor injury				
Vaginal and perineal injury, no	45 (11)	368 (89)	1.0	
Vaginal and perineal injury,	16 (17)	78 (83)	1.6 (0.9-2.6)	0.113
Anal-sphincter injury, no	57 (12)	433 (88)	1.0	
Anal-sphincter injury, yes	4 (29)	10 (71)	<b>2.5 (1.0-5.8)</b>	0.093

Abbreviations: CI = confidence interval, BMI = body mass index, EBRT = external beam radiation therapy

Significant results in bold. \*p-values from log-binomial model type III tests

\*Angina pectoris, cardiac infarction

**Table 1 (continued). Demographic and Clinical Characteristics for 519 Gynecological Cancer Survivors with and without****Defecation into Clothing without Forewarning after Pelvic Radiation Therapy**

Co-morbidities				
Previous abdominal surgery, no	31 (11)	261 (89)	1.0	
Previous abdominal surgery, yes	32 (14)	196 (86)	1.3 (0.8-2.1)	0.230
Diabetes mellitus, no	54 (12)	413 (88)	1.0	
Diabetes mellitus, yes	9 (18)	42 (82)	1.5 (0.8-2.9)	0.230
Heart failure, no	50 (10)	432 (90)	1.0	
Heart failure, yes	10 (36)	18 (64)	<b>3.4 (2.0-6.0)</b>	0.001
Other cardiovascular disease*, no	31 (10)	275 (90)	1.0	
Other cardiovascular disease, yes	29 (14)	175 (86)	1.4 (0.9-2.3)	0.164
Crohn's disease and or ulcerative colitis, treatment	57 (12)	424 (88)	1.0	
Crohn's disease and or ulcerative colitis, treatment	2 (33)	4 (67)	2.8 (0.9-9.0)	0.612
Irritable bowel syndrome, treatment for, no	52 (11)	428 (89)	1.0	
Irritable bowel syndrome, treatment for, yes	5 (20)	20 (80)	1.8 (0.8-4.2)	0.194
Hemorrhoids, treatment for, no	54 (12)	398 (88)	1.0	
Hemorrhoids, treatment for, yes	5 (10)	45 (90)	0.8 (0.4-2.0)	0.679
Lactose intolerance and/or gluten intolerance, no	53 (11)	429 (89)	1.0	
Lactose intolerance and/or gluten intolerance, yes	9 (28)	23 (72)	<b>2.6 (1.4-4.7)</b>	0.011
Pelvic organ prolapse, treatment for, no	60 (12)	436 (88)	1.0	
Pelvic organ prolapse, treatment for, yes	1 (9)	9 (91)	0.8 (0.1-4.9)	0.753
Rheumatism, no	52 (11)	424 (89)	1.0	
Rheumatism, yes	8 (24)	26 (76)	<b>2.2 (1.1-4.2)</b>	0.046
Neurological disorders <sup>†</sup> , no	58 (12)	440 (88)	1.0	
Neurological disorders, yes	2 (17)	10 (83)	1.4 (0.4-5.2)	0.612

Abbreviations: CI = confidence interval, BMI = body mass index, EBRT = external beam radiation therapy

Significant results in bold

\*Angina pectoris, cardiac infarction and hypertension

<sup>†</sup> Parkinson's disease, multiple sclerosis and epilepsy

**Table 2. Diagnosis and Treatment Characteristics for 519 Gynecological Cancer Survivors with and without Defecation into Clothing without Forewarning after Pelvic Radiation Therapy**

