Multimodal treatment for high-risk prostate cancer with high-dose intensity-modulated radiation therapy preceded or not by radical prostatectomy, concurrent intensified-dose docetaxel and long-term androgen deprivation therapy: results of a prospective phase II trial

Andrea Guttilla1*, Roberto Bortolus2, Gianluca Giannarini1,3, Pirus Ghadjar4, Fabio Zattoni1, Michele Gnech1, Vito Palumbo1, Francesca Valent5, Antonio Garbeglio6 and Filiberto Zattoni1

Abstract

Background: The optimal management of high-risk prostate cancer remains uncertain. In this study we assessed the safety and efficacy of a novel multimodal treatment paradigm for high-risk prostate cancer.

Methods: This was a prospective phase II trial including 35 patients with newly diagnosed high-risk localized or locally advanced prostate cancer treated with high-dose intensity-modulated radiation therapy preceded or not by radical prostatectomy, concurrent intensified-dose docetaxel-based chemotherapy and long-term androgen deprivation therapy. Primary endpoint was acute and late toxicity evaluated with the Common Terminology Criteria for Adverse Events version 3.0. Secondary endpoint was biochemical and clinical recurrence-free survival explored with the Kaplan-Meier method.

Results: Acute gastro-intestinal and genito-urinary toxicity was grade 2 in 23% and 20% of patients, and grade 3 in 9% and 3% of patients, respectively. Acute blood/bone marrow toxicity was grade 2 in 20% of patients. No acute grade ≥4 toxicity was observed. Late gastro-intestinal and genito-urinary toxicity was grade 2 in 9% of patients each. No late grade ≥3 toxicity was observed. Median follow-up was 63 months (interquartile range 31–79). Actuarial 5-year biochemical and clinical recurrence-free survival rate was 55% (95% confidence interval, 35-75%) and 70% (95% confidence interval, 52-88%), respectively.

Conclusions: In our phase II trial testing a novel multimodal treatment paradigm for high-risk prostate cancer, toxicity was acceptably low and mid-term oncological outcome was good. This treatment paradigm, thus, may warrant further evaluation in phase III randomized trials.

Keywords: Prostatic neoplasms, Radiotherapy, Intensity-modulated, Prostatectomy, Androgen deprivation therapy, Chemotherapy, Docetaxel, Toxicity, Clinical trial, Phase II
**Background**

Depending on the definition, high-risk localized or locally advanced prostate cancer (PCa) accounts for 15-40% of newly diagnosed cases [1,2]. Despite advances in patient selection and primary/adjuvant therapies, disease recurrence remains substantial, affecting >50% of patients within 10 years after treatment [3,4].

In the absence of adequately conducted randomized trials, optimal treatment remains controversial. External beam radiation therapy (EBRT) with long-term androgen deprivation therapy (ADT) has long been considered the standard of care over other modalities [5,6]. The survival benefit of long-term ADT combined with EBRT vs. EBRT alone has recently been confirmed in a meta-analysis [7]. Over the past decade, however, there has been a surge in the utilization of radical prostatectomy (RP) as primary treatment in selected patients, followed by EBRT in case of adverse pathological features [5,6]. It is currently accepted that a combined modality-regimen should be pursued in clinical trials, although sequence, timing and intensity of the single treatments are still under scrutiny [8].

**Table 1** Clinical and tumor characteristics of patients with high-risk localized or locally advanced prostate cancer enrolled in our phase II trial

| Variable                        | Cohort 1 (n=18) | Cohort 2 (n=17) |
|---------------------------------|-----------------|-----------------|
| Age, years, median (IQR)        | 59 (56–63)      | 69 (67–73)      |
| Serum PSA level, ng/ml, median (IQR) | 17 (8–26)      | 18 (9–28)      |
| Clinical T stage, n             |                 |                 |
| T2                              | 6               | 5               |
| T3                              | 12              | 11              |
| T4                              | 0               | 1               |
| Clinical N stage, n             |                 |                 |
| N0                              | 16              | 15              |
| N+                              | 2               | 2               |
| Biopsy Gleason score, n         |                 |                 |
| 6                               | 3               | 2               |
| 3+4                             | 1               | 2               |
| 4+3                             | 4               | 7               |
| 8-10                            | 9               | 6               |
| Pathological T stage*, n        |                 |                 |
| T2c                             | 2               | NA              |
| T3a                             | 3               |                 |
| T3b                             | 11              |                 |
| T4                              | 2               |                 |
| Positive lymph node status, n   | 4               | NA              |
| Positive surgical margins, n    | 8               | NA              |
| Pathological Gleason score, n   | 8               | 5               |
| 9                               | 13              |                 |
| Follow up, months, median (IQR) | 61 (45–80)      | 63 (26–77)     |

IQR = interquartile range; NA = not applicable; PSA = prostate specific antigen; *according to the 2002 TNM classification.

**Table 2** Acute adverse events recorded during concurrent high-dose intensity-modulated radiation therapy and docetaxel-based chemotherapy: allergy/immunology

| Grade | Allergic reactions | Highest |
|-------|-------------------|---------|
| Cohort 1 | 1 | 1 |
| Cohort 2 | 2 | 0 |
| 3 | 0 | 0 |
| 1 | 0 | 0 |

**Methods**

**Study population**

Eligible subjects were men aged ≥18 years with histologically confirmed PCa and high-risk localized (PSA level ≥20 ng/ml or clinical stage T2c or Gleason score ≥8) or locally advanced (clinical stage T3 or N+, any PSA level, any Gleason score) disease. All patients had abdominal computed tomography, chest X-ray and bone scan excluding distant metastases. Exclusion criteria were: Eastern Cooperative Oncology Group performance status >1; hematological abnormalities (absolute neutrophil count <1500/mm³, hemoglobin <8.0 g/dl, platelets <10⁵/mm³); liver function tests abnormalities (total bilirubin >1.2 mg/dl, alanine and aspartate transaminase >1.5× the upper normal limit, docetaxel is an active agent in the treatment of PCa but, until recently, its role has been limited to patients with metastatic castration-resistant disease [9,10]. However, in order to treat micrometastatic disease, an earlier use of chemotherapy within the curative treatment setting may be beneficial. Docetaxel is also known as a radiosensitizer, and has been shown to improve local control in combination with EBRT in the treatment of lung, head/neck and cervical cancers [11,12]. Furthermore, recent evidence suggests that a positive interaction effect exists also between docetaxel and ADT [13]. All these premises have made docetaxel a suitable agent to integrate into multimodal treatment paradigms for high-risk PCa.

Few studies have reported on the safe use and efficacy of docetaxel at a weekly dose of 20 mg/m² in combination with conventional EBRT in hormone-naïve or ablated high-risk PCa [14-16]. Very little, however, is known about the effect of docetaxel administered concurrently with state-of-the-art high-dose EBRT, e.g. enabled by intensity-modulated radiation therapy (IMRT), and long-term ADT.

In a phase II trial, we evaluated the safety and efficacy of a novel multimodal treatment paradigm for high-risk PCa including high-dose IMRT preceded or not by RP with concurrent intensified-dose docetaxel-based chemotherapy and long-term ADT.

**Table 2 Acute adverse events recorded during concurrent high-dose intensity-modulated radiation therapy and docetaxel-based chemotherapy: allergy/immunology**

**Table 2 Acute adverse events recorded during concurrent high-dose intensity-modulated radiation therapy and docetaxel-based chemotherapy: allergy/immunology**

| Grade | Allergic reactions | Highest |
|-------|-------------------|---------|
| Cohort 1 | 1 | 1 |
| Cohort 2 | 2 | 0 |
| 3 | 0 | 0 |
| 1 | 0 | 0 |
alkaline phosphatase >2.5× the upper normal limit); renal function tests abnormalities (creatinine >1.5× the upper normal limit); myocardial infarction or unstable angina within 1 year prior to study entry; congestive heart failure (New York Heart Association Class ≥2); uncontrolled chronic disease; severe infection; peripheral neuropathy; prior malignancy (except for non-melanoma skin cancer) within the last 5 years before study entry; prior pelvic external beam radiation therapy; prior chemotherapy; prior prostate surgery; prior treatment for prostate cancer.

The protocol was approved by the local ethics committee of the National Cancer Institute Aviano, and the study was conducted according to the Declaration of Helsinki and European Good Clinical Practice requirements. Written informed consent was obtained at study entry.

Study design and treatments

Patients were offered one of two multimodal therapy strategies depending on their age at PCa diagnosis. Patients aged ≤65 years were preferentially offered initial open non-nerve-sparing RP with extended pelvic lymph node dissection (PLND) according to a previously reported, anatomically defined template [17], adjuvant EBRT within 3–6 months postoperatively with concurrent docetaxel and long-term ADT (cohort 1). Older patients were preferentially offered primary EBRT with concurrent docetaxel and long-term ADT (cohort 2).

Surgery was performed at two centres (Padua and Pordenone).

| Table 3 Acute adverse events recorded during concurrent high-dose intensity-modulated radiation therapy and docetaxel-based chemotherapy: blood/bone marrow |
|-----------------------------------|-----------|-------------|-------------|-------------|
| Grade | Anemia | Leucopenia | Neutropenia | Thrombocytopenia | Highest |
|-------|--------|------------|-------------|-----------------|--------|
| Cohort 1 | 2      | 1          | 3           | 0               | 6      |
|       | 3      | 0          | 1           | 0               | 3      |
|       | 1      | 6          | 5           | 2               | 5      |
|       | 1      | 2          | 4           | 1               | 4      |
|       | 3      | 0          | 0           | 0               | 0      |
| Cohort 2 | 2      | 2          | 4           | 1               | 4      |
|       | 3      | 0          | 0           | 0               | 0      |

All patients were treated with IMRT in Aviano. A 4- to 6-field technique with 15- to 18-MV photon beams was used. No image-guided radiation therapy (IGRT) was used.

In patients of cohort 1, clinical target volume (CTV) included the area from the level of the caudal remnant of the vas deferens, which was intraoperatively marked with clips, to 10 mm inferior to the vesico-urethral anastomosis, from the posterior aspect of the pubis to the anterior rectal wall, and between the medial aspects of each obturator internus muscle. Planning target volume (PTV) consisted of CTV plus a 5-mm margin in all directions except posteriorly, where a 3-mm margin was set. No radiation to the pelvis was given. A total dose of 70 Gy in 35 fractions was prescribed. Dose constraints for organs at risk were: a) bladder: V65 Gy ≤50%, V40 Gy ≤70%; b) rectum: V65 Gy ≤35%, V40 Gy ≤55%; c) femoral heads: V50 Gy ≤10%.

In patients of cohort 2, CTV1 included the pelvic area encompassing external iliac, obturator, internal iliac and lower common iliac lymph node chains bilaterally, from the inferior margin of L5 to the inferior margin of the inferior pubic branch, and CTV2 included prostate and seminal vesicles. PTV1 consisted of CTV1 plus a 10-mm margin in all directions, and PTV2 consisted of CTV2 plus a 10-mm margin in all directions, except posteriorly, where a 6-mm margin was set. A total dose of 80 Gy in 40 fractions was prescribed in three phases: i) first phase: 26 Gy in 13 fractions to PTV1 + PTV2; ii) second phase: 30 Gy in 15 fractions to PTV2; iii) third phase: 24 Gy in 12 fractions to PTV1 + PTV2. Dose constraints for organs at risk were: a) bladder: V80

| Table 4 Acute adverse events recorded during concurrent high-dose intensity-modulated radiation therapy and docetaxel-based chemotherapy: constitutional symptoms |
|-----------------------------------|-------------|-------------|-------------|
| Grade | Fatigue | Fever | Hot flashes | Highest |
|-------|---------|-------|-------------|---------|
| Cohort 1 | 2 | 1 | 2 | 4 |
|       | 3 | 0 | 0 | 0 |
| Cohort 2 | 1 | 1 | 2 | 3 |
|       | 2 | 1 | 1 | 3 |

| Table 5 Acute adverse events recorded during concurrent high-dose intensity-modulated radiation therapy and docetaxel-based chemotherapy: dermatology/skin |
|-----------------------------------|-------------|-------------|-------------|
| Grade | Rash/dermatitis | Highest |
|-------|-----------------|---------|
| Cohort 1 | 2 | 0 | 0 |
|       | 3 | 0 | 0 |
| Cohort 2 | 1 | 1 | 1 |
|       | 2 | 0 | 0 |
|       | 3 | 0 | 0 |
Gy ≤15%, V70 Gy ≤35%, V50 Gy ≤60%; b) rectum: V70 Gy ≤20%, V60 Gy ≤35%, V50 Gy ≤50%; c) femoral heads: V50 Gy ≤5%.

Docetaxel was administered starting on the same week as IMRT in a standard 1-hour intravenous weekly dose (30 mg if body surface area <1.8 m² and 40 mg if body surface area ≥1.8 m²) for 7 and 8 weeks in patients of cohort 1 and 2, respectively. Premedication with intravenous methylprednisolone, chlorphenamine and ondansetron was given prior to each docetaxel infusion.

ADT, consisting of an injection of a 3-monthly LHRH-agonist, was started either immediately after surgery in cases of pathologically proven seminal vesicle and/or lymph node invasion or 1 to 2 months before IMRT in patients of cohort 1, and 2 to 3 months before treatment in patients of cohort 2. No antiandrogens were given. ADT was prescribed for 24 months in all patients.

Follow-up
During concurrent IMRT/docetaxel physical examination, complete blood count and toxicity assessment were obtained weekly. After completion, patients were followed with physical examination, urinalysis, serum PSA, liver and renal function tests 3-monthly for the first 2 years, 6-monthly for the following 3 years, and annually thereafter. Follow-up began either on the date of RP (cohort 1) or the date of IMRT completion (cohort 2), and continued until last observation or death.

Study endpoints
Primary endpoint was safety. Complications after RP were assigned according to the modified Clavien classification system [18]. Toxicity related to concurrent IMRT/docetaxel was scored according to the National Cancer Institute Common Terminology Criteria for Adverse Events version 3.0 [19]. Baseline gastro-intestinal (GI) and genito-urinary (GU) symptoms before IMRT/docetaxel initiation were investigated. Acute toxicity was defined as occurring during and until 3 months after IMRT/docetaxel, while late toxicity as occurring after 3 months until the time of administration of salvage treatment. Similarly to previous investigators [20], toxicity was defined as increase over baseline value for each symptom. All adverse events occurring in a given patient at any follow-up visit were recorded, but only the highest toxicity grade per patient was counted as event when calculating the toxicity rate. To explore toxicity change with time, the incidence of late toxicity at the last follow-up visit was also reported.

Secondary endpoint was efficacy. Biochemical recurrence was defined as an increase in PSA level above 0.2 ng/ml [5] for patients treated with RP or an increase of 2 ng/ml above nadir [21] for patients treated with primary IMRT. Clinical recurrence was defined as local or regional (pelvic lymph nodes) tumour growth, or distant metastases (including retroperitoneal lymph nodes). Biochemical recurrence-free survival (BRFS) and clinical recurrence-free survival (CRFS) were calculated from

Table 6 Acute adverse events recorded during concurrent high-dose intensity-modulated radiation therapy and docetaxel-based chemotherapy: gastrointestinal symptoms

| Grade | Nausea/vomiting | Diarrhea | Gastritis | Proctitis | Hematochezia | Highest |
|-------|-----------------|----------|-----------|-----------|--------------|---------|
| Cohort 1 | 2 | 2 | 3 | 2 | 2 | 2 | 8 |
| Cohort 2 | 2 | 2 | 3 | 1 | 1 | 1 | 7 |

Study endpoints
Primary endpoint was safety. Complications after RP were assigned according to the modified Clavien classification system [18]. Toxicity related to concurrent IMRT/docetaxel was scored according to the National Cancer Institute Common Terminology Criteria for Adverse Events version 3.0 [19]. Baseline gastro-intestinal (GI) and genito-urinary (GU) symptoms before IMRT/docetaxel initiation were investigated. Acute toxicity was defined as occurring during and until 3 months after IMRT/docetaxel, while late toxicity as occurring after 3 months until the time of administration of salvage treatment. Similarly to previous investigators [20], toxicity was defined as increase over baseline value for each symptom. All adverse events occurring in a given patient at any follow-up visit were recorded, but only the highest toxicity grade per patient was counted as event when calculating the toxicity rate. To explore toxicity change with time, the incidence of late toxicity at the last follow-up visit was also reported.

Secondary endpoint was efficacy. Biochemical recurrence was defined as an increase in PSA level above 0.2 ng/ml [5] for patients treated with RP or an increase of 2 ng/ml above nadir [21] for patients treated with primary IMRT. Clinical recurrence was defined as local or regional (pelvic lymph nodes) tumour growth, or distant metastases (including retroperitoneal lymph nodes). Biochemical recurrence-free survival (BRFS) and clinical recurrence-free survival (CRFS) were calculated from

Table 7 Acute adverse events recorded during concurrent high-dose intensity-modulated radiation therapy and docetaxel-based chemotherapy: genitourinary symptoms

| Grade | Frequency/urgency | Dysuria | Incontinence | Retention | Hematuria | Highest |
|-------|------------------|---------|--------------|-----------|-----------|---------|
| Cohort 1 | 1 | 3 | 6 | 3 | 1 | 2 | 1 | 3 | 4 | 1 | 0 | 2 | 2 | 0 | 1 | 1 | 9 | 5 |
| Cohort 2 | 2 | 3 | 5 | 2 | 0 | 1 | 1 | 0 | 2 | 2 | 2 | 3 | 5 | 2 | 0 | 2 | 2 | 8 | 6 |

B = baseline (before concurrent intensity-modulated radiation therapy and docetaxel-based chemotherapy); A = all acute adverse events; A* = acute adverse events of higher grade than baseline.
the date of RP or IMRT completion to the date of biochemical and clinical recurrence, respectively, or death from any cause.

Statistical analyses
A Simon’s two-stage design was used to calculate the optimal sample size [22]. The trial was designed to test an acute grade $\geq 3$ toxicity rate of $\leq 5\%$, considered as acceptable to design a phase III trial, against an acute grade $\geq 3$ toxicity rate of $\geq 20\%$, considered as unacceptable, with an 85% power and a 5% significance level. Stopping rule was grade $\geq 3$ toxicity in $\geq 4$ patients during the first stage (17 patients) or in $\geq 3$ patients in the second stage (18 patients). Interim analyses in the second stage were planned every 6 patients.

Continuous variables were non-normally distributed according to Kolmogorov-Smirnov test, and are reported as median with interquartile range (IQR). All analyses were based on intention to treat. Kaplan-Meier estimates were used to explore BRFS and CRFS. Analyses were performed with SAS Enterprise Guide v4.3 (SAS Institute Inc., Cary, NC, USA).

Results
Between January 2005 and March 2012, 35 men, 18 in cohort 1 and 17 in cohort 2, were enrolled. All patients had complete baseline and follow-up data and were available for the present analysis. Median age was 64 years (IQR 58-69). Median follow-up was 63 months (IQR 31-79). Details are reported in Table 1.

All patients in cohort 1 and 14 in cohort 2 completed IMRT to the prescribed dose. Three patients in cohort 2 received 78 Gy because of non-compliance. Eight and 27 patients received 30 mg and 40 mg docetaxel, respectively. Chemotherapy was interrupted in two patients in cohort 2 on initiation of the 2nd cycle because of grade 3 allergic reaction. All patients completed ADT for the prescribed duration.

Safety
In the first stage, three patients had grade $\geq 3$ toxicity, hence the trial was continued. In the second stage, two further patients had grade $\geq 3$ toxicity, hence the trial was completed.

Perioperative complications after RP and extended PLND were prolonged (>14 days) lymphorrhea (n = 1) (grade I “d”), pelvic hematoma (n = 1) and urinary tract infection (n = 2) (both grade II).

Before IMRT/docetaxel, GI symptoms were absent, while GU symptoms were present in 11 patients, five in cohort 1 and six in cohort 2.

Acute adverse events were common, but most of grade $\leq 2$. All grade 3 events resolved by the time of the subsequent chemotherapy cycle. No grade 4 adverse events were observed. GI and GU symptoms prevailed. Acute GI and GU toxicity was grade 2 in 23% and 20% of patients, and grade 3 in 9% and 3% of patients, respectively. Acute blood/bone marrow toxicity was grade 2 in 20% of patients. All details are reported in Tables 2,3,4,5,6,7.

The rate of late adverse events was low and no grade $\geq 3$ toxicity was observed. Late GI and GU toxicity was grade 2 in 9% of patients each. At the last follow-up visit, grade 2 GI and GU toxicity was present in 0% and 6% of pa-

Table 8 Late adverse events recorded after concurrent high-dose intensity-modulated radiation therapy and docetaxel-based chemotherapy: constitutional symptoms

| Grade | Fatigue | Hot flashes | Highest during follow-up | Highest at last follow-up |
|-------|---------|-------------|--------------------------|--------------------------|
| Cohort 1 | 2 | 1 | 3 | 3 | 2 |
| 3 | 0 | 0 | 0 | 0 | 0 |
| Cohort 2 | 2 | 1 | 1 | 1 | 1 |
| 3 | 0 | 0 | 0 | 0 | 0 |

Table 9 Late adverse events recorded after concurrent high-dose intensity-modulated radiation therapy and docetaxel-based chemotherapy: gastrointestinal symptoms

| Grade | Diarrhea | Gastritis | Proctitis | Hematochezia | Highest during follow-up | Highest at last follow-up |
|-------|----------|-----------|-----------|--------------|--------------------------|--------------------------|
| Cohort 1 | 2 | 0 | 1 | 1 | 1 | 1 |
| 3 | 0 | 0 | 0 | 0 | 0 | 0 |
| Cohort 2 | 2 | 1 | 0 | 2 | 2 | 3 |
| 3 | 0 | 0 | 0 | 0 | 0 | 0 |
tients. No patients underwent surgery for bladder outlet obstruction or urethral stricture. Details are reported in Tables 8,9,10.

Efficacy
One month after RP 12 patients had an undetectable (<0.01 ng/ml) PSA level. Three months after IMRT/docetaxel completion, all patients of cohort 1 had an undetectable PSA level, and all patients of cohort 2 achieved a post-treatment PSA level of ≤1 ng/ml, including 9 patients who achieved an undetectable level.

Fourteen patients (40%), eight in cohort 1 and six in cohort 2, had biochemical recurrence. Actuarial 5-year BRFS rate was 55% (95% CI, 35-75%), 46% (95% CI, 19-73%) for cohort 1 and 68% (95% CI, 42-95%) for cohort 2 (Figure 1a).

Ten patients (29%), five in each cohort, had clinical recurrence. Local recurrence, regional recurrence and distant metastases were detected in three, two and three patients, respectively. The two remaining patients had concomitant local and regional recurrence, and regional recurrence and distant metastases. Actuarial 5-year CRFS rate was 70% (95% CI, 52-88%), 74% (95% CI, 51-96%) for cohort 1 and 65% (95% CI, 36-94%) for cohort 2 (Figure 1b).

Salvage treatment was high-intensity focused ultrasound prostate ablation (n = 1), helical tomotherapy (n = 2), ADT (n = 7), chemotherapy (n = 3), or supportive care (n = 1). Four patients (all in cohort 1) died, two of PCa and two of cardiovascular disease in absence of disease recurrence and ADT.

Discussion
In the present study, we explored safety and efficacy of a novel multimodal treatment regimen for high-risk PCa based on high-dose IMRT preceded or not by RP, concurrent intensified-dose docetaxel and long-term ADT. Acute toxicity was common but mostly of mild to moderate severity, late toxicity rate was low with no severe adverse events, and mid-term oncological outcome was good.

Six trials have explored the role of concurrent EBRT and taxane-based chemotherapy in patients with high-risk PCas [14-16,23-25] (Table 11). Direct comparison between these and the present trial is hampered by differences in study design, type of EBRT and chemotherapeutic agent used, ADT duration, toxicity assessment and follow-up length. Notably, all previous studies but one [25] applied conventional-dose EBRT only (using either three-dimensional conformal radiation therapy [3D-CRT] or IMRT), which is now considered suboptimal. A meta-analysis, in fact, concluded that high-dose EBRT is associated with a significant improvement in BRFS compared to conventional-dose EBRT, regardless of PCa risk category [26]. Additionally, IMRT has been shown to be able to deliver a higher dose to the prostate while reducing GI and GU toxicity compared to 3D-CRT [27], thus it is currently regarded the most effective form of EBRT.

The only available trial in which high-dose IMRT was used [25] was a phase I trial, where 18 patients with high-risk PCas were treated with high-dose IMRT, concurrent docetaxel at increasing doses and long-term ADT. Only three patients received a dose of 20 mg/m² and none experienced ≥ grade 3 adverse events. In this context, our phase II trial, which used high-dose IMRT, has particular value. In our study, toxicity was common, but was mostly grade ≤2. Most symptoms were of the GI and GU tract. A strength of our analysis is the detailed reporting of both acute and late toxicity, which included the evaluation of baseline symptoms.

Our study included also high-risk PCa patients with prior RP, who may exhibit a different acute and late toxicity profile from their surgery-naïve counterparts, in spite of the lower radiation dose delivered. To the best of our knowledge, only one study [24] explored a similar treatment paradigm also in RP patients. Fifty-nine men with high-risk PCas, including 30 who had undergone RP, were treated with 3D-CRT and concurrent weekly paclitaxel. Although the trial was not designed to test differences between post-RP and non-post-RP patients, overall grade 3 GI and GU toxicity was more common in post-RP patients, whereas estimates of biochemical and clinical recurrence were similar.

Table 10 Late adverse events recorded after concurrent high-dose intensity-modulated radiation therapy and docetaxel-based chemotherapy: genitourinary symptoms

| Grade | Frequency/urgency | Dysuria | Incontinence | Retention | Hematuria | Highest during follow-up | Highest at last follow-up |
|-------|-------------------|---------|--------------|-----------|-----------|--------------------------|--------------------------|
| L     | L*                | L       | L*           | L         | L         | L                        | L                        |
| Cohort 1 | 2                | 3       | 1            | 1         | 0         | 2                        | 1                        |
| Cohort 2 | 2                | 2       | 1            | 1         | 1         | 2                        | 1                        |

L = all late adverse events; L* = adverse events of higher grade than baseline.
Our study is different because we used a more modern treatment regimen including high-dose IMRT and docetaxel, but was similarly not equipped for this scope, although both functional and oncological outcome were apparently comparable in the two cohorts.

Over the past few years, there has been an increasing interest towards higher-dose taxane-based chemotherapy. In the ongoing phase III Radiation Therapy Oncology Group-0521 trial (ClinicalTrials.gov, NCT00288080), which is evaluating whether adjuvant docetaxel to EBRT with ADT provides a survival advantage over EBRT with ADT alone in high-risk localized PCa, a high dosage of 75 mg/m² every 3 weeks for 6 weeks is used. In our study, an intensified dosage of docetaxel was used compared to previous trials. This was decided in an attempt to further improve the oncological outcome, given the tolerability of the 20 mg/m² dose established in the pivotal trial by Kumar et al. [14]. The above-mentioned limitations notwithstanding, toxicity profile in the present and previous trials is apparently comparable.

Concerning the oncological outcome, our results are comparable to those of the two trials with the longest

Figure 1 Kaplan-Meier estimates of a) biochemical recurrence-free survival and b) clinical recurrence-free survival.
| Study | Design | N patients | EBRT technique | EBRT dosage (Gy) | Taxane drug | Taxane dosage (mg/m²) | ADT duration (months) | Toxicity scoring system | Highest acute GU toxicity § | Highest acute GI toxicity § | Highest late toxicity § | Biochemical recurrence | Follow-up, median (months) |
|-------|--------|------------|----------------|-----------------|-------------|----------------------|----------------------|------------------------|------------------------|------------------------|------------------------|--------------------------|--------------------------|
| Kumar [14] | Phase I | 22 | 3D-CRT | 70.2 | Docetaxel | 5 (n=3), 8 (n=3), 12 (n=3), 16 (n=5), 20 (n=6), 25(n=2) | None | CTC v2.0, RTOG † | Grade 2 | Frequency/urgency | Diarrhea (n=2) | Urinary Retention (n=1) | 5/8 | 8 |
| Sanfilippo [23] | Phase I/II | 22 | 3D-CRT | 63 (n=3), 66.6 (n=7), 70.2 (n=4), 73.8 (n=8) | Paclitaxel | 30 | 9 | CTC v2.0 | Grade 2 | Frequency/urgency | Diarrhea (n=4) | Frequency | Grade 3 | 6/22 | 38 |
| Perrotti [15] | Phase I/II | 20 | IMRT | 72 | Docetaxel | 20 | None | CTC, RTOG † | Grade 2 | Frequency | Diarrhea (n=8) | none | Grade 2 | 3/20 | 11.7 |
| Bolla [16] | Phase II | 50 | 3D-CRT (n=45), IMRT (n=5) | 70 | Docetaxel | 20 | <36 (n=6), 36 (n=43), >36 (n=1) | CTC v2.0, RTOG † | Grade 3 | Dysuria (n=2) | Proctitis (n=1) | Proctitis (n=1) | Grade 3 | 3/20, 76.3, 74.9* |
| Hussain [24] | Phase I | 59 | 3D-CRT | 70.2 (n=29), 64.8 (n=30)* | Paclitaxel | 40 (n=10), 50 (n=31), 60 (n=18) | 4 (n=29), 24 (n=30) | CTC v2.0 | Grade 2 | Frequency/urgency | Diarrhea (n=9) | NA | 13/20, 11/30* |
| Chen [25] | Phase I | 18 | IMRT | 78 | Docetaxel | 10 (n=9), 15 (n=6), 20 (n=3) | 24 | CTCAE v3.0 | Grade 2 | Frequency | Diarrhea (n=2) | NA | 3/18 | 26 |
| Present series | Phase II | 35 | IMRT | 80 (n=17), 70 (n=18)* | Docetaxel | 30 mg (n=8), 40 mg (n=27) | 24 | CTCAE v3.0 | Grade 3 | Grade 2 | Urinary retention (n=1) | Diarrhea | 6/17, 8/18* | 63 |

3D-CRT = three-dimensional conformal radiation therapy; ADT = androgen deprivation therapy; CTC = Common Toxicity Criteria of the National Cancer Institute; CTCAE = Common Terminology Criteria for Adverse Events of the National Cancer Institute; EBRT = external beam radiation therapy; GI = gastrointestinal symptoms; GU = genitourinary symptoms; IMRT = intensity-modulated radiation therapy; NA = not assessed; NR = not reported; RTOG = Radiation Therapy Oncology Group/European Organization for Research and Treatment of Cancer toxicity criteria. *Patients with previous radical prostatectomy; † late toxicity; ‡ clinical disease-free survival was 66.72% at 5 years; § when two or more events, only the most common was reported.
Finally, irradiation to pelvic lymph nodes in addition to similar randomized trial [30], surgical margin status and clinical recurrence remains under debate. In another, the impact of positive surgical margins on biochemical outcomes were not available. However, even currently, vant EBRT in subgroup analyses. Our study started in following RP for high-risk PCa [28] showed that patients with positive surgical margins benefit most from adjuvant EBRT with ADT alone, with appropriate oncological outcomes as the primary endpoint. In fact, our treatment paradigm might prove beneficial if hard endpoints (e.g. metastasis-free and cancer-free survival) and a sufficiently long-term follow-up are considered. On the other hand, a recent study on ultra-high-dose (86.4 Gy) IMRT for localized PCa [29] has reported higher BRFS rates (67.9%) in high-risk patients at a similar follow-up (5.5 years). One may speculate that IMRT delivered at this very high dose can achieve a better improvement, at least, in local control, than docetaxel-based chemo-radiation. However, whether concurrent docetaxel confers a true benefit has to be seen in the pattern of clinical recurrence, where the incidence of regional and distant metastases is expected to be lower. Future research in this area is eagerly awaited.

To the best of our knowledge, two phase II/III trials and one phase III trial testing EBRT with ADT plus concurrent docetaxel vs EBRT with ADT alone in high-risk PCa are currently recruiting patients (ClinicalTrials.gov, NCT00116142, NCT01603420 and NCT01811810). However, in none of these trials the experimental arm entails a combination of high-dose IMRT, intensified-dosage docetaxel-based chemotherapy and long-term ADT, such as in our trial.

Our study is not devoid of limitations. First, no control group was available. Second, toxicity was graded according to a physician-based assessment and health-related quality of life was not assessed. Third, IGRT may have contributed to improve the toxicity profile. Fourth, the choice between RP and RT was based on an arbitrary age cut-off with its inherent limitations. Fifth, as per our trial protocol post-RP EBRT was given to patients with either extracapsular disease, seminal vesicle invasion or positive surgical margins, which are all established risk factors for local recurrence. The latest results of a large randomized trial on adjuvant EBRT vs observation following RP for high-risk PCa [28] showed that patients with positive surgical margins benefit most from adjuvant EBRT in subgroup analyses. Our study started in 2005, i.e. at a time when mature data on post-RP EBRT outcomes were not available. However, even currently, the impact of positive surgical margins on biochemical and clinical recurrence remains under debate. In another similar randomized trial [30], surgical margin status was not an independent predictor of biochemical outcome. Finally, irradiation to pelvic lymph nodes in addition to the prostate in cN0 high-risk PCa, as in our cohort 2 patients, remains controversial. One option would be a staging PLND, whereby pN0 patients might be spared irradiation and pN1 patients might benefit from irradiation with long-term ADT [5].

Conclusions
In a phase II trial including patients with high-risk localized or locally advanced PCa we showed acceptably low toxicity and good efficacy of a multimodal treatment including high-dose IMRT preceded or not by RP, concurrent intensified-dose docetaxel and long-term ADT. Thus, this treatment paradigm may warrant further evaluation in phase III randomized trials.

Abbreviations
3D-CRT: Three-dimensional conformal radiation therapy; ADT: Androgen deprivation therapy; BRFS: Biochemical recurrence-free survival; CRFS: Clinical recurrence-free survival; CTV: Clinical target volume; EBRT: External beam radiation therapy; GI: Gastro-intestinal; GU: Genito-urinary; IGRT: Image-guided radiation therapy; IMRT: Intensity-modulated radiation therapy; IQR: Interquartile range; PCa: Prostate cancer; PLND: Pelvic lymph node dissection; PTV: Planning target volume; RP: Radical prostatectomy.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
RB and Fiz: study conception and design. AGu, FaZ, MG and VP: acquisition of data. AGu, GG, PG, AGa, RB and Fiz: analysis and interpretation of data. AGu, GG and PG: drafting of the manuscript. AGu and Fiz: surgery. RB: radiation therapy planning. FV: statistical analyses. RB and Fiz: critical revision of the manuscript for important intellectual content, and supervision. All authors read and approved the final manuscript.

Acknowledgement
This work was presented in part at the 27th European Association of Urology Annual Congress, 24–28 February 2012, Paris, and at the 54th American Society for Radiation Oncology Annual Meeting, 28–31 October 2012, Boston, MA, USA.

Author details
1Department of Oncological, Surgical and Gastroenterological Sciences, Urology Clinic, University of Padua, Padua 35128, Italy. 2Department of Radiation Oncology, National Cancer Institute Aviano, Aviano, PN, Italy. 3Department of Urology, University of Bern, Inselspital, Bern, Switzerland. 4Department of Radiation Oncology, University of Bern, Inselspital, Bern, Switzerland. 5Department of Epidemiology, Service, Direzione Centrale Salute, Integrazione Socioassistenza e Politiche Sociali, Regione Autonoma Friuli Venezia Giulia, Udine, Italy. 6Department of General Surgery, Division of Urology, Azienda Ospedaliera “Santa Maria degli Angeli”, Pordenone, Italy.

Received: 25 September 2013 Accepted: 30 December 2013 Published: 14 January 2014

References
1. Yossepowitch O, Eggener SE, Serio AM, Carver BS, Bianco FJ Jr, Scardino PT, Eastham JA: Secondary therapy, metastatic progression, and cancer-specific mortality in men with clinically high-risk prostate cancer treated with radical prostatectomy. Eur Urol 2008, 53:950–959.
2. Cooperberg MR, Broering JM, Carroll PR: Time trends and local variation in primary treatment of localized prostate cancer. J Clin Oncol 2010, 28:1117–1123.
3. Horwitz EM, Bae K, Hanks GE, Porter A, Grignon DJ, Breieron HD, Venkatesan V, Lawton CA, Rosenthal SA, Sandler HM, Shipley WU: Ten-year follow-up of radiation therapy oncology group protocol 92–02: a phase III trial of
