Cytoreductive Radiotherapy Combined with Abiraterone in Metastatic Castration-Resistance Prostate Cancer: A Single Center Experience

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Research

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Abstract

**Background:** To investigate the potential benefit of cytoreductive radiotherapy (cRT) in metastatic castration-resistant prostate cancer (mCRPC) patients receiving abiraterone.

**Methods:** From February 2014 to February 2019, 149 mCRPC patients treated with abiraterone were identified. Patients receiving cRT before abiraterone failure (AbiRT group) were matched by one-to-two propensity score to patients without cRT before abiraterone failure (non-AbiRT group).

**Results:** The median follow-up was 23.5 months. Thirty patients (20.1%) were in the AbiRT group, whereas 119 patients (79.9%) were in the non-AbiRT group. The 2-year overall survival (OS) of patients managed by AbiRT, cRT after abiraterone failure, and no cRT was 89.5%, 72.0% and 72.0%, respectively ($P = 0.001$). On multivariate analysis, only AbiRT (HR, 0.17; 95%CI, 0.05–0.58; $P = 0.004$) and prognostic index (HR 2.71; 95% CI, 1.37–5.35; $P = 0.004$) were significant factors. After matching, AbiRT continued to be associated with improved OS (median OS not reached vs. 44.0 months, $P = 0.009$). Subgroup analysis revealed that patients aged $\leq$ 65 years (HR, 0.09; 95%CI, 0.01–0.65; $P = 0.018$), PSA $\leq$ 20 ng/mL (HR 0.29; 95% CI, 0.09–0.99; $P = 0.048$), chemotherapy-naïve upon abiraterone treatment (HR 0.20; 95% CI, 0.06–0.66; $P = 0.008$) and in intermediate prognosis groups (HR 0.13; 95% CI, 0.03-0.57; $P = 0.007$) had improved OS with AbiRT.

**Conclusions:** cRT before resistance to abiraterone may improve survival in selected mCRPC patients: relatively young, chemotherapy-naïve, with a relatively low PSA level at the diagnosis of mCRPC.

**Background**

Prostate cancer is the second most commonly diagnosed cancer in men worldwide [1]. In China, prostate cancer used to be a rare disease, but a sustained rapid increase of incidence since 2000 has made prostate cancer the sixth most common cancer and the tenth leading cause of death in the Chinese men [2]. The proportion of early-stage and localized prostate cancer is rather low in China, and the estimated 5-year overall survival (OS) is only 66.4% [2]. Many Chinese patients present with metastatic disease at diagnosis, and prolonging the survival of patients with metastatic prostate cancer has become a great challenge in general practice.

Nearly all metastatic prostate cancer will progress into an aggressive state known as metastatic castration-resistant prostate cancer (mCRPC). Without effective treatment, the median OS of these patients is only 9–30 months [3]. In recent years, major advances in therapeutic agents has significantly improved the survival of mCRPC patients. Abiraterone is one of the standard of care for mCRPC. Abiraterone plus prednisone has shown remarkable efficacy and safety in chemotherapy-naïve and chemotherapy-treated mCRPC patients [4, 5]. However, heterogenous responses to abiraterone exist due to the polyclonal nature of metastatic sites, and acquired resistance to abiraterone eventually develop after 6–14 months [4, 5]. Thus, increasing studies are investigating the potential benefit of adding local therapy to systemic therapy in metastatic prostate cancer.
Current cytoreductive treatment strategies for metastatic prostate cancer includes prostate-directed therapy and metastasis-directed therapy. The STAMPEDE trial and the HORRAD trial demonstrates that prostate-directed radiotherapy could prolong survival in metastatic hormone-sensitive prostate cancer (mHSPC) with low metastatic burden [6, 7]. Beyond local cytoreduction of primary sites, reducing the metastatic burden through local therapy may also bring survival benefit, as observed in ovarian, kidney and some gastrointestinal cancers [8–10]. In prostate cancer, metastasis-directed radiotherapy is increasingly endorsed in patients with limited metastatic lesions early in their chain of progression. Stereotactic body radiotherapy (SBRT) to all metastatic sites is likely to delay disease progression and prolong systemic in oligometastatic mHSPC, as supported by the results of some phase II trials such as STOMP, ORIOLE and POPSTAR [11–13].

