Radiotherapy dose-distribution to the perirectal fat space (PRS) is related to gastrointestinal control-related complications

S.L. Gulliford a,*, S. Ghose b, M.A. Ebert c,d, A. Kennedy c, J. Dowling e, J. Mitra b, D.J. Joseph c,f, J.W. Denham g

a Joint Department of Physics, Institute of Cancer Research and Royal Marsden National Health Service Foundation Trust, Sutton, United Kingdom
b School of Medicine and Public Health, University of Newcastle, New South Wales, Australia
c School of Surgery, University of Western Australia, Western Australia, Australia
d Australian e-Health Research Centre, CSIRO, Brisbane, Queensland, Australia
e School of Radiography, University of Western Australia, Western Australia, Australia
f School of Medicine and Public Health, University of Newcastle, New South Wales, Australia

e-mail address: sarahg@icr.ac.uk (S.L. Gulliford).

Original Research Article

Introduction

The range of rectal/bowel symptoms reported following prostate radiotherapy is diverse including rectal bleeding and control-related symptoms such as loose stools and urgency. The dosimetric relationship to the specific toxicity of rectal bleeding has been comprehensively studied and characterised [1]. For other endpoints the aetiology and relationship with dosimetry is less well defined and the subject of ongoing investigations [2–4]. How ever several studies reporting the rectal toxicity from large prospective clinical trials found differences in the anatomical sub-regions and dosimetric variables which related to individual toxicity outcomes [5–9]. A study by Smeenk et al. which considered the dosimetric relationship between the anal wall and pelvic floor muscle groups and incontinence-related toxicity demonstrated specific dose–response relationships with individual muscle groups [10]. Buettner at al [11] demonstrated that spatial descriptors of the dose received by the surface of anal canal (defined as the caudal 3 cm of the anorectum) were correlated to sphincter control (LENT SOM) [12]. It is apparent that different manifestations of toxicity are related to different underlying pathophysiology, including inflammatory responses and epithelial damage [9,10].

It is well recognised that rectal dose volume histograms (DVHs) obtained during prostatic irradiation differ from those derived during the radiotherapy planning process [13]. However, the surrounding region, the perirectal fat space (PRS), is thought to remain relatively immobile. If this is true, then it may also be true that the DVH of the PRS derived during planning will correlate more satisfactorily with subsequent radiation induced dysfunctional rectal symptoms than the rectal DVH generated during planning.

Moreover, if peri-rectal fat is as radiosensitive as other fatty tissue regions in the body, it is possible that a course of prostatic irradiation will reduce the elasticity of peri-rectal fat, which may in its own right alter rectal function adversely. Therefore, in this study...
we test the hypothesis that DVHs of the PRS obtained at planning correlates better with the severity of dysfunctional rectal symptoms and their underlying injuries than planning rectal DVHs.

**Methods and materials**

**Data source and description**

The RADAR trial (Randomised Androgen Deprivation and Radiotherapy, TROG 03.04) [14] examined the influence of duration of androgen deprivation (AD) with or without bisphosphonates, adjuvant with radiation therapy, for treatment of prostate carcinoma. 1071 participants were accrued from 23 centres across Australia and New Zealand between 2003 and 2007.

All participants received centre-nominated radiation therapy to the prostate with 46 Gy 3D conformal external beam radiation therapy (EBRT – “Phase 1”) followed by either a 19.5-Gy high dose-rate (HDR) brachytherapy boost or EBRT to either 66, 70, 74, or 78 Gy (at clinician discretion – “Phase 2”). Phase 1 was determined by PTV1, being CTV plus 10 mm margin in all directions except posteriorly where it was 5 mm. Phase 2 was determined by PTV2, being CTV plus 0–10 mm margin in all directions except posteriorly where it was 0–5 mm. Fractionation is shown in Table 1. No participants receiving the HDR boost were considered in this current study. Image guidance was via bony anatomy only.

Rectal dose volume constraints, derived from results presented by Boersma et al. [15], were applied during treatment planning. They were 65, 70, and 75 Gy to a maximum 40%, 30%, and 5% of rectum, respectively.