	Survivors with Defecation into Clothing without Forewarning	Survivors without Defecation into Clothing without Forewarning	Unadjusted Relative Risks	p-value*
	N=63 n (%)	N=456 n (%)	(95 % CI)	
Diagnoses				0.092
Endometrial cancer	30 (9)	292 (91)	1.0	
Cervical cancer	19 (17)	93 (83)	<b>1.8 (1.1-3.1)</b>	
Ovarian and Fallopian tube cancer	4 (10)	38 (90)	1.0 (0.4-2.8)	
Uterine Sarcoma	7 (26)	20 (74)	<b>2.8 (1.4-5.7)</b>	
Vaginal cancer	2 (18)	9 (82)	2.0 (0.5-7.2)	
Vulvar cancer	1 (20)	4 (80)	2.1 (0.4-12.8)	
Stage of malignancy				0.075
I	29 (9)	286 (91)	1.0	
II	22 (18)	98 (82)	<b>2.0 (1.2-3.3)</b>	
III	10 (14)	63 (86)	1.5 (0.8-2.9)	
IV	0 (0)	3 (100)	0.0 (-)	
Unknown	2 (25)	6 (75)	2.7 (0.8-9.5)	
Surgery				0.006
None	15 (29)	37 (71)	1.0	
TAH +/- SOE +/- omentectomy	42 (11)	333 (89)	<b>0.4 (0.2-0.6)</b>	
TAH +/-SOE +/- omentectomy +/- lymph node sampling	1 (4)	27 (96)	<b>0.1 (0.0-0.9)</b>	
Radical hysterectomy + pelvic lymphadenectomy	4 (7)	50 (93)	<b>0.3 (0.1-0.7)</b>	
Vulvar resection +/- lymph node resection	1 (20)	4 (80)	0.7 (0.1-4.2)	
Other	0 (0)	5 (100)	0.0 (-)	
EBRT				0.011
Doses (Gy)				
≤ 40	22 (9)	217 (91)	1.0	
40-45	8 (9)	79 (91)	1.0 (0.5-2.2)	
45-55	26 (15)	147 (85)	<b>1.6 (1.0-2.8)</b>	
> 55	7 (35)	13 (65)	<b>3.8 (1.8-7.8)</b>	
Field technique				0.682
Four-field box	50 (12)	371 (88)	1.0	
Two opposing fields	13 (13)	84 (87)	1.1 (0.6-2.0)	
Target area				0.774
Pelvic field	49 (12)	364 (88)	1.0	
Abdominal field	9 (15)	52 (85)	1.2 (0.6-2.4)	
Pelvic field + paraaortic lymph nodes	3 (9)	31 (91)	0.7 (0.2-2.3)	
Pelvic or vulvar field + inguinal lymph nodes	2 (18)	9 (82)	1.5 (0.4-5.5)	
Brachytherapy isotope				0.052
Ir	35 (10)	327 (90)	1.0	
Ra +/- Cs	0 (0)	15 (100)	0.0 (-)	
Cs	9 (18)	42 (82)	1.8 (0.9-3.6)	
Time relation				0.056
No brachytherapy	19 (21)	72 (79)	1.0	
Preoperative	2(6)	32 (94)	0.3 (0.1-1.1)	
Postoperative	33 (9)	319 (91)	<b>0.4 (0.3-0.8)</b>	
During EBRT (irradiation alone)	9 (21)	33 (79)	1.0 (0.5-2.1)	
Treatment				0.385
Chemotherapy, no	38 (11)	315 (89)	1.0	
Chemotherapy concurrent, yes	4 (14)	24 (86)	1.3 (0.5-3.5)	

Chemotherapy sequential, yes	21 (15)	117 (85)	1.4 (0.9-2.3)	0.026
Brachytherapy, no	19 (21)	72 (79)	1.0	
Brachytherapy dose Ir $\leq$ 0-11.25	33 (10)	302 (90)	0.5 (0.3-0.8)	
	<b>Survivors with Defecation into Clothing without Forewarning</b>	<b>Survivors without Defecation into Clothing without Forewarning</b>	<b>Unadjusted Relative Risks</b>	<b>p-value*</b>
Brachytherapy dose Ir > 11.25 Gy or Ra +/- Cs	11 (12)	82 (88)	0.6 (0.3-1.1)	0.147
Time since EBRT treatment (years)				
2-5	19 (9)	184 (91)	1.0	
5-10	36 (15)	201 (85)	<b>1.6 (1.0-2.7)</b>	
10-14	8 (10)	71 (90)	1.1 (0.5-2.4)	

*Abbreviations:* CI = confidence interval, BMI = body mass index, EBRT = external beam radiation therapy, Ir = Iridium, Ra = Radium, Cs = Cesium

Significant results in bold. \*p-values from log-binomial model type III tests

**Table 3. Potential Risk factors for Defecation into Clothing without Forewarning after Pelvic Radiation Therapy in 519 Gynecological Cancer Survivors**

