review

Ulthahypofractionation of localized prostate cancer – Statement from the DEGRO working group prostate cancer

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Abstract

Due to its low fractionation sensitivity also known as ‘alpha/beta ratio’ in relation to its surrounding organs at risk, prostate cancer is predestined for hypofractionated radiation schedules assuming an increased therapeutic ratio compared to normofractionated regimen. While moderate hypofractionation (2.2-4Gy) has been proven to be non-inferior to normal fractionation in several large randomized trials for localized prostate cancer, level I evidence for ultrahypofractionation (>4Gy) was lacking until recently. An accumulating body of non-randomized evidence has recently been strengthened by the publication of two randomized studies comparing ultrahypofractionation with a normofractionated schedule, the Scandinavian HYPO-RT trial by Widmark et al. and the first toxicity results of the PACE-B trial.

In this review, we aimed to give a brief overview of the current evidence of ultrahypofractionation, make an overall assessment of the level of evidence and provide recommendations and requirements that should be followed before introducing ultrahypofractionation in routine clinical use.
Introduction

External beam radiation (EBRT) is one of the mainstays in the treatment of prostate cancer of all risk groups to all patients who are in the decision making process of which treatment to choose. This decision has become more complex recently since many – equally effective – treatment alternatives are available, including active surveillance or deferred treatment for low risk (LR) disease. The fact that LR prostate cancer bears a high risk of overtreatment is now unanimously addressed in current relevant guidelines and translates in de-escalated treatment regimens where potential side effects are very carefully weighed against the benefits of a given therapy. In contrast, high-risk (HR) prostate cancer still represents a potentially lethal disease demanding more aggressive treatment.

In that light, ultrahypofractionation qualifies as a viable option in the primary treatment of localized prostate cancer, since it can be tailored to the risk status in terms of fractional and total dose, with or without androgen deprivation therapy. In a situation where LINAC capacities are limited in many countries (or reduced as a side effect of the current COVID-19 epidemic) possibilities to reduce treatment time or fractions without compromising outcome are highly sought after. At the same time ultrahypofractionation offers a high level of patient convenience due to low overall treatment time without an excess of toxicity. Thus, it is viewed as an attractive alternative to surgery.

In this review we recapitulate the more recent literature (randomized evidence and metaanalyses) on ultrahypofractionation, put it in context with current recommendations and provide principles which should be followed before introducing ultrahypofractionation into clinical routine.

Terminology

*Extreme* or *ultra-*hypofractionation is commonly used synonymously with stereotactic body radiation therapy (SBRT) and stereotactic ablative body radiation (SABR), although the former terms strictly refer to the fraction size whereas the latter also refer to the platform of beam delivery and radiation technique. We therefore chose to use the term *ultrahypofractionation* for all forms of delivery of more than 4Gy per fraction.

The American Society for Radiation Oncology (ASTRO), American Society of Clinical Oncology (ASCO) and American Urological Association (AUA) hypofractionation guideline
[1] defines moderate hypofractionation as 2.4–3.4 Gy/fraction and ultrahypofractionated radiotherapy as doses per treatment of 5.0Gy/fraction or higher, thus leaving a ‘grey zone’ between 3.4 and 5Gy.

The Prostate Cancer Expert Panel of the German Society of Radiation Oncology (DEGRO) and the Working Party Radiation Oncology of the German Cancer Society (DKG-ARO) use a definition of 2.2-4Gy/fraction for moderate and beyond 4Gy/fraction for ultrahypofractionation [2].

Ultrahypofractionation is usually delivered using high precision techniques (LINAC based or CyberKnife) aided by daily image guidance including adequate motion management strategies allowing for small PTV margins and high dose conformation.

Radiobiology
While for most cancer types a normofractionation schedule of 1.8-2Gy per day / five times a week represents the sweet spot in terms of tumor control and toxicity, some tumors exhibit a higher sensitivity to fraction doses and might therefore benefit from hypofractionated schedules. This property is reflected by a low alpha/beta (α/β) value and can be quite accurately described with the so-called linearquadratic model [3]. The α/β value is a measure of fractionation sensitivity and is related to the inherent capacity of tumor cells to repair sublethal DNA damage inflicted by ionizing radiation.