Given the promising results of cytoreductive radiotherapy in mHSPC, we hypnotized that cytoreductive radiotherapy might as well have a role in the event of mCRPC, a more terminal state. In this retrospective study, we sought to investigate the potential benefit of cytoreductive radiotherapy (cRT) in mCRPC patients treated with abiraterone.

Methods

Patient selection and baseline evaluation

This study was approved by the institutional review board. We retrospectively reviewed 320 prostate cancer patients treated with abiraterone plus prednisone between February 2014 and February 2019 in our institution. Inclusion criteria were mCRPC patients receiving abiraterone with or without radiotherapy. Patients were excluded if they were HSPC or nonmetastatic at the time of abiraterone treatment. Patients lost to follow-up less than 3 months after abiraterone treatment, who discontinued abiraterone treatment for nancial burden, or who lacked data for risk stratification were also excluded. Finally, a total of 149 patients were included in the analyses (Fig. 1).

Oligometastasis was de ned as \( \leq 5 \) metastatic lesions to lymph nodes and/or bones without visceral metastasis [14]. Risk stratification was determined by a prognostic index developed from the COU-AA-301 study [15], which has been validated in both chemotherapy-naive and chemotherapy-treated mCRPC patients [16, 17]. This model comprises six risk factors: lactate dehydrogenase > upper limit of normal (ULN); Eastern Cooperative Oncology Group performance status of 2; liver metastases; albumin \( \leq 4 \) g/dL; alkaline phosphatase > ULN; time from start of initial androgen-deprivation therapy (ADT) to treatment initiation is \( \leq 36 \) months. Patients were categorized into good (0–1 risk factor), intermediate (2–3 risk factors) and poor (4–6 risk factors) prognostic groups.

Treatment

Patients treated with cRT before abiraterone failure were placed in the AbiRT group. The comparison group was patients who did not receive cRT before abiraterone (non-AbiRT group). cRT was defined as
ablative radiotherapy to lesions that accounted for > 50% of the total tumor burden, which was the sum of the longest unidimensional diameter of target lesions according to Response Evaluation and Criteria in Solid Tumors v1.1. All patients had ADT together with 1000 mg of abiraterone once-daily and 5 mg of prednisone twice-daily. Abiraterone was not withheld or its dose was not reduced during radiotherapy.

Primary sites were treated with intensity-modulated radiotherapy, and all metastatic sites were treated with SBRT. Patients underwent simulation with contrast-enhanced CT (slice thickness = 3 mm) with site-specific immobilization. Planning CT images were fused with magnetic resonance images of the skeletal segment interested. Contouring was in accordance with the corresponding recommendations set by the Radiation Therapy Oncology Group (RTOG). For the prostate, the clinical target volume (CTV) included the prostate gland with or without seminal vesicles, and the planning target volume (PTV) was yielded by expansion by 5 mm (3 mm posteriorly). Regional lymph nodes were not contoured unless radiographically positive. Distant metastases were treated with SBRT. The CTV equals the gross tumor volume. The PTV was defined as the CTV plus a margin of ≤ 5 mm.

The prescription dose for the prostate gland and pelvic lymph nodes was 60–67.5 Gy and 45–60 Gy in 25 fractions, respectively. The dosage regimen for metastatic lesions was 18–35 Gy in 1–5 fractions. Dose constraints for normal tissue were in accordance with RTOG guidelines. Treatment was delivered by a linear accelerator using 6–8 MV photons. Image guided radiation therapy using cone-beam CT was performed before each treatment.

Outcomes

For patients treated with radiotherapy, the PSA tests were carried out 1 month after radiotherapy, and every 3 months thereafter. For patients treated with abiraterone alone, PSA was generally tested every 3 months. Radiological evaluation was ordered at the discretion of physicians. OS was defined as the time from the diagnosis of mCRPC until last follow-up or death from any cause. Progression-free survival (PFS) was measured from the start of radiotherapy until PSA or radiographic progression or death. Biochemical and radiographic progression was assessed according to the Prostate Cancer Clinical Trials Working Group 2 (PCWG2). PSA response was defined as a decline in PSA levels of > 50% from baseline, measured twice 4 weeks apart.