	<b>Defecation into Clothing without Forewarning n/N (%)</b>	<b>Unadjusted RR (95 % CI)</b>	<b>Unadjusted OR (95 % CI)</b>	<b>Adjusted OR* (95 % CI)</b>
<b>Delivery with birth weight &gt; 4 kg</b>				
0	49/416 (12)	1.0	1.0	1.0
1	5/65 (8)	0.7 (0.3-1.6)	0.6 (0.2-1.6)	0.7 (0.2-1.7)
≥ 2	9/35 (26)	<b>2.2 (1.2-4.1)</b>	<b>2.6 (1.1-5.9)</b>	<b>2.6 (1.0-6.3)</b>
Missing values	3			
<b>Heart failure</b>				
No	50/482 (10)	1.0	1.0	1.0
Yes	10/28 (36)	<b>3.4 (2.0-6.0)</b>	<b>4.8 (2.1-11.0)</b>	<b>4.1 (1.7-10.0)</b>
Missing values	9			
<b>Lactose intolerance and/or gluten</b>				
No	53/482 (11)	1.0	1.0	1.0
Yes	9/32 (28)	<b>2.6 (1.4-4.7)</b>	<b>3.2 (1.4-7.2)</b>	<b>2.9 (1.2-6.9)</b>
Missing values	5			

*Abbreviations:* RR = relative risk, CI = confidence interval, OR = odds ratio

Significant results in bold

\*Adjustment for high birth weight, heart failure and lactose and/or gluten intolerance

**Table 4. Mean Dose to Organs at Risk and Risk of Defecation into Clothing without Forewarning after Pelvic Radiation therapy**