Whether hypofractionation is beneficial depends on the relation of α/β values of the target in relation to its surrounding normal tissues. For prostate cancer cells, very low α/β values of about 1.5Gy have been derived from multiple preclinical and clinical studies [4-9]. Late toxicity of the bladder and rectum has been estimated to have an α/β value of 5.6Gy [10,11] and 3Gy [12,3], respectively. Therefore, in theory, hypofractionated radiation schedules should have a beneficial effect on the therapeutic ratio.

More recent data have shown that, in addition to fraction dose, overall treatment time seems to play a major role [13] which has been neglected in the aforementioned calculations of the α/β ratio ([14]reviewed in [15]). When a time factor is accounted for in the calculation, slightly higher α/β values will result so that many authors nowadays endorse values of approximately 2-2.7Gy. In a recent meta-analysis, Vogelius and Bentzen calculated α/β values based on 13 randomized trials with and without the presence of a time-factor of 0.31Gy loss.
per day, yielding $\alpha/\beta$ values of 1.2Gy and 2.7Gy, respectively. Of note, the higher $\alpha/\beta$ derived from hypofractionated dose escalation studies might in part be contributed to the fact, that the dose response relationship starts to max out at approximately EQD2 80Gy – a dose which is superseded by most ultrahypofractionation regimen [16]. It also needs to be emphasized that when comparing EQD2’s of different fractionation regimen using the time corrected $\alpha/\beta$ value, only regimen with the same overall treatment time should be compared. For that reason, we chose to use an $\alpha/\beta$ value of 2Gy in the present manuscript in order to appreciate that ultrahypofractionation regimen have a considerably reduced overall treatment time (mostly roughly 2 weeks) compared to normofractionated as well as moderately hypofractionated regimen.

Cost effectiveness
In addition to its potential benefit in regard of the therapeutic ratio, ultrahypofractionation may reduce treatment cost for prostate cancer, which, due to its high prevalence, has a major impact on general health care expenses.

It has been shown that ultrahypofractionation is associated with less overall treatment cost than normofractionated 3D conformal or IMRT [17] as well as moderate hypofractionation [18]. In a recent systemic review comprising 12 studies Abreha et al. [19] performed a model-based cost-effectiveness analysis confirming that ultrahypofractionation is the most effective treatment in terms of overall treatment cost, including prostatectomy.

However, most available studies are based on the US Medicare system. For Europe, treatment costs can differ dramatically but there is reason to assume that the relations between different modalities remain similar [20].

Moderate hypofractionation
For low and intermediate risk (IR) prostate cancer, moderate hypofractionation has been shown to be non-inferior to normo-fractionated treatment in several prospective randomized trials [21-24] and is now strongly recommended in the primary setting by NCCN guidelines [25] and viewed as a viable alternative in current EAU [21] and German S3 Guidelines (https://www.leitlinienprogramm-onkologie.de/leitlinien/prostatakarzinom/). The latter two explicitly advise its performance only by experienced teams using high-quality EBRT (IGRT
and IMRT) in carefully selected patients with strong adherence to published phase III protocols.
For HR patients, the benefit of hypofractionated radiotherapy is less clear. Analyses from 3 large meta-analyses [7,8,6] comprising more than 20000 patients have yielded low α/β values for all risk groups, which led Fowler et al. to conclude that ‘the low α/β ratio is an intrinsic property of all prostate cancer cells irrespective of their Gleason score or grading’ [26]. However, clinical data to support this notion are still lacking. In the three large non-inferiority trials [22-24], HR patients were underrepresented. In the superiority-design HYPRO trial [25] HR patients were included, but in that trial the primary endpoint (improved biochemical control at 5 years) did not reach significance and toxicity was slightly higher in the hypofractionated (and dose escalated) arm.

**Literature review**

**RANDOMIZED EVIDENCE:**

**The Scandinavian trial (HYPOR-RT-PC)**
The Scandinavian non-inferiority design HYPO-RT-PC trial by Widmark et al. [26] randomized men with IR to HR prostate cancer to receive either 42.7Gy in seven fractions, 3 days per week, or conventionally fractionated radiotherapy (78Gy in 39 fractions, 5 days per week). After a median follow up of 5 years failure free survival was identical (84%) in both arms. Acute RTOG G2 or worse genitourinary (GU) toxicity was slightly but not significantly increased in the ultrahypofractionated arm at the end of treatment (28% vs 23%, p=0.057). A significant rise was only seen at 1-year of follow-up (6% vs 2%, p=0.0037) disappearing completely at timepoints thereafter (5-year rate: 5% in both arms). There was no difference in gastrointestinal (GI) toxicity at any time point and no differences in toxicity after 5 years. Of note, the Widmark trial features some peculiarities and differences compared to the abundant but retrospective trial protocols that have accumulated in the past decade. These differences need to be critically reviewed before routine clinical application:
1) It excluded LR and included intermediate and high risk patients: to our knowledge, the Widmark study is the only ultrahypofractionation study in which LR patients were excluded. In addition, a particular subset of HR patients (PSA <20ng/ml and T3a) were included (11%) which is in stark contrast to the low percentage of HR patients treated within the published retrospective series.

2) It used 7 instead of the commonly reported 5 fractions, which is unique among the ultrahypofractionation trials.

3) No androgen deprivation therapy (ADT) was given: In both arms, ADT was withheld to all patients, which might have had a negative impact on progression free survival (PFS) as well as overall survival (OS) for (unfavorable) IR and HR patients. For these risk groups, there is no evidence that either hypofractionation or dose escalation (or both) can compensate for the lack of ADT which is known to improve both, biochemical control as well as OS [27].

4) The LINAC-based radiation technique did not have to meet highest standards. Neither MR imaging for contouring nor treatment by IMRT were mandatory. In fact, 80% of patients were treated with conventional 3D planning (commented in [28]).

5) Contouring and margins: Seminal vesicles were not included in the CTV which is questionable since EORTC guidelines recommend including the proximal 1-2cm for IR and HR patients, respectively [29]. PTV margins were rather large (7mm) even though image guidance was used using either beam cath (10%) or gold fiducials (90%).

PACE-B
Early toxicity results of the randomized PACE-B trial have recently been published [30].
In that non-inferiority trial, men with LR or IR prostate cancer (Gleason 7b excluded) received either conventional or moderately hypofractionated radiotherapy (78 Gy in 39 fractions in 7-8 weeks or 62 Gy in 20 fractions over 4 weeks, respectively), or SBRT (36.25 Gy in five fractions over 1–2 weeks). ADT was not permitted.
41% of patients in the SBRT arm were treated with CyberKnife 58.3% with a conventional LINAC using volumetric arc therapy (VMAT). IGRT and intra-fraction motion-control were mandatory.
In terms of acute toxicity, there was no significant difference between arms, but a slight trend in favor of the SBRT Arm (23% vs 27%). That is in contrast to the Widmark trial, where a trend for increased acute toxicity was seen in the ultrahypofractionated arm.

METAANALYSES OF NON-RANDOMIZED, PROSPECTIVE DATA

Well over 10,000 patients have been treated within ultrahypofractionated non-randomized, prospective protocols with large variations in fraction size, total dose and radiation technique. The most relevant studies based on quality and patient cohort size have been summarized and re-analyzed by three large pooled analyses [31-33].

King et al. 2013 [31]
The first pooled analysis by King et al. included 1100 patients from 8 institutions who have been treated within prospective phase II trials using CyberKnife with a median follow up of 36 months. They received a median dose of 36.25Gy in 4-5 fractions. LR (58%), IR (30%) and HR (11%) patients were included. A short-course of ADT was given to 14% of patients. The five-year biochemical relapse free survival (bRFS) rate was 93% for all patients and 95%, 84% and 81% for LR, IR- and HR patients, respectively (p < 0.001). Toxicity was not reported.

Kishan et al. 2019 [32]
The second analysis is a cohort study which analyzed individual patient data from 12 phase II trials comprising 2142 men with low and IR prostate cancer, treated with either CyberKnife (7/12 studies) or a conventional LINAC (5/12 studies). 55.3% of patients had LR disease, 32.3% had favorable IR disease, and 12.4% unfavorable IR disease. HR patients were excluded. The follow up period was quite long with a median of 6.9 years. Seven-year biochemical free survival (bRFS) amounted to 95.5% for LR disease and 89.8% for IR disease. The crude incidence of acute grade 3 or higher toxic events was 0.60% for GU and 0.09% for GI side effects.

Jackson et al. [33]
The most recent and more extensive review has been undertaken by Jackson et al. comprising 6116 patients from 38 prospective studies. There was a large patient overlap with the patient collectives of the former two analyses by King and Kishan et al. A meta-analysis using random effect modeling was performed on a study-level basis. Only studies reporting the same outcome at the same time point were pooled which is an inherent limitation.

On a patient level 45% had LR, 47% had IR and 8% HR disease. Median follow up was 39 months, but 5- and 7-years bRFS rates and toxicities have been reported, not complying with the RTOG-ASTRO Phoenix consensus which recommends the reported date of control be listed as 2 years short of the median follow up [34].

Combined acute ≥G3 toxicity was below 1%. Late ≥G3 GU and GI toxicity was 2.0% and 1.1%, respectively and did not change when only studies with a median FU of ≥5yr were analyzed. There was a significant publication bias, which – when corrected for – increased toxicity rates by 1% to 2%. Interestingly, there was an association of dose and late grade ≥G3 GU toxicity but not with ≥G3 GI toxicity. The authors conclude that ultrahypofractionation could be considered a standard radiotherapeutic strategy for localized prostate cancer – maybe a premature statement given how underrepresented HR patients were in that study.

TREATMENT OF THE PRIMARY IN LOW-VOLUME METASTATIC DISEASE SETTING
Two recent prospective randomized trials (HORRAD [35] and STAMPEDE [36]) have addressed the role of RT to the prostate in metastatic disease. Ultrahypofractionation is an appealing option in this scenario and has been used optionally in the STAMPEDE trial in which 48% of patients were treated with 36Gy in 6 weekly fractions corresponding to an EQD2 with β 2Gy of 72Gy. The STAMPEDE sub-group analysis of low-volume metastatic disease demonstrated a survival advantage in favor of the RT arm (hazard ratio 0.68; 95% CI 0.52-0.90). The HORRAD trial showed a similar but non-significant trend towards an improved OS by RT (hazard ratio 0.68; 95% CI 0.42-1.10). As a result, the 2019 European Association of Urology and National Comprehensive Cancer Network guidelines now include RT to the prostate as an option in the setting of low-volume metastatic disease [37].

Interpretation
An accumulating body of retrospective evidence for the safety and efficacy of SBRT for low and IR prostate cancer has recently been strengthened by the publication of two randomized studies comparing SBRT with a normo-fractionated schedule, i.e. the Scandinavian trial by Widmark et al. [26] and the first toxicity results of the PACE-B trial [30]). Since the PACE study has not yet reached sufficient follow up to report on outcome or late toxicity, the Scandinavian trial thus far is the only randomized trial comparing an ultrahypofractionated to a normofractionated schedule with reported long term (i.e. 5 years) outcome and toxicity. It is therefore the only level Ib evidence (according to the Oxford Centre for Evidence-Based Medicine. (https://www.cebm.net/index.aspx?o=5653) available and has raised the grade of recommendation from C to B. This could justify the clinical use of ultrahypofractionation outside of clinical trials for low and intermediate risk prostate cancer.

HR patients should continue to be treated within clinical trials due to several reasons: First, the body of evidence for LR and IR prostate cancer is overwhelmingly larger than for HR prostate cancer patients constituting well below 8% of studied patients. In addition, it remains disputable whether Gleason 8-10 prostate cancer cells feature an equally low α/β value although this has been postulated [38]. Third, ultrahypofractionation of the prostate hampers the simultaneous coverage of pelvic lymph-nodes. Pelvic RT in HR patients is endorsed by many institutions, although its value is still controversially discussed. However, an ultrahypofractionated boost with reduced dose after whole pelvic RT might be an attractive concept [39,40].

ADT has been used inconsistently in the available ultrahypofractionation trials. However, at the present time there is no evidence that high dose can compensate for the lack of ADT in higher risk prostate cancer. Therefore, ADT should be administered according to the current guidelines i.e. short term ADT (4-6 months) for unfavorable IR and long term ADT (18-36 months) for HR prostate cancer). For favorable IR the omission of ADT seems appropriate in a dose escalated setting [27].

For ultrahypofractionation, the ideal fractionation regimen has not yet been established. A variety of schedules have been published and can be considered safe (see table 1 for select examples and corresponding EQD2 for different α/β values). With the exception of the
Scandinavian trial by Widmark et al, most larger series used doses of 36.25 Gy to 40 Gy in 5 fractions administered every other day. In the meta-analysis by Jackson et al, the median fraction size was 7.4 Gy. For unfavorable IR patients slightly higher total doses may be considered. The 7 fraction schedule by Widmark et al. has been tested in a randomized prospective fashion and can be considered a viable option for IR patients.

For treatment of the primary in the metastatic setting ultrahypofractionation is an attractive alternative to moderate hypofractionation and has been tested in the STAMPEDE trial. The prolongation of the overall treatment time to 6 weeks is rather unusual for a ultrahypofractionated regimen and may be the reason why in this trial, the alternatively used moderately hypofractionated radiation schedule of 55 Gy in 20 fractions (daily) showed a slightly better outcome (not significant, HR 0.86 vs 1.01).

Compared to radiation schedules in the non-metastatic setting, the resulting BEDs of these doses are rather conservative which is owed to the palliative setting where the maxim ‘primum non nocere’ is to be followed strictly. In general, further studies are needed to establish an appropriate fractionation schedule in the metastasized setting, but if ultrahypofractionation is used, the DEGRO Prostate Cancer Expert Panel favors every-other-day-schedules over once-weekly regimen and recommends aiming for a total dose of at least EQD2\^{α/β2} 72 Gy.

**Summary**

Retrospective as well as randomized prospective data with a follow up of 5 years or more are now available and have shown comparable results to recent moderate hypofractionation trials in terms of both, biochemical control and toxicity. Although level Ib evidence for ultrahypofractionation is now available, the DEGRO prostate cancer expert panel does not yet recommend ultrahypofractionation for HR patients on a routine basis. However, for centers that are experienced in SBRT and wish to offer ultrahypofractionation to select LR and IR patients, this seems justified outside of a clinical trial (Grade B recommendation).

In interpretation of the published data, the following principles should be followed when administering ultrahypofractionation in prostate cancer in clinical routine:
1. Ultrahypofractionation is a treatment alternative, amongst moderate hypofractionation and normofractionation, which can be offered outside clinical trials to LR patients who are not suitable for active surveillance and for IR patients including both favorable and unfavorable IR.

2. ADT should be administered according to current guidelines for normo- and moderately hypofractionated regimen i.e. short term ADT for unfavorable IR (and long term ADT for HR patients). For favorable IR dose escalated RT alone appears to be an appropriate treatment.

3. Ultrahypofractionation should be administered in a setting of high technical standards. We consider MR-based planning, IMRT/VMAT and daily IGRT with basic intrafraction control (i.e. imaging after 3min of treatment time) as minimum requirements in order to safely achieve PTV margins of approximately 3mm. In a LINAC based setting short treatment times need to be pursued using single arc VMAT and/or flattening filter free (FFF) techniques.

4. Dose schedules should strictly adhere to published concepts of larger studies. A maximum fraction size of 8Gy and a maximum EQD2$^{\alpha/\beta}$ of 100Gy should not be exceeded. In the curative, non-metastatic setting, a minimum EQD2$^{\alpha/\beta}$ of 83.3Gy (e.g. 5x7.25Gy) is required. See table 1 for different fractionation schedules and corresponding EQD2 at different $\alpha/\beta$ values.

5. For treatment of the primary in the metastatic setting ultrahypofractionation may be used as an alternative to moderate hypofractionation. However, slightly de-escalated schedules with an EQD2$^{\alpha/\beta}$ of approximately 72Gy-76Gy should be used.

6. Patients should be followed up by the treating facility for at least 5 years. Inclusion in a registry study is recommended.
Table 1. Treatment schedules and corresponding EQD2 for different α/β values.

|                      | dose/fx | # fx | total dose | α/β 1.5 | α/β 2 | α/β 3 | α/β 10 |
|----------------------|---------|------|------------|---------|-------|-------|--------|
| HYPO RT-PC           | 6.1     | 7    | 42.7       | 92.72   | 86.47 | 77.71 | 57.29  |
| PACE                 | 7.25    | 5    | 36.25      | 90.63   | 83.83 | 74.31 | 52.11  |
| Median fx Jackson et al. | 7.4    | 5    | 37         | 94.09   | 86.95 | 76.96 | 53.65  |
| Mantz et al. 2014    | 8       | 5    | 40         | 108.57  | 100.00| 88.00 | 60.00  |
| Fuller et al. 2014   | 9.5     | 4    | 38         | 119.43  | 109.25| 95.00 | 61.75  |
| STAMPEDE primary metast. | 6      | 6    | 36         | 77.14   | 72.00 | 64.80 | 48.00  |
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