All patients were assessed at randomization (baseline) and then routinely followed in clinic every 3 months for 18 months, then at 6 months up to 5 years post randomization and then annually. At these visits, toxicity was assessed according to Late Effects of Normal Tissues Subjective, Objective, Management, and Analytic (LENT SOMA) scales [12]. Participant treatment planning data (CT images, planned dose distributions, delineate anatomy, beam configurations and treatment demographic data) were archived in a database using the SWAN software system [16], enabling subsequent query and arbitrary analysis.

| Trial arm½, n (%) | Whole group | No control-related rectal-toxicity | Control-related rectal toxicity |
|-------------------|-------------|---------------------------------|-------------------------------|
| N                 | 68          | 34                              | 34                            |
| **Trial arm**     |             |                                 |                               |
| A                 | 17 (25%)    | 9 (26%)                         | 8 (24%)                       |
| B                 | 17 (25%)    | 11 (32%)                        | 6 (18%)                       |
| C                 | 16 (24%)    | 6 (18%)                         | 10 (29%)                      |
| D                 | 18 (26%)    | 8 (24%)                         | 10 (29%)                      |
| **Prescription dose group, n (%)** |             |                                 |                               |
| 66 Gy             | 10 (15%)    | 6 (18%)                         | 4 (9%)                        |
| 70 Gy             | 35 (51%)    | 13 (38%)                        | 22 (65%)                      |
| 74 Gy             | 23 (34%)    | 15 (44%)                        | 8 (22%)                       |
| Rectal volume, mean (SD) | 77.1 (40.7) cm³ | 84.5 (48.0) cm³ | 69.7 (30.8) cm³ |
| **Risk group**    |             |                                 |                               |
| Intermediaterune | 47 (69%)    | 20 (59%)                        | 27 (71%)                      |
| High              | 21 (31%)    | 14 (41%)                        | 7 (21%)                       |
| BMI, mean (SD)    | 27.6 (3.4)  | 27.9 (3.4)                      | 27.4 (3.5)                    |
| PRS volume, mean (SD) | 140.7 (51.4) cm³ | 149.3 (49.0) cm³ | 132.2 (53.1) cm³ |
| Age at treatment, mean (SD) | 70.6 (6.1) | 71.4 (5.5)                      | 69.7 (6.6)                    |
| ≥G2 peak rectal bleeding, n (%) | 23 (34%) | 5 (15%)                         | 18 (55.9%)                    |

A: 6 months androgen suppression; B: 6 months androgen suppression + zoledronic acid; C: 18 months androgen suppression; D: 18 months androgen suppression + zoledronic acid; 

b 33 fractions, 2 Gy/fraction, PTV1 within 95% isodose.

c 74 Gy = 74 Gy+c.

d Phase 1 = 30 fractions, 2 Gy/fraction, PTV1 within 95% isodose. Phase 2 = additional dose in 2 Gy/fraction, PTV2 within 95% isodose.

| Definition of the PRS region |
|-------------------------------|

The PRS is here defined as the region of tissue, mostly fat, which the rectum can expand into or contract from [17]. Although the spatial extent of the fat region is relatively apparent on CT images, the extent of the fat which, when irradiated, could lead to rectal toxicity is ambiguous. As such, for this investigation, the entire region of fat adjacent to the prostate, rectum and bladder, though excluding any of the interior of those structures, was initially incorporated into defining the PRS. In order to optimise potentially causal dosimetric correlations, a sub-section of this region was then examined for statistical analysis as described below.

**Segmentation of PRS**

Due to the complexity and convoluted nature of the PRS region, manual segmentation on a large number of cases was considered infeasible. An auto-segmentation method was established based on manual delineation by a single observer (JD) on a series of ten test cases. A thorough description of the auto-segmentation process has been presented elsewhere [18].

In summary, on each analysed CT image set, the auto-segmentation involved defining a probability map for the PRS region based on non-rigid registration to the test cases. The voxels within the volume of interest were then labelled using an expectation maximisation clustering. The resulting structure was represented as a binary mask on each patient’s CT images. For the purpose of correlating PRS dosimetric factors with toxicity, the structure was refined by only including defined PRS image-pixels within 50 mm of the previously-delineated anorectum structure [8], as well as caudal to the bladder neck, and excluding any pixels within the outer wall of the rectum, as delineated at patient treatment planning. This ensured that the fat region immediately adjacent to the rectum was included in the dosimetric analysis excluding the fat region posterior to the bladder. It was also desired that the superior–inferior extent of the region should encompass the borders of the coplanar beams oriented about the cranio-caudal axis.