Statistical analysis

To address the imbalance of potential confounders, we used propensity score matching to compare the OS between the AbiRT and non-AbiRT group. The propensity score was estimated as the probability of receiving AbiRT from a logistic regression model. Variables that were prognostic for OS were explored. The propensity-score model included age, Gleason score, PSA level at mCRPC, prognostic group, synchronous metastasis, oligometastasis, chemotherapy-naïve upon abiraterone treatment. One-to-two matching without replacement was implemented using nearest-neighbor matching. The caliper width was 0.2 times the standard deviation of the logit of the propensity score.
Categorical data were compared using the chi-squared test. OS and PFS were estimated using the Kaplan–Meier method and compared using the log-rank test. The Cox proportional hazard model was used for univariate and multivariate analyses. Variables that were significant in the univariate analysis or clinically relevant were included in the multivariate analysis. A \( P \) value of less than 0.05 was considered significant. All tests were two-sided. Descriptive analysis and survival analyses were carried out by SPSS v23 (IBM, Armonk, NY, USA). Matching of propensity scores was done by Python (www.Python.org).

Results

The baseline characteristics of the entire cohort are summarized in Table 1. The median PSA level at the time of mCRPC diagnosis was 16.8 ng/mL (range 0.2–3411.0 ng/mL). Seventy-nine patients (53.0%) had a Gleason score of 9 or 10. Synchronous metastasis was present in 127 patients (85.2%). Fifty-eight patients (38.9%) had oligometastasis at the time of mCRPC diagnosis. The number of patients allocated to good, intermediate and poor prognostic groups was 65 (43.6%), 75 (50.3%) and 9 (6.0%), respectively. Fifty-four patients received docetaxel (65–75 mg/m\(^2\)) every 3 weeks, with 26 cases (17.4%) before abiraterone treatment. Forty patients (26.8%) underwent cRT, 30 (75.0%) of whom were irradiated before abiraterone failure. The remaining 10 patients (25.0%) received cRT with concurrent secondary hormone therapies after abiraterone failure.
Table 1
Baseline characteristics of the entire cohort (N = 149)

| Characteristics                | No. (%) |
|--------------------------------|---------|
| Age, median (range), years     | 68 (45–86) |
| PSA at mCRPC                   |         |
| ≤ 20 ng/mL                     | 80 (53.7) |
| > 20 ng/mL                     | 69 (46.3) |
| Gleason score                  |         |
| ≤ 8                            | 70 (47.0) |
| 9–10                           | 79 (53.0) |
| ECOG                           |         |
| 0–1                            | 105 (70.5) |
| > 1                            | 44 (29.5) |
| Prognostic index               |         |
| Good                           | 65 (43.6) |
| Intermediate                   | 75 (50.3) |
| Poor                           | 9 (6.0) |
| Synchronous metastasis         | 127 (85.2) |
| Oligometastasis                | 58 (38.9) |
| Chemo-naïve                    | 123 (82.6) |

PSA prostate-specific antigen, mCRPC metastatic castration-resistant prostate cancer, Chemo-naïve chemotherapy-naïve upon abiraterone treatment

Thirty patients (20.1%) were in the AbiRT group, whereas 119 patients (79.9%) were in the non-AbiRT group. Compared with the non-AbiRT group, patients in the AbiRT group were more likely to have oligometastasis (P = 0.002), with a lower PSA level at mCRPC diagnosis (P = 0.005). Other baseline characteristics including age, Gleason score, synchronous metastasis, chemotherapy-naïve upon abiraterone treatment and prognostic index were similar (Table 2).
Table 2
Comparison of baseline characteristics in the unmatched and the matched data

| Characteristics          | Unmatched data | Matched data |  |
|-------------------------|----------------|--------------|---------------|
|                         | No. (%)        | No. (%)      | *P*           |
| AbiRT (N = 30)          | Non-AbiRT (N = 119) |                |               |
| Age, years              |                |              | 0.224         |
| ≤ 65                    | 15 (50.0)      | 45 (37.8)    | 0.440         |
| > 65                    | 15 (50.0)      | 74 (62.2)    |               |
| PSA at mCRPC            |                |              | 0.005         |
| ≤ 20 ng/mL              | 23 (76.6)      | 57 (47.9)    | 0.667         |
| > 20 ng/mL              | 7 (23.3)       | 62 (52.1)    |               |
| Gleason score           |                |              | 0.435         |
| ≤ 8                     | 16 (53.3)      | 54 (45.4)    | 0.886         |
| 9–10                    | 14 (46.6)      | 65 (54.6)    |               |
| Prognostic index        |                |              | 0.107         |
| Good                    | 17 (56.6)      | 48 (40.3)    | 0.227         |
| Intermediate/Poor       | 13 (43.3)      | 71 (59.7)    |               |
| Oligometastasis         | 19 (63.3)      | 39 (32.8)    | 0.002         |
| Synchronous metastasis  | 24 (80.0)      | 103 (86.6)   | 0.908         |
| Chemo-naïve             | 28 (93.3)      | 95 (79.8)    |               |

*PSA prostate-specific antigen, mCRPC metastatic castration-resistant prostate cancer, Chemo-naïve chemotherapy-naïve upon abiraterone treatment*

At a median follow-up of 23.5 months, 54 patients (36.2%) died. Seven patients (4.7%) were lost to follow-up. The median OS of the entire cohort was 38.4 months. The median OS of patients undergoing cRT was not reached, compared with 31.4 months in patients who did not have cRT (*P* = 0.001). The 2-year OS rates of patients managed by AbiRT, cRT after abiraterone failure, and no cRT was 89.5%, 72.0% and 72.0%, respectively (*P* = 0.001) (Fig. 2). The median PFS following radiotherapy in the AbiRT group was 12.2 months, and 23 (76.6%) patients had a PSA response after radiotherapy. The median OS of the AbiRT group was not reached whereas, in the non-AbiRT group, the median OS was 31.8 months (*P* = 0.000). The 2-year OS rates of the AbiRT group and non-AbiRT group were 89.5% and 73.5%, respectively (*P* = 0.000) (Fig. 3). Upon univariate analysis, AbiRT, oligometastasis, intermediate/poor group according
to the prognostic index, PSA > 20 ng/mL and chemotherapy-naïve upon abiraterone treatment were significant prognostic factors for OS (Table 3). Upon multivariate analysis, the AbiRT group [hazard ratio (HR), 0.17; 95% confidence interval (CI), 0.05–0.58; \( P = 0.004 \)] and intermediate/poor grouping for the prognostic index (HR 2.71; 95% CI, 1.37–5.35; \( P = 0.004 \)) were significant prognostic factors (Table 3).

### Table 3
Univariate and multivariable analyses of factors predictive of overall survival

| Variables                  | Univariate analysis | Multivariate analysis |
|----------------------------|---------------------|-----------------------|
|                            | HR (95% CI)         | \( P \)               | HR (95% CI)         | \( P \)               |
| AbiRT                      |                     |                       |
| Yes vs No                  | 0.15 (0.05, 0.48)   | 0.001                 | 0.17 (0.05, 0.58)   | 0.004                 |
| Oligometastasis            |                     |                       |
| Yes vs No                  | 0.52 (0.30, 0.93)   | 0.028                 | 0.75 (0.39, 1.44)   | 0.387                 |
| PSA > 20 ng/mL             |                     |                       |
| Yes vs No                  | 2.16 (1.23, 3.79)   | 0.007                 | 1.66 (0.93, 2.96)   | 0.089                 |
| Prognostic index           |                     |                       |
| Interm/poor vs favorable   | 3.11 (1.64, 5.92)   | 0.001                 | 2.71 (1.37, 5.35)   | 0.004                 |
| Chemo-naïve at Abi         |                     |                       |
| Yes vs No                  | 0.45 (0.25, 0.81)   | 0.008                 | 0.81 (0.41, 1.61)   | 0.548                 |

*PSA* prostate-specific antigen, *Interm* intermediate, *Chemo-naïve* chemotherapy-naïve upon abiraterone treatment, *HR* hazard ratio, *CI* confidential interval

After propensity score matching with a caliper of 0.21, 30 patients in the AbiRT group were matched to 58 patients in the non-AbiRT group. The difference between the baseline characteristics was not significant after matching (Table 2). The survival advantage of AbiRT remained significant. The median OS was not reached in the AbiRT group, compared with 44.0 months in the non-AbiRT group (\( P = 0.009 \)). The 2-year OS of the AbiRT group and non-AbiRT group was 89.5% and 79.3%, respectively (Fig. 3).

The AbiRT group was associated with improved OS in the subgroups of age \( \leq 65 \) years old, PSA \( \leq 20 \) ng/mL and chemotherapy-naïve upon abiraterone treatment (Fig. 4). The benefit of AbiRT was not evaluated in patients with poor prognosis according to the prognostic index given that there were only nine patients in this subgroup. In the subgroup with an intermediate prognosis according to the prognostic index, application of AbiRT was associated with a reduction of death of 85% (HR, 0.15; 95%CI, 0.03–0.63; \( P = 0.010 \)); this advantage was not observed in patients with a good prognosis (0.23; 0.03–1.79; 0.161).
Radiotherapy was well tolerated, with no grade 3 or higher toxicities reported. Acute side effects included gastrointestinal (GI) toxicity (9 patients), genitourinary (GU) toxicity (11 patients), thrombocytopenia (1 patient), neutropenia (1 patient) and hypokalemia (1 patient). Late GI and GU toxicity were observed in 7 and 6 patients, respectively. Two patients developed vertebral compression fractures, and neither of them were symptomatic.

**Discussion**

Evolving novel hormonal agents have improved the systemic control and prolonged survival of mCRPC patients, and increasing focus is being placed on the potential benefits of cytoreductive local therapy [18]. This study provides valuable insights regarding the value of cytoreductive radiotherapy in mCRPC. In this comparison of survival outcomes between the two groups, cRT before abiraterone failure was associated with a remarkable improvement in OS, and adoption of cRT required careful consideration when systemic control could no longer be maintained by abiraterone therapy. Patients aged ≤ 65 years, with PSA ≤ 20 ng/mL and who were chemotherapy-naïve may be potential candidates for cytoreductive therapies.

Addition of radiotherapy to abiraterone, a novel androgen receptor axis-targeted therapy (ARAT), may provide additional advantages given the potential synergistic interaction between these two treatment modalities [19]. Experiments suggest that androgen-receptor signaling can increase the expression of DNA repair genes and promote radioresistance by accelerating repair of the DNA damage induced by ionizing radiation [20]. New-generation ARAT can result in downregulation of expression of the DNA repair genes, thereby potentiating the effect of ionizing radiation [21]. In clinical studies, a delay of disease progression has been observed after delivering radiotherapy to mCRPC patients treated with abiraterone [22–24]. Similarly, our study showed that the survival benefit was most pronounced if cRT was combined with abiraterone. Ongoing clinical trials (NCT03449719 and NCT03556904) will help to decide whether combining radiotherapy and abiraterone could provide additional advantage in mCRPC.

cRT can discontinue direct seeding of new metastases as well as stopping supportive interactions between primary and metastatic sites [25, 26]. cRT can also eliminate resistant clones and release a wider range of tumor antigens [27]. Emerging clinical evidence suggests that cRT can significantly prolong PFS and even OS in mHSPC [6, 11–13]. In contrast, cRT for mCRPC seems less attractive because the value of local therapy at such a late stage is questionable. Several retrospective studies show that radiotherapy to oligoprogressive sites could delay disease progression in mCRPC [22, 24, 28]. In the study by Yildirim et al., prolonged abiraterone use was also observed in mCRPC patients receiving prostate-directed therapy, but no significant improvement of OS was found (24.1 vs. 21.4 months; P = 0.08) [23]. As half of the patients in this study were previously treated with chemotherapy, we speculate that patients are more likely to have survival benefit when cytoreductive local therapy is adopted at an early stage of mCRPC [14]. In our study, patients undergoing cRT before abiraterone failure enjoyed a remarkable improvement in OS, while the OS of patients receiving cRT after abiraterone failure remained poor even if cytoreduction had been applied. These results imply that the power of cytoreductive local therapy is restricted at a terminal stage of disease, especially if multiple lines of treatment have failed.
In subgroup analyses, the survival benefit from AbiRT was observed in patients in the intermediate prognostic group instead of in those in the good prognostic group. These data suggested that abiraterone treatment alone could elicit satisfactory control for some low-risk patients, whereas intensive therapy (e.g., AbiRT) was worth trying for intermediate-risk patients because abiraterone treatment alone might be not sufficient for these patients. Interestingly, oligometastasis could not be used to identify potential candidates for local radiotherapy in our study. Current definition of oligometastasis only offers an assessment of tumor burden in a snapshot, but the underlying clinical pathways that lead to an oligometastatic state is undefined. The European Society for Radiotherapy and Oncology and European Organisation for Research and Treatment of Cancer has classified the oligometastases into 9 distinct situations \[29\]. Oligometastasis can be induced by multiple lines of systemic therapies in mCRPC, and patients who are heavily pretreated are not in their early chain of progression despite having limited metastases. These data suggest that oligometastasis, a generally accepted indication for local cytoreductive therapy in newly diagnosed mHSPC, may not be applicable in mCRPC. Thus, the decision regarding local intervention requires combined interpretation of the patient’s general condition, disease state and the profile of systemic treatment \[14\]. Patients who are not heavily-pretreated, and those who are relatively young with good treatment tolerance as well as a low level of PSA, may benefit from aggressive local therapy in mCRPC.