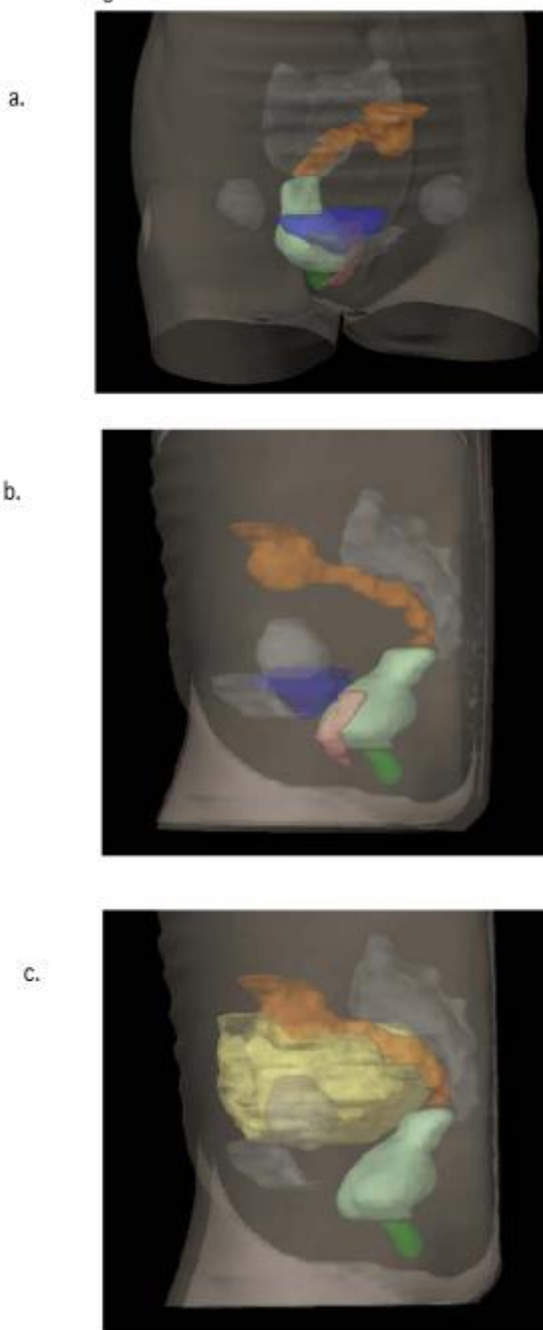
		All survivors No=519		
Mean absorbed EBRT dose to OARs (Gy)	n/N (%)	Unadjusted RR (95 % CI)	Unadjusted OR (95 % CI)	Adjusted OR* (95 % CI)
<b>Anal-sphincter region</b>				
≤ 35	27/317 (9)	1.0	1.0	1.0
> 35-40	14/92 (15)	<b>1.8 (1.0-3.3)</b>	<b>1.9 (1.0-3.9)</b>	<b>2.3 (1.1-4.7)</b>
> 40-45	12/79 (15)	1.9 (0.9-3.4)	1.9 (0.9-4.0)	<b>2.3 (1.0-5.1)</b>
> 45-50	3/28 (17)	2.0 (0.7-5.8)	2.1 (0.6-7.9)	2.8 (0.7-10.6)
> 50	7/11 (64)	<b>7.5 (4.2-13.3)</b>	<b>18.8 (5.2-68.3)</b>	<b>22.7 (5.7-89.7)</b>
<b>Rectum</b>				
≤ 35	3/45 (7)	1.0	1.0	1.0
> 35-40	23/217 (11)	1.6 (0.5-5.1)	1.7 (0.5-5.8)	1.6 (0.4-6.1)
> 40-45	10/91 (11)	1.6 (0.5-5.7)	1.7 (0.5-6.6)	1.9 (0.4-8.1)
> 45-50	18/136 (13)	2.0 (0.6-6.4)	2.1 (0.6-7.6)	2.6 (0.7-10.4)
> 50	9/27 (33)	<b>5.0 (1.5-16.9)</b>	<b>7.0 (1.7-29.0)</b>	6.8 (0.4-6.1)
<b>Sigmoid</b>				
≤ 35	4/62 (6)	1.0	1.0	1.0
> 35-40	21/224 (9)	1.5 (0.5-4.1)	1.5 (0.5-4.5)	1.3 (0.4-4.0)
> 40-45	19/151 (13)	2.0 (0.9-8.2)	2.1 (0.7-6.4)	2.3 (0.7-7.3)
> 45-50	11/62 (18)	2.8 (0.9-8.2)	3.1 (0.9-10.4)	3.2 (0.9-11.1)
> 50	8/19 (42)	<b>6.5 (2.2-19.3)</b>	<b>10.5 (2.7-41.2)</b>	<b>9.3 (2.2-39.2)</b>
<b>Small intestines</b>				
≤ 35	4/56 (7)	1.0	1.0	1.0
> 35-40	25/228 (11)	1.5 (0.6-4.2)	1.6 (0.5-4.8)	1.5 (0.5-4.9)
> 40-45	12/134 (9)	1.3 (0.4-3.7)	1.3 (0.4-4.2)	1.3 (0.4-4.8)
> 45-50	14/64 (22)	<b>3.1 (1.1-8.8)</b>	<b>3.6 (1.1-11.8)</b>	<b>4.2 (1.2-14.9)</b>
> 50	7/20 (35)	<b>4.9 (1.6-15.0)</b>	<b>7.0 (1.8-27.6)</b>	<b>7.1 (1.6-31.9)</b>
<b>Number of OARs with mean dose &gt; 50 Gy</b>				
0	49/456 (11)	1.0	1.0	1.0
1	4/15 (27)	2.5 (1.0-6.0)	3.0 (0.9-9.9)	2.6 (0.7-9.1)
2	1/7 (14)	1.3 (0.2-8.3)	1.4 (0.2-11.7)	2.2 (0.3-19.1)
3	2/11 (18)	1.7 (0.5-6.1)	1.8 (0.4-8.8)	2.5 (0.5-12.2)
4	6/9 (67)	<b>6.2 (3.6-10.6)</b>	<b>16.6 (4.0-68.5)</b>	<b>18.5 (4.1-83.1)</b>

Abbreviations: EBRT = external beam radiation therapy, OAR = organ at risk, Ir = Iridium, BT = brachytherapy, RR = relative risk, CI = confidence interval, OR = odds ratio

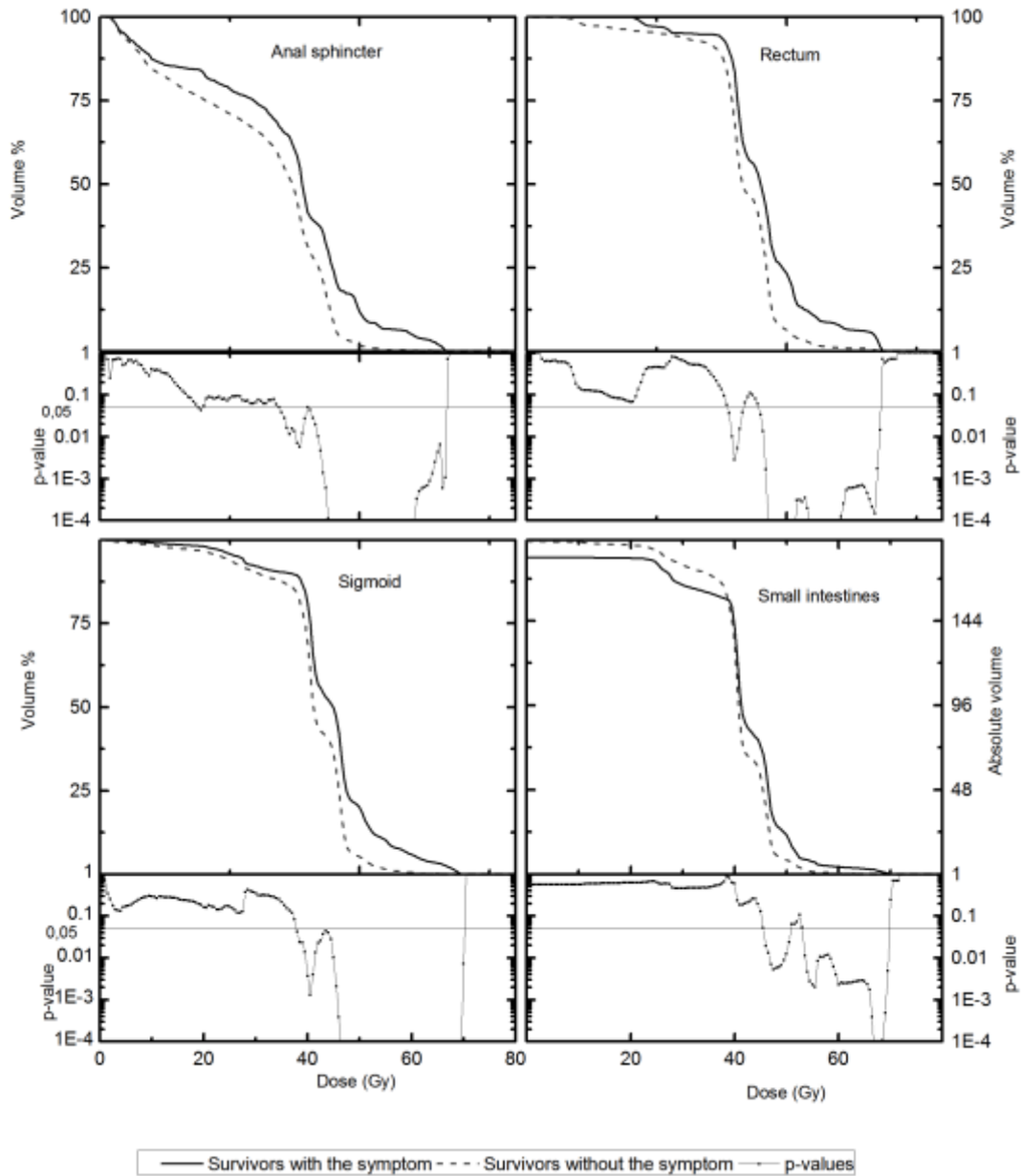
Significant results in bold. Doses corrected to 2 Gy per fraction

\*Adjustment for high birth weight, heart failure and lactose and/or gluten intolerance

Figure 1



**Figure 1.** a. Organs at risk, frontal view b. Organs at risk, lateral view c. Organs at risk, lateral view; Dark green – anal sphincter; Light green – rectum. Orange – sigmoid; Yellow – small intestines. Pink – vagina. Blue – urinary bladder; Grey – sacrum, femoral heads, pubic bone



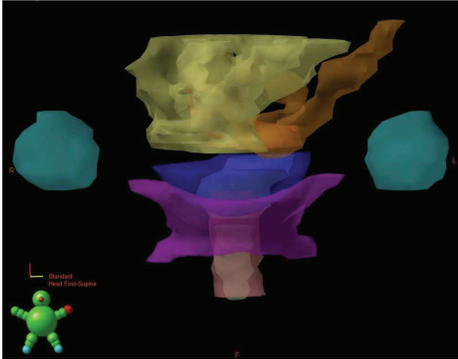
**Figure 2.** Mean dose-volume histograms over the anal-sphincter region, the rectum, the sigmoid and the small intestines with p-values. DVH for small intestines shows absolute volumes, while for the other structures relative volumes are shown.

**Gynecological Cancer Survivors Treated with  
Pelvic External Beam Radiation Therapy  
in Stockholm and Gothenburg  
Sweden 1991-2003**

Contouring Manual for Pelvic Organs at risk

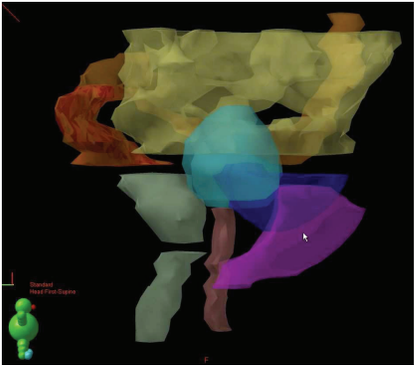
1

Pelvic organs at risk



2

Organs at risk small pelvis right lateral view



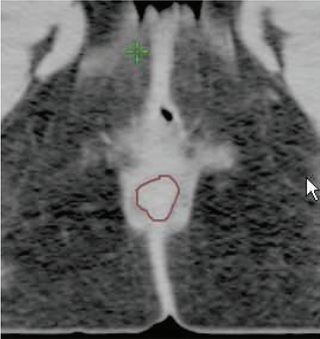
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Anal-sphincter region caudal border



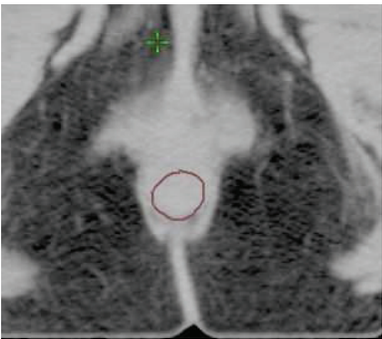
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Anal-sphincter region



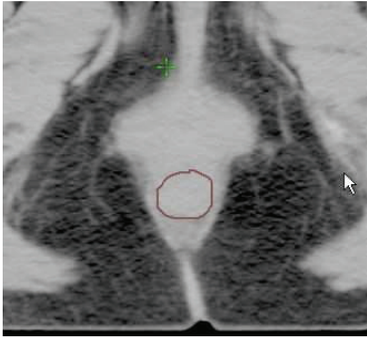
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Anal-sphincter region



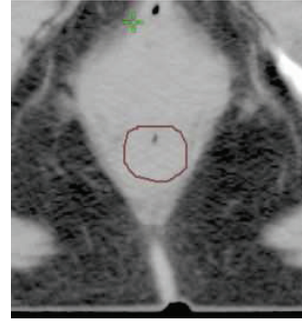
6

Anal-sphincter region



7

Anal-sphincter region cranial border



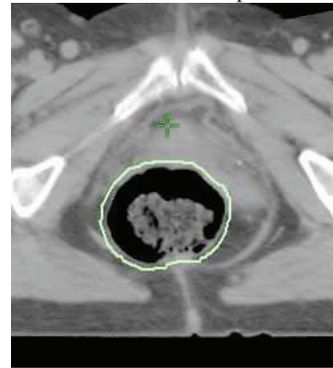
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Rectum caudal border



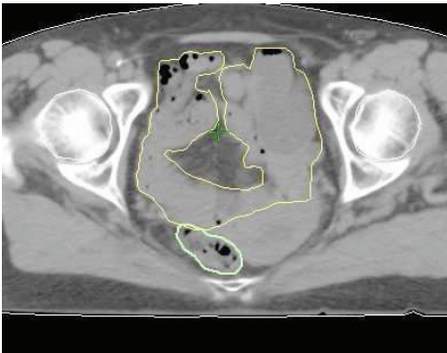
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Rectum middle part



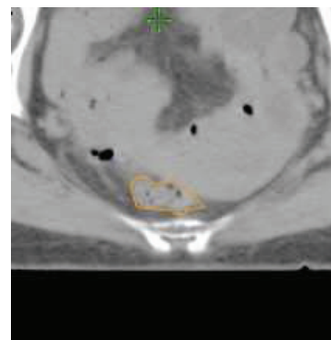
10

Rectum cranial border (green)



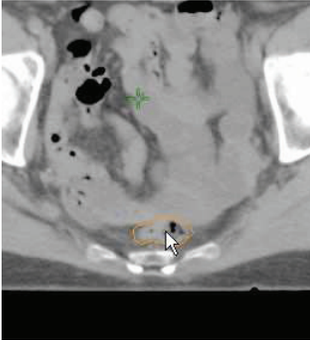
11

Sigmoid



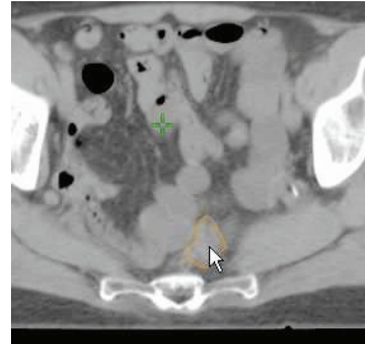
12

Sigmoid



13

Sigmoid



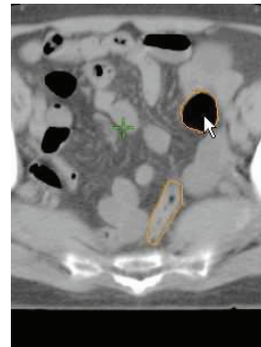
14

Sigmoid



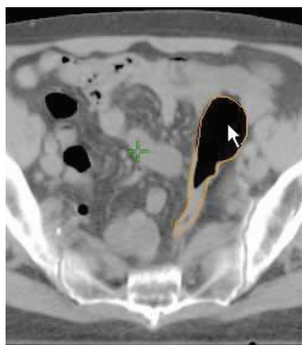
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Sigmoid



16

Sigmoid



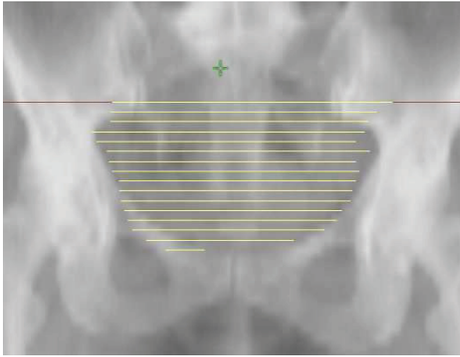
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Sigmoid



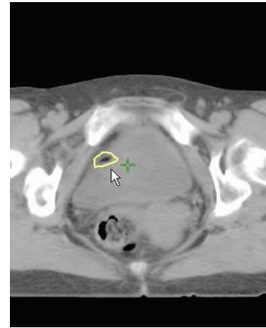
18

Small intestines in pelvis minor frontal view



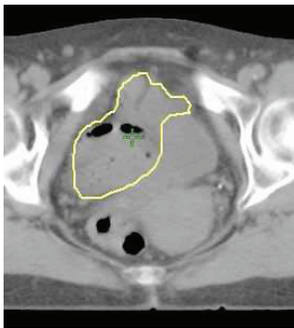
19

Small intestines caudal border



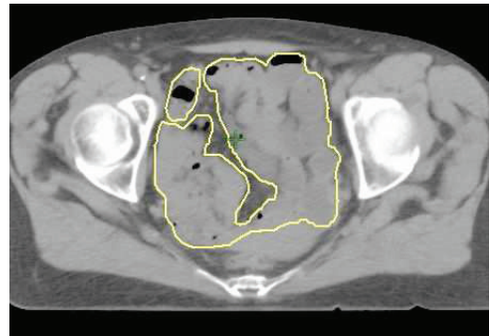
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Small intestines



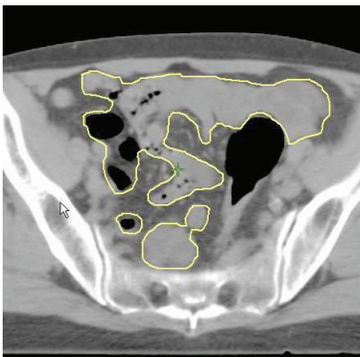
21

Small intestines



22

Small intestines pelvis minor cranial border



23

Vagina, urinary bladder, rectum and anal-sphincter region right lateral view



24