The auto-segmentation process was computationally intensive and, as such, a subset of 100 of the available RADAR patients (treated entirely with EBRT) was selected for auto-segmentation. This
was a case–control analysis representing patients with control-related toxicity (requiring LENT SOMA ‘stool frequency’ ≥ grade 2 and ‘urgency and tenesmus’ ≥ grade 2 at any time throughout a minimum 60 months follow-up) and without control-related toxicity (grade = 0 throughout a minimum 60 months follow-up). The anorectum and anal canal were also outlined separately for comparison. The anorectum was delineated as the outer rectal wall from the ischial tuberosities until the level where the rectum turns horizontally into the sigmoid colon, of which the anal canal was considered to be the caudal 3 cm [5].

Once segmented, cohorts with balanced characteristics were defined based on prescription dose, rectal and PRS volume and age at treatment.

Derivation of DVHs

Dose distributions from each treatment phase were combined on a voxel-by-voxel basis. DVH for the anorectum, anal canal and refined PRS regions were independently calculated as defined in Kennedy et al. [19]. Since the PRS is an undescribed structure in terms of radiobiology, physical dose was used for the study. The DVH data were imported, with toxicity data, to Matlab version 2013a (Mathworks, Natick, MA). The relationship between dosimetry and toxicity was explored using Atlas of Complication Incidence (ACI) [20,21] which were generated using a grid of 5 Gy dose and 10% volume bins. The denominator in each grid square indicates the number of patients whose DVH passed through whilst the numerator indicates the number of those patients who reported complications. ACI were generated to present the incidence of control-related toxicity for each of the outlined structures. For completeness ACI were also generated for rectal bleeding ≥ G2.

Statistical considerations

A spearman’s correlation matrix was generated to assess correlations between dosimetric descriptors of the 3 structures considered in this study PRS, anorectum and anal canal. Non parametric comparisons between the dosimetry of patients who did/did not report toxicity were made using Wilcoxon rank sum test. A Holm–Bonferroni correction was made to account for multiple testing of different dose levels. All statistical analysis was undertaken using R [22].

Results

Of the 100 patients chosen for auto-segmentation, 79 datasets were considered sufficiently well delineated for inclusion in the analysis. Fig. 1 present examples of the automatically defined PRS regions. 34 patients who did not report rectal control related toxicity had successful PRS delineation and these were balanced against 34 patients who reported rectal control-like toxicity. Table 1 details the patient characteristics of the 68 patients included in the dosimetric analysis. There were no significant differences between the groups with and without control-related toxicity in terms of prescription, rectal and PRS volume, BMI or age at treatment. The correlation matrix (Appendix Fig. A5) indicates a high degree of correlation between the dosimetric variables of a particular structure but low correlation between structures. The ACI relating the dose distribution to the PRS with rectal control-related toxicity is presented in Fig. 2. Fig. 3 shows the ACI for the subgroup of patients who reported control-related toxicity but who did not report rectal bleeding. The ACI relating the dose distribution to the PRS with rectal bleeding (≥G2) is shown in Fig. 4. Wilcoxon rank sum test results are presented in Table 2 where a number of dose levels were shown to be related to control-related toxicity for the PRS, when including all patients and also when excluding patients with rectal bleeding. However no results remained statistically significant after applying the Holm–Bonferroni correction. There were no statistically significant results when relating the PRS to Grade 2 rectal bleeding.

The ACI for the anorectum (Appendix Figs. A1 and A2) do not demonstrate a clear dosimetric relationship with either rectal bleeding or control-related toxicity. However, the ACI for the anal canal (Appendix Figs. A3 and A4) indicate a dose–response for both

![Fig. 1. Examples of autosegmented and processed PRS regions shown as a light-grey mask on axial, coronal and sagittal reconstructions.](image-url)
Fig. 2. Atlas of complication incidence (ACI) relating the perirectal space (PRS) with control-like rectal toxicity described using LENT/SOMA. The denominator in each box indicates the number of patients whose DVH passes through whilst the numerator details how many of those patients reported control-like rectal toxicity. Hot (red) regions of the colour scale indicate high incidence and cold (blue) regions indicate low incidence. The bottom right hand box indicates overall incidence in the cohort (shaded green). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

Fig. 3. Atlas of complication incidence (ACI) relating the perirectal space (PRS) with control-like rectal toxicity described using LENT/SOMA. Patients who reported rectal bleeding were excluded.

Fig. 4. Atlas of complication incidence (ACI) relating the perirectal space (PRS) with Grade 2 rectal bleeding (LENT/SOMA).
Results presented in Table 2 strongly support the hypothesis that the PRS behaves as a parallel-responding structure with significant dependence on the mean but not maximum dose. It must be noted that the dose distribution of the PRS is not highly correlated with the relationship with control-related toxicity. It has been shown that the dose distribution to the anal canal is related to control-related symptoms (personal communication 2017 Drs. Jervoise Andreyev and Andrew Wotherspoon).

To our knowledge, this is the first study of the dosimetric relationship between the PRS and control-related gastrointestinal toxicity. The dataset utilised was selected on the basis of availability, number of available participants, completeness and extent of follow-up. It must be acknowledged that the use of relatively dated treatment techniques, without image-guidance other than for bony anatomy, will likely influence applicability to contemporary treatment techniques, without image-guidance other than for bony anatomy, will likely influence applicability to contemporary treatments. The use of soft-tissue image-guidance and more conformal delivery techniques are known to impact on delivered dose distributions and toxicity incidence [26,27]. Validation in relevant datasets is desirable, including assessment of the mobility of the PRS on inter-fraction images. Further study of the individual structures within the PRS may provide more specific information relating dosimetry to toxicity.

**Conflict of interest statement**

The authors do not declare any conflict of interests relating to this manuscript submission.

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**Discussion**

Technological improvements enable radiotherapy delivery to be optimised to individual anatomy and function. This provides an opportunity to capitalise on an improved understanding of dose–response for discrete treatment complications. This study has focused on elucidating a more complete aetiology for a subset of gastrointestinal complications, utilising recent developments in non-linear image registration and autosegmentation.

The ACI and statistical analysis indicate that the strongest relationship between the outlined structures and control-related toxicity is described by the dose distribution to the PRS. Although the definition of the region is still ambiguous, it is hoped that development of voxel-level investigations as a means of refining the definition will develop a consensus of the structure delineation. Associated analysis, including assessment of inter-observer agreement, is underway.

The ACI relating anorectal DVH with rectal bleeding did not demonstrate a clear dose response. This study presents results on a small cohort reflecting the development efforts in autosegmentation of the PRS and case selection to specifically explore the relationship with control related toxicity. It has been shown that the dose distribution of the PRS is not highly correlated with that of the anorectum and anal canal. However, the results from a previous study utilising all available data from the RADAR study [8] do demonstrate a relationship between mid–high doses and rectal bleeding and between lower doses to the anal canal and urgency. Previous publications [11,23] have also indicated that the dose distribution to the anal canal is related to control-related symptoms. These results appear to be corroborated by the anal canal atlas presented in this study.

The ACI relating PRS DVH to control-like toxicities (Fig. 2) shows a clear pattern of increasing incidence rates with increasing dose and volume. This is apparent even when isolated just to patients without incidence of rectal bleeding (Fig. 3). The statistical results presented in Table 2 strongly support the hypothesis that the PRS behaves as a parallel-responding structure with significant dose–volume parameters across a broad range of doses, and a significant dependence on the mean but not maximum dose. It must be highlighted that duration of AD was significantly different between the patient groups (see Table 1). Although an attempt was made to match the groups based on their toxicity incidence using prescription dose, rectal volume and age at treatment, there were not sufficient patient numbers to allow control of all other factors. Previous analyses of the entire RADAR cohort have not uncovered significant impacts of AD duration or age [24,25]. Similarly note that no rectal filling protocol was specified for the trial and uniform proportions of any applied protocols between the toxicity groups cannot be guaranteed.

Given the role of the PRS in facilitating rectal motility, compliance and control, and the potential for fatty atrophy and fibrosis on irradiation, there is reason to hypothesise a causal relationship between dose to the PRS and subsequent control-related gastrointestinal symptoms. Moreover, a large number of sympathetic, parasympathetic and non-autonomic nerve fibres are to be found in the perirectal fat space. Radiation injury to the vasa nervorum may therefore directly lead to nerve dysfunction and contribute to control-related symptoms (personal communication 2017 Drs. Jervoise Andreyev and Andrew Wotherspoon).

**Table 2**

Wilcoxon rank sum test p values, relating individual dose metrics for the PRS, anorectum and anal canal to rectal control-like symptoms and bleeding. Results with an (uncorrected) p value < 0.05 shown in bold.

| Dose metric | Perirectal space DVH | Ano-rectum DVH | Anal canal DVH |
|-------------|----------------------|----------------|----------------|
|              | Other toxicity | Other toxicity | Other toxicity | Other toxicity | Other toxicity | Other toxicity |
|             | (RB < G2) | Rectal bleeding G2 | (RB < G2) | Rectal bleeding G2 | (RB < G2) | Rectal bleeding G2 |
| n = 68 | n = 44 | n = 68 | n = 68 | n = 44 | n = 68 | n = 68 |
| V5 | 0.163 | 0.620 | 0.088 | 0.154 | 0.421 | 0.454 | 0.123 | 0.168 | 0.328 |
| V10 | 0.031 | 0.293 | 0.184 | 0.421 | 0.413 | 0.974 | 0.085 | 0.110 | 0.280 |
| V15 | 0.025 | 0.168 | 0.571 | 0.611 | 0.620 | 0.568 | 0.173 | 0.143 | 0.433 |
| V20 | 0.029 | 0.070 | 0.964 | 0.677 | 0.544 | 0.107 | 0.195 | 0.276 | 0.332 |
| V25 | 0.034 | 0.063 | 0.894 | 0.579 | 0.314 | 0.050 | 0.071 | 0.235 | 0.411 |
| V30 | 0.014 | 0.030 | 0.854 | 0.312 | 0.103 | 0.073 | 0.048 | 0.200 | 0.302 |
| V35 | 0.037 | 0.013 | 0.448 | 0.296 | 0.114 | 0.157 | 0.066 | 0.302 | 0.326 |
| V40 | 0.015 | 0.005 | 0.362 | 0.083 | 0.047 | 0.264 | 0.057 | 0.338 | 0.130 |
| V45 | 0.072 | 0.146 | 0.904 | 0.424 | 0.677 | 0.864 | 0.185 | 0.922 | 0.028 |
| V50 | 0.107 | 0.192 | 0.954 | 0.308 | 0.732 | 0.964 | 0.164 | 0.748 | 0.013 |
| V55 | 0.089 | 0.175 | 0.954 | 0.258 | 0.788 | 0.814 | 0.164 | 0.902 | 0.013 |
| V60 | 0.089 | 0.209 | 0.814 | 0.206 | 0.864 | 0.411 | 0.112 | 0.795 | 0.005 |
| V65 | 0.189 | 0.291 | 0.774 | 0.118 | 0.677 | 0.181 | 0.085 | 0.406 | 0.011 |
| V70 | 0.931 | 0.535 | 0.748 | 0.602 | 0.553 | 0.255 | 0.740 | 0.806 | 0.316 |
| Mean | 0.025 | 0.067 | 0.995 | 0.134 | 0.248 | 0.794 | 0.085 | 0.418 | 0.071 |
| Max | 0.289 | 0.872 | 0.653 | 0.374 | 0.569 | 0.379 | 0.492 | 0.473 | 0.087 |
Fig. A1. Atlas of complication incidence (ACI) relating the anorectum with control-like rectal toxicity described using LENT/SOMA. The denominator in each box indicates the number of patients whose DVH passes through whilst the numerator details how many of those patients reported control-like rectal toxicity. Hot (red) regions of the colour scale indicate high incidence and cold (blue) regions indicate low incidence. The bottom right hand box indicates overall incidence in the cohort (shaded green). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

Fig. A2. Atlas of complication incidence (ACI) relating the anorectum with Grade 2 rectal bleeding described using LENT/SOMA.
**Fig. A3.** Atlas of complication incidence (ACI) relating the anal canal with control-like rectal toxicity described using LENT/SOMA.

**Fig. A4.** Atlas of complication incidence (ACI) relating the anal canal with Grade 2 rectal bleeding described using LENT/SOMA.
Fig. A5. Correlation matrix for dosimetric parameters of the PeriRectal Space (PRS), Anorectum (AR) and Anal Canal (AC).

References

[1] Michalski JM et al. Radiation dose–volume effects in radiation-induced rectal injury. Int J Radiat Oncol Biol Phys 2010;76(3 Suppl): S123–9.
[2] Thor M et al. Relationships between dose to the gastro-intestinal tract and patient-reported symptom domains after radiotherapy for localized prostate cancer. Acta Oncol 2015;54(9):1326–34.
[3] Krol R et al. Systematic review: anal and rectal changes after radiotherapy for prostate cancer. Int J Colorectal Dis 2014;29(3):273–83.
[4] Valdagni R, Rancati T. Reducing rectal injury during external beam radiotherapy for prostate cancer. Nat Rev Urol 2013;10(6):345–57.
[5] Peeters ST et al. Localized volume effects for late rectal and anal toxicity after radiotherapy for prostate cancer. Int J Radiat Oncol Biol Phys 2006;64 (4):1151–61.
[6] Fellin G et al. Long term rectal function after high-dose prostate cancer radiotherapy: results from a prospective cohort study. Radiother Oncol 2014;110(2):272–7.
[7] Cicchetti A et al. Modelling late stool frequency and rectal pain after radical radiotherapy in prostate cancer patients: results from a large pooled population. Phys Med 2016;32(12):1690–7.
[8] Ebert MA et al. Gastrointestinal dose–histogram effects in the context of dose-volume-constrained prostate radiation therapy: analysis of data from the RADAR prostate radiation therapy trial. Int J Radiat Oncol Biol Phys 2015;91 (3):595–603.
[9] Fiorino C et al. Late fecal incontinence after high-dose radiotherapy for prostate cancer: better prediction using longitudinal definitions. Int J Radiat Oncol Biol Phys 2012;83(1):38–45.
[10] Smeenk RJ et al. Dose–effect relationships for individual pelvic floor muscles and anorectal complaints after prostate radiotherapy. Int J Radiat Oncol Biol Phys 2012;83(2):636–44.
[11] Buettner F et al. The dose–response of the anal sphincter region – an analysis of data from the MRC RT01 trial. Radiother Oncol 2012;103(3):347–52.

[12] Lent soma scales for all anatomic sites. Int J Radiat Oncol Biol Phys 1995;31(5):1049–91.

[13] Hatton JA et al. Does the planning dose–volume histogram represent treatment doses in image-guided prostate radiation therapy? Assessment with cone-beam computerised tomography scans. Radiother Oncol 2011;98(2):162–8.

[14] Denham JW et al. Short-term androgen suppression and radiotherapy versus intermediate-term androgen suppression and radiotherapy, with or without zoledronic acid, in men with locally advanced prostate cancer (TROG 03.04 RADAR): an open-label, randomised, phase 3 factorial trial. Lancet Oncol 2014;15(10):1076–89.

[15] Boersma LJ et al. Estimation of the incidence of late bladder and rectum complications after high-dose (70–78 Gy) conformal radiotherapy for prostate cancer, using dose-volume histograms. Int J Radiat Oncol Biol Phys 1998;41(1):83–92.

[16] Ebert MA et al. Detailed review and analysis of complex radiotherapy clinical trial planning data: evaluation and initial experience with the SWAN software system. Radiother Oncol 2008;86:200–10.

[17] Chen N et al. Radiologic and anatomic study of the extraperitoneal space associated with the rectum. AJR Am J Roentgenol 2010;194(3):642–52.

[18] Gloso S, Denham JW, Ebert MA, Kennedy A, Mitra J, Dowling JA. Multi-atlas and unsupervised learning approach to perirectal space segmentation in CT images. Aust Phys Eng Sci Med 2016;39(4):933–41.

[19] Kennedy AM, Lane J, Ebert MA. An investigation of the impact of variations of DVH calculation algorithms on DVH dependant radiation therapy plan evaluation metrics. J Phys Conf Ser 2014;489(1):012053.

[20] Jackson A, Yorke ED, Rosenzweig KE. The atlas of complication incidence: a proposal for a new standard for reporting the results of radiotherapy protocols. Semin Radiat Oncol 2006;16(4):260–8.

[21] Otter S et al. Evaluation of the risk of grade 3 oral and pharyngeal dysphagia using atlas-based method and multivariate analyses of individual patient dose distributions. Int J Radiat Oncol Biol Phys 2015;93(3):507–15.

[22] R Development Core Team. R: a language and environment for statistical computing. Vienna, Austria: R Foundation for Statistical Computing; 2008.

[23] Heemskerken WD et al. 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;61(4):1011–8.

[24] Denham JW et al. Rectal and urinary dysfunction in the TROG 03.04 RADAR trial for locally advanced prostate cancer. Radiother Oncol 2012;105(2):184–92.

[25] Yahya N et al. Impact of treatment planning and delivery factors on gastrointestinal toxicity: an analysis of data from the RADAR prostate radiotherapy trial. Radiother Oncol 2014;9:282.

[26] Fonteyne V et al. Rectal toxicity after intensity modulated radiotherapy for prostate cancer: which rectal dose volume constraints should we use? Radiother Oncol 2014;113(3):393–403.

[27] Gill S et al. Acute toxicity in prostate cancer patients treated with and without image-guided radiotherapy. Radiat Oncol 2011;6:145.