Our study has several limitations. First, it is a retrospective study. However, given the lack of evidence on the role of cytoreductive radiotherapy in the general situation other than oligometastasis and oligoprogession, our study provides valuable information for current practice. Second, excluding patients with missing data might have led to selection bias (though there is no evidence that these patients were more or less prone to be omitted based on the scarce information in the medical records). Third, our patients represent a heterogenous cohort of mCRPC patients who received different treatment at different time points.

**Conclusions**

The present study evaluated the survival outcomes of cytoreductive radiotherapy in mCRPC patients treated with abiraterone. The findings from our study support the use of cytoreductive radiotherapy before the resistance to abiraterone in selected mCRPC patients, preferably in relatively young and chemotherapy-naïve patients with a relatively low PSA at the time of mCRPC.

**List Of Abbreviations**

OS: overall survival  
mCRPC: metastatic castration-resistant prostate cancer  
mHSPC: metastatic hormone-sensitive prostate cancer  
ADT: androgen-deprivation therapy
Declarations

Ethics approval and consent to participate

This project was approved by the Ethical Committee of Sun Yat-Sen University Cancer Center and informed consent was waived by the committee.

Consent for publication

Not applicable.

Availability of data and materials

The datasets used and/or analysed during the current study are available from the corresponding author on reasonable request.

Competing interests

The authors declare that they have no competing interests.
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Authors' contributions

LY and LW participated in study design, statistical analysis and manuscript drafting. DP collected the clinical data. ZEZ, MLX, HSJ, LBJ, CWF, WJH and ZFJ contributed to the operation work during treatment. LYH and HLR designed the study, reviewed and revised the manuscript. All authors read and approved the final manuscript.

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References

1. Bray F, Ferlay J, Soerjomataram I, Siegel RL, Torre LA, Jemal A. Global cancer statistics 2018: GLOBOCAN estimates of incidence and mortality worldwide for 36 cancers in 185 countries. CA Cancer J Clin. 2018;68(6):394-424.
2. Zheng R, Sun K, Zhang S, Zeng H, Zou X, Chen R. Report of cancer epidemiology in China 2015. Chin J Oncol. 2019;41(1):19-28.
3. Kirby M, Hirst C, Crawford ED. Characterising the castration-resistant prostate cancer population: a systematic review. Int J Clin Pract. 2011;65(11):1180-92.
4. Ryan CJ, Smith MR, Fizazi K, Saad F, Mulders PF, Sternberg CN, et al. Abiraterone acetate plus prednisone versus placebo plus prednisone in chemotherapy-naive men with metastatic castration-resistant prostate cancer (COU-AA-302): final overall survival analysis of a randomised, double-blind, placebo-controlled phase 3 study. Lancet Oncol. 2015;16(2):152-60.
5. Fizazi K, Scher HI, Molina A, Logothetis CJ, Chi KN, Jones RJ, et al. Abiraterone acetate for treatment of metastatic castration-resistant prostate cancer: final overall survival analysis of the COU-AA-301 randomised, double-blind, placebo-controlled phase 3 study. Lancet Oncol. 2012;13(10):983-92.
6. Parker CC, James ND, Brawley CD, Clarke NW, Hoyle AP, Ali A, et al. Radiotherapy to the primary tumour for newly diagnosed, metastatic prostate cancer (STAMPEDE): a randomised controlled phase 3 trial. Lancet. 2018;392(10162):2353-66.
7. Boeve LMS, Hulshof M, Vis AN, Zwinderman AH, Twisk JWR, Witjes WPJ, et al. Effect on Survival of Androgen Deprivation Therapy Alone Compared to Androgen Deprivation Therapy Combined with Concurrent Radiation Therapy to the Prostate in Patients with Primary Bone Metastatic Prostate Cancer.
Cancer in a Prospective Randomised Clinical Trial: Data from the HORRAD Trial. Eur Urol. 2019;75(3):410-8.

8. Dabestani S, Marconi L, Hofmann F, