Rationale and development of image-guided intensity-modulated radiotherapy post-prostatectomy: the present standard of care?

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Abstract: The indications for post-prostatectomy radiotherapy have evolved over the last decade, although the optimal timing, dose, and target volume remain to be well defined. The target volume is susceptible to anatomical variations with its borders interfacing with the rectum and bladder. Image-guided intensity-modulated radiotherapy has become the gold standard for radical prostate radiotherapy. Here we review the current evidence for image-guided techniques with intensity-modulated radiotherapy to the prostate bed and describe current strategies to reduce or account for interfraction and intrafraction motion.

Keywords: radiotherapy, prostate cancer, post-prostatectomy, image-guided radiation therapy

Introduction

Over the last three decades, external beam radiotherapy has evolved as a result of improvements in radiotherapy planning software, delivery, and introduction of computed tomography (CT)/magnetic resonance imaging (MRI)-based planning. Prostate radiotherapy has also evolved with these technological advancements, enabling dose escalation.¹⁻³ Comparative data suggest that intensity-modulated radiotherapy (IMRT) reduces particularly gastrointestinal (GI) toxicity compared with three-dimensional conformal radiotherapy (3D-CRT).⁴ IMRT allows not only sparing of organs at risk, but also irradiation of different tumor targets at various dose levels, known as simultaneous integrated boost techniques.⁵ Optimizing the conformality of radiation dose requires increased set-up precision. Therefore, methods to improve precision of planned dose delivery have been developed with image guidance, in addition to ways to stabilize the target volume.

Prostate cancer is the second most common cancer in men worldwide, and more than 1.1 million cases of prostate cancer were recorded in 2012. This accounts for 15% of the cancers diagnosed in men, with almost 70% of the cases occurring in the more developed regions.⁶

Radical prostate radiotherapy and radical prostatectomy (RP) are considered the mainstay of management for localized prostate cancer. The number of RPs performed worldwide is increasing.⁷⁻⁸ This increase is due to more patients being diagnosed with localized prostate cancer as a result of increased availability of prostate-specific antigen (PSA) testing,⁹ reduced morbidity associated with RP performed by “high volume” surgeons,¹⁰ and the developing role of RP as part of multi-modality treatment for patients with high-risk prostate cancer.¹¹⁻¹³
Recurrent disease occurs within 10 years in approximately one third of patients who have an RP.\textsuperscript{14–16} Recurrence risk is greatest among men with adverse pathological features such as positive surgical margins, seminal vesicle invasion, extraprostatic extension, and higher Gleason scores.\textsuperscript{17} There are established descriptions for adjuvant radiotherapy (ART) and salvage radiotherapy (SRT) post-prostatectomy. ART is given to patients with an undetectable PSA at high risk of recurrence because of adverse pathological features. Whereas SRT is given to patients with biochemical recurrence, defined as PSA $\geq 0.2$ ng/mL with a second confirmatory level of $>0.2$ ng/mL, but with no evidence of distant metastatic disease.\textsuperscript{18}

Three randomized trials have addressed the significance of ART, demonstrating a near 20\% absolute benefit for biochemical progression-free survival at 5 years after ART compared with a “wait and see” policy for patients with pT3± involved surgical margins.\textsuperscript{19–21} Systematic reviews\textsuperscript{22–25} have confirmed the benefit of ART. There are no randomized prospective studies available to prove the benefit of SRT for biochemical progression-free survival, local or systemic failure, or survival. However, observational studies using multivariate analyses have identified factors predictive of PSA recurrence and disease outcome.\textsuperscript{26–30}

The definitive answer as to whether immediate adjuvant treatment is superior to early SRT is currently unknown, but is being evaluated in prospective randomized trials. These trials include the international Medical Research Council RADICALS trial (NCT00541047)\textsuperscript{31} where patients can be randomized to either early postoperative radiotherapy or deferred radiotherapy given at the time of biochemical failure. This is defined as either two consecutive rising PSA levels and deferred radiotherapy given to patients with an undetectable PSA at high risk of recurrence because of adverse pathological features. Whereas SRT is given to patients with biochemical recurrence, defined as PSA $\geq 0.2$ ng/mL with a second confirmatory level of $>0.2$ ng/mL, but with no evidence of distant metastatic disease.\textsuperscript{18}

In the RADICALS trial (NCT00541047)\textsuperscript{32} led by the Trans-Tasman Radiation Oncology Group, patients with adverse prognostic factors were randomized to either receive ART or early SRT triggered by PSA rising to $>0.2$ ng/mL. A further trial, currently recruiting in France (NCT00667069), is randomizing patients to receive either immediate ART or delayed radiotherapy until biochemical relapse (defined as PSA $>0.2$ ng/mL but $\leq 2$ ng/mL), with both arms receiving 6 months of triptorelin, a luteinizing hormone releasing hormone analog.

Although the clinical indications for radiotherapy post prostatectomy are becoming established, there are many aspects of post-prostatectomy radiotherapy where evidence needs to be considered and consensus gained. In this review, we consider how to define the post-prostatectomy clinical target volume (CTV) and evidence to guide the selection of appropriate dose, radiotherapy planning, and delivery techniques.

**Post-prostatectomy clinical target volume**

To determine the optimal CTV, it is important to appreciate the most common sites of local relapse post-prostatectomy. Nearly two thirds of local relapses identified on imaging and/or biopsy occur at the vesicourethral anastomosis, with the bladder neck and retrotrigone area making up the significant remainder.\textsuperscript{33–35} A recent study using 1.5 or 3 Tesla (T) MRI scans from 113 patients with local recurrence assessed the locations of their recurrences to suggest an optimal target volume.\textsuperscript{36} The CTV proposal included 97\% of suspected tumor recurrences, which were found at the anastomotic site (78.8\%), bladder neck (15.3\%), and retrovesical area (5.9\%). In the cranial direction, 106 (87.3\%) lesions were located within 3 cm of the inferior border of the pubic symphysis, with 12 (10.2\%) lesions located below this anatomical point. In the transverse plane, 112 lesions (94.9\%) were located within 10 mm of the midline.

During the last decade, MRI has become commonly used in the planning of prostate radiotherapy, and compared with planning CT provides improved soft tissue resolution, allowing more consistent delineation of the prostate apex, anterior rectal wall,\textsuperscript{37} and penile bulb.\textsuperscript{38} These last two structures are also important when outlining the prostate bed. Additionally, identification of the vesicourethral anastomosis is better on T2-weighted MRI sagittal slices, where it is recognized as the disruption in the high signal of urine immediately below the urethral sphincter.\textsuperscript{39}

Choline positron emission tomography (PET) has been evaluated in patients with biochemical recurrence post-prostatectomy.\textsuperscript{40–42} In a study comparing multi-parametric MRI (mp-MRI) at 3 T with endorectal coil (EC) and $^{18}$F-choline PET/CT for detecting local recurrence after RP, 84 patients were allocated into two groups dependent on PSA and maximal transverse dimension of local recurrence.\textsuperscript{43} The superiority of mp-MRI was greater in group A (lesion size range 5–7.2 mm; PSA level range 0.8–1.4 ng/mL) than in group B (lesion size range 7.6–19.4 mm; PSA level range 1.3–2.5 ng/mL); the areas under the receiver operating characteristic curves for mp-MRI and PET/CT were 0.833 and 0.562 in group A and 0.971 and 0.837 in group B, respectively.

CTV definition in the postoperative setting is complicated due to changes in anatomy caused by surgery and the limited information on the preoperative location of the prostate.
The pelvic anatomy after robotic-assisted RP has been shown to be considerably different from that after open prostatectomy, with the size of the trigonal musculature defect being more pronounced, total urethral length statistically longer, and larger separation for the vesicorectal distance after robotic-assisted RP.44

Substantial interphysician variation in CTV delineation for post-prostatectomy radiotherapy exists, but this can be reduced by use of a contouring protocol.45 To date, four consensus articles have been published,39,46–48 and Table 1 summarizes their guidelines on CTV delineation. A comparison of these guidelines found that treatment volumes differed significantly between guidelines and that the European Organisation for Research and Treatment of Cancer (EORTC) volume was significantly smaller than the other guideline-produced target volumes.49

Use of multi-parametric MRI in CTV delineation

These consensus guidelines have been assessed in relation to the preoperative MRI contour of the prostate and gross visible tumor. CTVs according to the four consensus guidelines were applied and expanded to create a planning target volume (PTV).50 Irrespective of the guideline used, the consensus CTVs did not completely cover the pre-resection extent of the prostate seen on the MRI in any of the 20 patients evaluated. On average, 38% of the prostate volume and 41% of the gross tumor volume (GTV) on preoperative MRI were not included in the CTV. The Radiation Therapy Oncology Group (RTOG) and Princess Margaret Hospital guidelines are similar with respect to prostate and tumor coverage and yielded the best overall results. The EORTC guidelines provided the least overall coverage. The prostate base and mid zones were the predominant site of inadequate coverage.

A further appraisal by the same group compared a CTV based upon each patient's co-registered preoperative MRI and a CTV produced using RTOG guidelines, with respect to target volumes and doses to the rectum and bladder.51 The CTV produced using the preoperative MRI volume was a mean 18.6% larger than the CTV produced using the RTOG guidelines. The mean Jaccard Index, representing the intersection volume between CTVs, was 0.72 and 0.84 for PTVs, with both methods achieving similar rectal and bladder constraints as defined by the MRC RADICALS trial51 and QUANTEC criteria.52

In post-prostatectomy patients, mp-MRI has been shown to be an effective tool for evaluation of the prostatic fossa, with the addition of dynamic contrast-enhanced MRI improving the diagnostic performance in detecting local recurrence.53,54 However, these studies were conducted with a 1.5 T MRI scanner using an EC. MRI with EC causes a distortion of local anatomy and does not allow image fusion with the radiotherapy planning CT. A study assessing dynamic contrast-enhanced MRI scans acquired on a 1.5 T system without EC has shown that it can detect local recurrence with an estimated accuracy of 83% at low PSA levels (mean 0.74±0.64 ng/mL).

A pre-radiotherapy PSA cut-off value of ≥0.54 ng/mL predicted a positive result on dynamic contrast-enhanced MRI.55 This has been further supported by a recent trial using 3 T MRI without EC, which showed that the probability of radiographic local recurrence was significantly higher in patients with PSA >0.5 ng/mL.56 However, these are retrospective studies, and no histopathological confirmation was undertaken to confirm the interpretation of the mp-MRI findings. Additionally, to our knowledge, there are no comparative studies evaluating the value of increasing the field strength of the magnet and using an EC in detecting local recurrence. Barchetti and Panebianco have published a thorough appraisal of the data regarding mp-MRI in identification of local recurrence after RP.57 The further development of MRI and more widespread utilization of higher strength magnets may improve the precision of localization of recurrence and comprehensive studies are awaited with interest.

Boost to GTV

The finding of local relapse on mp-MRI enables a GTV to be defined and this has enabled a dose-adapted approach to treating the prostate bed with a boost to the GTV.58

The principle of a boost to the dominant nodule in radical prostate radiotherapy has been evaluated, and mp-MRI for SRT planning purposes has the potential to identify the suspected residual disease or locoregional recurrence. The ability to boost the suspected site of relapse has been evaluated in a retrospective study where patients received a boost of 10 Gy, with a median dose of 64 Gy to the prostatic bed. Boost and grade ≥2 acute genitourinary (GU) toxicity were independently correlated with late grade ≥2 GU toxicity on multivariate analysis. However, no significant difference in 3-year biochemical recurrence-free survival was observed between the boost and no-boost groups.59

Low dose rate brachytherapy in the salvage treatment of local recurrence after RP has been shown to be technically feasible, either as single modality treatment or to deliver a boost to the macroscopic disease within the prostate.
Table 1 Summary of the four consensus guidelines and guidance from the RADICALS trial for post-prostatectomy target delineation

<table>
<thead>
<tr>
<th>Guideline</th>
<th>Inferior</th>
<th>Superior</th>
<th>Lateral</th>
</tr>
</thead>
<tbody>
<tr>
<td>Princess Margaret Hospital Wiltshire et al⁹</td>
<td>8 mm below the VUA or the top of the PB, whichever is most superior</td>
<td>Superior surgical clips if present, or 5 mm above the inferior border of the vas deferens. Retained SV included when pathologically involved</td>
<td>Caudal: medial border of the levator ani and obturator internus. Cranial: Sacrorecto-genitopubic fascia</td>
</tr>
<tr>
<td>Australian and New Zealand Radiation Oncology Genito-Urology Group Sidhom et al⁸⁰</td>
<td>5–6 mm below the VUA, but should include all surgical clips inferiorly. If VUA not clearly defined, then slice above the PB</td>
<td>Encompass all of the SV bed as defined by non vascular clips and should include distal portion of the vas deferens. If SV pathologically involved, include any residual SV</td>
<td>Medial border of the levator ani muscle or obturator internus muscle</td>
</tr>
<tr>
<td>Radiation Therapy Oncology Group Michalski et al⁷⁷</td>
<td>8–12 mm below VUA, may include more if concern for apical margin. Can extend to slice above PB if VUA not well visualized</td>
<td>Level of cut end of vas deferens or 3–4 cm above top of symphysis. Include SV remnants if pathologically involved</td>
<td>Below superior edge of symphysis pubis: levator ani muscles, obturator internus Above superior edge of symphysis pubis: Sacrorecto-genitopubic fascia</td>
</tr>
<tr>
<td>EORTC (“identified areas of greatest risk of relapse”)*⁹⁰</td>
<td>Apex*: -15 mm cranially from the PB +5 mm in all directions***</td>
<td>Bladder neck*: +5 mm in all directions*** Original site of the base of SV should be included. If SV involved, include original position ± the remnants</td>
<td>Up to the neurovascular bundles (if removed up to the ilio-obturatic muscles) +5 mm in all directions***</td>
</tr>
<tr>
<td>Poortmans et al⁹⁰</td>
<td>5 mm cranial to the superior border of the PB</td>
<td>If SV low risk and pathology uninvolved: base of SV</td>
<td>Medial border of obturator internus and levator ani muscles</td>
</tr>
<tr>
<td>RADICALS guidance 2007</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Parker et al³¹</td>
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</table>

Notes: *If there is concern that extraprostatic disease at base may extend to the obturator internus; **supplementary 5 mm in the posterior and lateral directions in the presence of incompletely resected extracapsular nodal extension, but excluding the rectal wall; supplementary 5 mm in the direction of microscopically involved tumor margins as reported by the pathologist (except the rectal wall). ***Except the rectal wall.

Abbreviations: CTV, clinical target volume; EORTC, European Organisation for Research and Treatment of Cancer; PTV, planning target volume; PB, penile bulb; VUA, vesicourethral anastomosis; SV, seminal vesicles; SD, standard deviation.

High dose rate (HDR) brachytherapy with or without IMRT has also been evaluated as a treatment option for patients with local recurrence. Five patients were treated with IMRT to 45–50.4 Gy in 25–28 fractions to the prostate bed followed by two 9.5 Gy HDR brachytherapy fractions separated by 1–2 weeks, with the remaining patient treated with HDR brachytherapy (38 Gy in four fractions over 3 days). No patients had late grade 2 GI toxicity, with one patient developing late grade 2 urinary incontinence. It must be noted, however, that the median follow-up in this study was 9 months. Despite these studies do demonstrate the potential use of brachytherapy for biopsy, ultrasound, or MRI-proven local recurrence; however, they are single-center experiences. With improving diagnostic capabilities for local recurrence and MRI-ultrasound fusion brachytherapy techniques, comprehensive studies with brachytherapy in the salvage setting should be undertaken with long-term effectiveness, toxicity, and quality of life endpoints.

Radiotherapy dose and toxicity

Dose escalation

Among observational studies, radiotherapy doses ranged from 50 to 78 Gy, with SRT doses being somewhat higher than for ART. Although radiotherapy dose escalation improves freedom from biochemical recurrence (BCR) in radical prostate radiotherapy, the optimal post-prostatectomy dose has not yet been determined from a randomized trial. The predominant treatment failure site in patients post-prostatectomy is local. The American Society for Radiation Oncology and American Urological Association panel view is that 64–65 Gy is the minimum dose that should be delivered post-prostatectomy, but decisions regarding dose should be made by the treating physician.
### Summary of the four consensus guidelines and guidance from the RADiCALS trial for post-prostatectomy target delineation

<table>
<thead>
<tr>
<th>Anterior</th>
<th>Posterior</th>
<th>CTV** (cm³), mean ± SD</th>
<th>PTV** (cm³), mean ± SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Caudal: posterior edge of the symphysis pubis up to the top of the symphysis pubis</td>
<td>Caudal: anterior border of the rectal wall and levator ani</td>
<td>104±25</td>
<td>350±50</td>
</tr>
<tr>
<td>Cranial: posterior 1.5 cm of the bladder wall</td>
<td>Cranial: mesorectal fascia</td>
<td></td>
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<tr>
<td>Lower border of CTV to 3 cm superior, posterior aspect of the symphysis pubis</td>
<td>Levator ani and anterior rectal wall. More superiorly, anterior mesorectal fascia</td>
<td>88±16</td>
<td>325±32</td>
</tr>
<tr>
<td>More superiorly: posterior 1.5 cm of the bladder</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Below superior edge of symphysis pubis: posterior edge of pubic bone</td>
<td>Below superior edge of symphysis pubis: Anterior rectal wall</td>
<td>102±24</td>
<td>351±46</td>
</tr>
<tr>
<td>Above superior edge of symphysis pubis: posterior 1–2 cm of bladder wall</td>
<td>Above superior edge of symphysis pubis: Mesorectal fascia</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anastomosis and urethral axis +5 mm in all directions***</td>
<td>Up to but not including the outer rectal wall, cranially including the most posterior part of the bladder neck +5 mm in all directions***</td>
<td>60±17</td>
<td>254±53</td>
</tr>
<tr>
<td>Caudal (less than 2 cm above anastomosis): posterior aspect of symphysis pubis</td>
<td>Anterior rectal wall</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cranial (more than 2 cm above anastomosis): posterior one third of bladder wall</td>
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</tbody>
</table>

**Despite early ART at a standard dose of 60–64 Gy in 2 Gy per fraction, 20.9%–34.9% of patients showed BCR during follow-up.** In a retrospective cohort, around 75% of patients received a dose >66 Gy and BCR was seen at a median of 26.4 months in 19.7% of patients. Univariate analysis showed that T4 tumor stage, a preoperative PSA value >10 ng/mL, and a radiotherapy dose <70 Gy were significant factors for BCR. Cozzarini et al undertook a retrospective analysis where patients were grouped according to dose delivered to the prostate bed, ie, <70.2 Gy (median 66.6 Gy) or ≥70.2 Gy (median 70.2 Gy in 1.8 Gy fractions). Multivariate analysis confirmed that a dose ≥70.2 Gy was independently related to both BCR-free survival and disease-free survival, with similar results obtained after exclusion of patients receiving any androgen deprivation.

A further analysis evaluated the association between SRT dose and BCR, and their results suggested a dose response, with doses higher than 66.6 Gy resulting in decreased risk of BCR. Following SRT, reported 5-year biochemical control rates range from 25% to 70%. RETROGRADE analyses have identified a number of factors that may influence the efficacy of SRT, including dose. A systematic review by King included 41 published SRT studies, with a median dose of 64.6±3.1 Gy (range 60–74.8 Gy). There was a significant association between dose and relapse-free survival (rho =0.42, P=0.0052), with an observed improvement in relapse-free survival of 2% for each incremental Gy of dose.

A systematic review and regression meta-analysis from 25 studies including 3,828 patients, generated tumor control probability and normal tissue complication probability models. They estimated that with a pre-SRT PSA level of 0.4 ng/mL, an approximately 50% chance of 5-year biochemical control probability.
progression-free survival could be achieved with an SRT dose of 60 Gy. However, if the PSA level before SRT was 1.0 ng/mL, a dose of approximately 70 Gy was required to achieve similar disease control, with severe late toxicity rates at that dose potentially reaching 10%. There are many confounding factors, including the use of hormonal therapy and a variety of treatment planning and delivery techniques, that preclude high quality evidence level guidance to be drawn.

A randomized Phase III trial (SAKK 09/10) for SRT is currently recruiting, where patients are randomized to either receive 64 Gy or 70 Gy.71 The target volume delineation is to be performed according to EORTC guidelines and patients are excluded if they have macroscopic local recurrence.

Hypofractionation
Radiobiological studies have suggested that the estimated $\alpha/\beta$ ratio of prostate carcinoma is lower than for most other solid tumors, and around 1.5–3 Gy.74–77 Hypofractionated regimens may therefore give tumor control advantages over the more traditional 2 Gy per fraction schedules. This has been investigated internationally in several large Phase III trials in radical prostate radiotherapy.78–80 A few studies have considered this in post-prostatectomy radiotherapy,81–86 and Table 2 summarizes these data. However, most of these studies have only reported on acute toxicity, and hypofractionation does pose an increased risk for late toxicity as the fraction size and total dose increases. A recent retrospective analysis to assess predictors of severe (grade $\geq 3$) late urinary toxicities after post-prostatectomy radiotherapy with conventional (1.8 Gy per fraction) and hypofractionated (median 2.5 Gy per fraction) radiotherapy found on multivariate analyses, acute grade $\geq 2$ toxicity and hypofractionation to independently predict late grade $\geq 3$ toxicity. This was in both the adjuvant and salvage settings, with different radiotherapy techniques and non-standardized reporting of toxicity.87

**Table 2** Comparison of acute toxicity for hypofractionated and conventionally fractionated post-prostatectomy radiotherapy

<table>
<thead>
<tr>
<th>Reference</th>
<th>Patients (n)</th>
<th>Trial design</th>
<th>Total dose/single dose fractionation (EQD2)</th>
<th>Treatment technique</th>
<th>Acute GI toxicity (%)</th>
<th>Acute GU toxicity (%)</th>
<th>Scoring system</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Conventionally fractionated post-prostatectomy radiotherapy</strong></td>
<td></td>
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<tr>
<td>De Meerleer et al $^7$</td>
<td>135</td>
<td>Retrospective</td>
<td>Median 75 Gy/2 Gy</td>
<td>IMRT, regular iGRT</td>
<td>G2: 15</td>
<td>G2: 28</td>
<td>In-house</td>
</tr>
<tr>
<td>Cozzarini et al $^9$</td>
<td>153</td>
<td>Retrospective</td>
<td>Median 66.6 Gy/1.8 Gy</td>
<td>Conventional non-conformal, 3D-CRT</td>
<td>G2/3: 17.5</td>
<td>G2: 10.5</td>
<td>RTOG</td>
</tr>
<tr>
<td></td>
<td>181</td>
<td></td>
<td>Median 70.2 Gy/1.8 Gy</td>
<td></td>
<td>G2/3: 14</td>
<td>G3: 4</td>
<td></td>
</tr>
<tr>
<td>Nath et al $^9$</td>
<td>50</td>
<td>Retrospective</td>
<td>Median 68 Gy/1.8–2 Gy</td>
<td>IMRT, daily iGRT</td>
<td>G2: 8</td>
<td>G2: 14</td>
<td>NCI CTCAE</td>
</tr>
<tr>
<td>Riou et al $^0$</td>
<td>57</td>
<td>Retrospective</td>
<td>Mean 68 Gy/2 Gy</td>
<td>IMRT, iGRT</td>
<td>G2: 4</td>
<td>G2: 7</td>
<td>NCI CTCAE</td>
</tr>
<tr>
<td>Bellavita et al $^4$</td>
<td>182</td>
<td>Retrospective</td>
<td>Median 66.6 Gy/1.8–2 Gy</td>
<td>3D-CRT</td>
<td>G2: 39</td>
<td>G2: 21</td>
<td>RTOG</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>G3: 1</td>
<td>G3: 0</td>
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<tr>
<td><strong>Hypofractionated post-prostatectomy radiotherapy</strong></td>
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<td></td>
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<tr>
<td>Massacesi et al $^4$</td>
<td>49</td>
<td>Prospective, non-randomized Phase II</td>
<td>62.5 Gy/2.5 Gy (71.4 Gy)</td>
<td>SIB-IMRT</td>
<td>G2: 32.6</td>
<td>G2: 9.6</td>
<td>RTOG</td>
</tr>
<tr>
<td>Cozzarini et al $^4$</td>
<td>50</td>
<td>Prospective, non-randomized Phase III</td>
<td>58 Gy/2.9 Gy (72.9 Gy)</td>
<td>Tomotherapy, daily iGRT</td>
<td>G2: 4</td>
<td>G2: 10</td>
<td>RTOG</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>G3: 0</td>
<td>G3: 2</td>
<td></td>
</tr>
<tr>
<td>Kruser et al $^4$</td>
<td>108</td>
<td>Retrospective</td>
<td>65 Gy/2.5 Gy (74.3 Gy)</td>
<td>Tomotherapy, daily iGRT, endorectal balloon</td>
<td>G2: 14</td>
<td>G2: 7</td>
<td>Modified RTOG</td>
</tr>
<tr>
<td>Katayama et al $^4$</td>
<td>39</td>
<td>Prospective, non-randomized Phase II</td>
<td>54 Gy/3 Gy (69.4 Gy)</td>
<td>Tomotherapy, daily iGRT</td>
<td>G3: 0</td>
<td>G3: 1</td>
<td>RTOG</td>
</tr>
<tr>
<td>Gladwish et al $^4$</td>
<td>30</td>
<td>Prospective, non-randomized Phase III</td>
<td>51 Gy/3 Gy (65.6 Gy)</td>
<td>IMRT, daily iGRT, fiducial-based</td>
<td>G2: 0</td>
<td>G2: 3</td>
<td>NCI CTCAE</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>G3: 3</td>
<td>v. 3.0</td>
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</tr>
</tbody>
</table>

Note: EQD2, 2 Gy-equivalent dose (assumed $\alpha/\beta$ ratio of 1.5 Gy).

**Abbreviations:** GI, gastrointestinal; GU, genitourinary; iGRT, image-guided radiotherapy; IMRT, intensity modulated radiotherapy; NCI CTCAE, National Cancer Institute Common Terminology Criteria for Adverse Events; RTOG, Radiation Therapy Oncology Group; 3D-CRT, three-dimensional conformal radiotherapy; SIB-IMRT, simultaneous integrated boost intensity modulated radiotherapy; G, grade.
urinary toxicity compared with 3D-CRT in patients treated at the same dose and to the same volume definition.⁹⁸

A few retrospective studies have investigated the use of IMRT for post-prostatectomy radiotherapy.⁷⁷,⁹⁵,⁹⁰ Bastash et al⁹¹ reported the effects on erectile function of dose-escalated prostate bed IMRT in a retrospective series. Despite the high dose (mean 69.6 Gy) to the prostate bed, this had no negative effect on erectile function for the patients who remained potent after nerve-sparing prostatectomy; the follow-up in this group was 19.5 months.

Ost et al⁹² reported the clinical results of adjuvant IMRT (median dose to PTV: 74 Gy in 2 Gy per fraction) in 104 patients. With respect to acute and late toxicity, no patients developed grade 3 GI toxicity, with 8% and 4% of patients developing acute and late grade 3 GU toxicity respectively.

Toxicity with dose escalation in SRT has been retrospectively evaluated, and with a median dose of 70.2 Gy, grade 3/4 late GU and GI toxicity was 29.6% and 5.6%, respectively. There was a non-significant trend towards reduced late GI toxicity with IMRT.⁹³

IMRT creates an opportunity to reduce side effects or attempt dose escalation to increase the probability of tumor control. A Phase II trial has recently been published, where patients who received 66 Gy in 2 Gy dose per fraction with IMRT were evaluated with health-related quality of life outcomes.⁹⁴ Expanded Prostate Cancer Index Composite (EPIC) scores were collected prospectively and showed transient declines in the EPIC GI domain summary score and GU irritative subscale, with complete recovery occurring between 3 and 12 months after radiotherapy, and remaining stable compared with baseline at 5-year follow-up. Sexual health-related quality of life remained stable at 5 years, with an improving trend in bother subscale.

At the Memorial Sloan-Kettering Cancer Center, a retrospective analysis of late toxicity (5-year actuarial rates) was undertaken of patients predominantly treated with either 3D-CRT or IMRT to a dose of ≥70 Gy with a median follow-up of 60 months.⁹⁵ Two hundred and five patients (72%) were treated with doses ≥70 Gy. IMRT was independently associated with a reduction in late grade ≥2 GI toxicity compared with 3D-CRT (1.9% versus 10.2%, respectively; ⁹⁷ P=0.02), despite the fact that those patients treated with IMRT were more likely to be treated with a higher dose than those receiving 3D-CRT. This supports the idea that IMRT improves the therapeutic ratio associated with SRT. However, IMRT was not associated with a reduction in risk of grade ≥2 GU toxicity, urinary incontinence, or grade 3 erectile dysfunction.

**Radiotherapy planning technique**

The CTV has an irregular shape, with its borders associated with the rectum and bladder, making it particularly difficult to sculpt the dose away from the organs at risk (OAR). Koontz et al⁹⁶ demonstrated that IMRT provides better high-dose sparing of the OAR than 3D-CRT. In addition, Cozzarini et al⁹⁷ demonstrated a benefit for helical tomotherapy compared with 3D-CRT with regards to rectal sparing.

As previously discussed, there is no clear consensus regarding CTV delineation for postoperative prostate radiotherapy. Analysis of dose volume histograms (DVH) was performed with the four published guidelines for CTV delineation⁹⁸-⁹⁹ and the dose constraints proposed in QUANTEC³¹ and the RADICALS trial. Comparison between 3D-CRT and tomotherapy IMRT showed that the latter reduced OAR irradiation; however, despite using IMRT, a significant percentage of cases did not meet the OAR dose constraints.⁹⁶ There is currently no agreement regarding the superiority of advanced forms of external beam radiotherapy.

A radiotherapy quality assurance program undertaken as part of the SAKK 09/10 trial had 43% of the centers using 3D-CRT, with the remaining centers using IMRT or volumetric modulated arc therapy (VMAT). CTVs were outlined using the EORTC guidelines, and the rectal and bladder wall DVH parameters with IMRT or VMAT versus 3D-CRT plans were not significantly different.⁹¹ A study quantifying the differences in treatment delivery efficiency and dosimetry between step and shoot IMRT, VMAT, and helical tomotherapy has shown that VMAT improves the efficiency of delivery for equivalent dosimetric quality compared with IMRT and helical tomotherapy in prostate bed ± whole pelvis radiotherapy.⁹⁷

**Image-guided radiotherapy**

Variation in location of the prostate bed is significantly influenced by the changing shape and volume of the rectum and bladder during treatment. Rectal volumes can vary significantly throughout treatment from −40% to +60% compared with planning, with bladder volumes fluctuating up to 200 cm³.³⁹ These day-to-day fluctuations can have a substantial dosimetric effect on both the prostate bed and OAR.⁹⁸ The greatest potential for geographical miss has been seen when either the bladder increases in size or the rectum becomes smaller.¹⁰⁰ Uncertainties due to patient set-up errors and prostate bed motion¹⁰¹ require a margin around the CTV to create the PTV. Minimizing these uncertainties can facilitate smaller CTV to PTV margins, thereby reducing the dose to OAR.
The accuracy of bony anatomy as a surrogate for prostate bed position is inadequate to account for intrafraction motion. Relative to bony anatomy, prostate bed displacement exceeded 5 mm in 21% of treatments of 20 patients in the superior–inferior direction and 9% in the anterior–posterior position. During treatment, the target exceeded the 5 mm tracking limit for at least 30 seconds in 11% of all fractions, generally in the anterior–posterior or superior–inferior direction. In the anterior–posterior direction, target motion was more likely to move posteriorly towards the rectum than anteriorly.\(^{102}\)

A predominant source of intrafraction error is prostate bed motion, and there is an interest in determining the best treatment margin to account for daily set-up uncertainty and prostate bed motion. There is a lack of data regarding the best treatment margin to account for daily set-up uncertainty and organ motion. Surgical clips have been used as fiducial markers for the prostate bed, with their positional shifts captured and recorded by kV images.\(^{103,104}\) However, surgical clips have often been found to be positioned outside the prostate bed, poorly visualized, and closely clustered.\(^{105}\)

Implantation of gold seeds\(^{106}\) and electromagnetic transponders\(^{107}\) into the prostate bed has been described and data regarding prostate bed motion published.\(^{102,108,109}\) The stability and interobserver variability of fiducial markers and position of surgical clips in the prostate bed has been considered, and fiducial markers were seen to give less interobserver variability in matching and less variation in position than surgical clips.\(^{105}\) The largest discrepancy in matching between surgical clips and fiducial markers was in the anterior–posterior axis. Migration of gold seed fiducials over a course of radiotherapy has been shown to be minimal, with the mean measured differences in inter-marker distance between the start and end of radiotherapy being 0.4 mm.\(^{110}\) However, implantation of the fiducial markers is an invasive procedure with associated morbidity.\(^{106}\)

Table 3 summarizes the findings from studies evaluating interfraction prostate bed motion, which is defined as the change in position of the prostate bed in relation to pelvic bony anatomy. There has been a suggestion from the data that an anisotropic PTV margin should be used in post-prostatectomy radiotherapy.\(^{111}\) A differential superior and inferior prostate bed motion has been reported, with the greatest movement occurring in the anterior–posterior plane in the upper prostate bed; however, in this study, only one surgical clip in the superior and in the inferior section was selected. The variability in movement between these clips implies that prostate bed tilt is a factor in interfraction motion and may not be easily corrected for with standard online matching techniques.

Significant intrafraction prostate bed motion and deformation has been found, as the boundaries are largely defined by the interfaces of the bladder and rectum, which are influenced by intrafractional variations in filling. Diot et al\(^{112}\) performed a study analyzing the effects of anatomical interventions, such as adjusting bladder filling, evacuation of stool, or insertion of a rubber catheter into the rectum to deal with excess gas. The indications for these interventions were determined on sagittal images when the anterior rectal wall displacement was greater than 5 mm or by a change in volume of the bladder by a factor of 2 compared with the planning CT. The comparison of pre and post intervention localization suggests that fluctuations in the volume and placement of the bony anatomy surrounding the prostate bed.

<table>
<thead>
<tr>
<th>Reference</th>
<th>Patients/images</th>
<th>Interfraction PBM</th>
<th>Lateral mm (SD)</th>
<th>Superior-inferior mm (SD)</th>
<th>Anterior–posterior mm (SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Schiffner et al(^{10})</td>
<td>Gold seeds, EPID</td>
<td>Mean:</td>
<td>0.3 (0.9)</td>
<td>0.4 (2.4)</td>
<td>−1.1 (2.1)</td>
</tr>
<tr>
<td>Sandhu et al(^{14})</td>
<td>Surgical clips, kV</td>
<td>Mean magnitude:</td>
<td>1 (1.7)</td>
<td>2.4 (2.1)</td>
<td>2.7 (2.1)</td>
</tr>
<tr>
<td>Ost et al(^{15})</td>
<td>Anterior rectal wall, CBCT</td>
<td>Mean:</td>
<td>0.01</td>
<td>0.58</td>
<td>2.19</td>
</tr>
<tr>
<td>Huang et al(^{10})</td>
<td>Surgical clips, CBCT</td>
<td>Calculated margin*</td>
<td>1.78</td>
<td>3.27</td>
<td>7.88</td>
</tr>
<tr>
<td>Bell et al(^{11})</td>
<td>Surgical clips, CBCT</td>
<td>Calculated margin*</td>
<td>3.24</td>
<td>5.49</td>
<td>8.36</td>
</tr>
<tr>
<td>Alander et al(^{19})</td>
<td>Gold seeds, CBCT</td>
<td>Mean:</td>
<td>Upper 0.1 (0.12)</td>
<td>0.28 (0.26)</td>
<td>0.5 (0.5)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Lower 0.08 (0.1)</td>
<td>0.18 (0.17)</td>
<td>0.18 (0.16)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Calculated margin**</td>
<td>1.4</td>
<td>3.9</td>
</tr>
</tbody>
</table>

Notes: PBM, motion of the either the gold seeds, surgical clips, or anterior rectal wall in relation to bony pelvic anatomy and is the mean of the individual patient means, unless otherwise stated. Mean magnitude: average of absolute values of all measurements in a given plane. Margin recipe used: \(+2.5\Sigma + 0.7\pi\); **1.96\Sigma + 0.7\pi.**

Abbreviations: PBM, prostate bed motion; CBCT, cone-beam computed tomography; CTV, clinical target volume; PTV, planning target volume; IGRT, image-guided radiotherapy; SD, standard deviation.
shape of the rectum and bladder increase the variability of the localization data most significantly in the anterior–posterior direction, with lesser effect in the superior–inferior direction. The dosimetric impact on performing these interventions to reduce treatment volume deformations due to bladder and rectal filling showed no significant difference for PTV coverage, or rectal or bladder sparing. However, this concept should be re-examined for hypofractionated treatments, as the gains from correcting one fraction would contribute to a larger portion of the treatment, and therefore may be more dosimetrically relevant. Other studies have used the anterior rectal wall as a surrogate for prostate bed motion.

Post-prostatectomy IMRT has enabled improved toxicity profiles compared with conventional radiotherapy techniques. With the increasing use of IMRT, enabling dose escalation and the associated steep dose gradients, it is critical to accurately localize the target for precise treatment delivery. Image-guided radiotherapy (IGRT) either using soft tissue, surgical clips, fiducial markers have been applied in prostate bed radiotherapy. The associated consequences in toxicity with IGRT have been evaluated in retrospective studies using 3D-CRT or IMRT.

Studies using cine-MRI to assess intrafraction motion identified rectal filling as a predictor of prostate motion. A filled rectum is associated with mobile gas pockets, leading to rectal motion. Therefore, a device inserted into the rectum should minimize a change in rectal filling or gas position and hence stability of rectum and therefore prostate or prostate bed.

The use of endorectal balloons (ERBs) was first reported in prostate radiotherapy in 1979, and has been investigated regarding its potential immobilizing properties and dosimetric consequences. Definite conclusions from a systematic review are difficult as there were many variables between the trials; however, it was felt that as the ERB is situated directly adjacent to the anterior rectal wall and is visualized on portal imaging, it can assist in localizing the prostate and thus in reducing the CTV to PTV margins. There was a rectal and anal wall sparing effect for the intermediate and high-dose regions even with IMRT, and the dosimetric consequence on the target volume of having an air-filled balloon in the rectum has been addressed and shown not to Underdose the prostate.

In the last 5 years, reports of ERBs in the post-prostatectomy setting have been published. Improvement in DVH with ERB has been seen in a planning study by Smeenk et al. They found significantly reduced anal wall DVH and to a lesser extent rectal DVH with the ERB. The mean dose to the anal wall was reduced by 6 Gy. However, a study assessing dosimetric stability to the CTV with ERBs did not observe any improvement, although there was improved geometric stability of the rectum. This observation may be due to deformation of the CTV caused by the ERB.

A recently published study by the same group compared geometric variations in CTV and OAR during prostate bed radiotherapy with and without the use of ERB. Cone-beam CTs were reviewed and CTV contoured and subdivided into superior and inferior CTV with the whole rectal volume subdivided into superior and inferior rectum and anal volume. The concordance index of cone beam CT treatment volumes compared with planning volumes was calculated and displacements measured. Rectal stability was improved with the use of ERB (concordance index improvement from 0.41 to 0.71), which translated into greater CTV stability with the improved concordance index in the ERB group. However, displacements of the center of volumes (centroids) for the superior and inferior CTV were not significantly different between the two groups. This study also looked at bladder filling and found that the ERB negated the impact of bladder filling on CTV stability, postulating that this was due to the ERB physically compressing the bladder anteriorly against the pubic symphysis for the majority of the CTV. However, they did not report on centroid displacements in the superior–inferior direction, which may be more susceptible due to positioning interfraction variation with the ERB.

A retrospective analysis of the largest reported cohort using ERBs after RP was by Ishiyama et al. They reported acceptable late RTOG GI and GU toxicity, with the highest late GU toxicity being grade 2 in 13% and grade 3 in 6% of patients. The highest late GI toxicity was grade 2 in 6% and grade 3 in 3% of patients. One hundred and seven patients were assessed, and the prescribed mean dose to the CTV was 70 Gy in 32 fractions (EQD2 = 73.9 Gy based on α/β = 1.5). However, this study has not recorded patient-reported outcomes, which may be a better measure of treatment-related toxicities than physician-reported assessments.

A novel rectal obturator, ProSpare™, has been developed at the Institute of Cancer Research, London, UK, as a single-use device made from high impact polystyrene, which is inserted by the patient just prior to radiotherapy. It has radiopaque markers encased in the anterior and posterior wall of
the device, allowing clear identification of the anterior rectal wall. It has venting holes in the tip of the device and a venting line along the central join of the device to allow passage of rectal gas through the device and deflate any rectal gas bubbles on insertion. It has been evaluated in prostate radiotherapy and shown to be an effective minimally invasive daily online image-guided tool and rectal spacer. 130–132 A Phase II trial (POPS; Post-Operative ProSpare) randomizing patients to receive post-prostatectomy IMRT with or without ProSpare will start recruiting in the UK by the end of 2015.

Conclusion

Despite the publication of four guidelines, there is a lack of consensus on CTV definition. Information from imaging such as mp-MRI should be considered when defining the CTV and incorporated into the contouring guidelines, to enable a personalized approach to an optimal target volume. It is unclear whether a differential dose in ART and SRT is needed to improve biochemical failure-free survival and whether this translates into an overall survival benefit. The long-term efficacy and tolerability data for hypofractionated schedules are not yet established.

IMRT and IGRT enable dose escalation to the target volume and permit delivery of a simultaneous integrated boost to a GTV; randomized trials are needed to determine if there is a clinical outcome benefit with acceptable toxicity. More data in larger patient groups on IGRT post-prostatectomy and prostate bed motion are needed before CTV-to-PTV margins can be modified. Rectal stabilizing methods are available and should be considered, as this may result in a reduction in prostate bed deformation and motion during treatment.

Toxicity with dose escalation and IMRT techniques appears acceptable, although GU toxicity remains the dose-limiting factor. Increased knowledge of the dose-response relationship is needed regarding OAR or specific regions within these organs associated with GU toxicity.

As the current randomized controlled trials mature, we shall have better information to determine the indications for radiotherapy and its timing and the need for androgen deprivation therapy. However, challenges remain to providing the evidence base to refine the definition of radiotherapy planning targets, optimal treatment planning, and delivery strategies.

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Disclosure

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References


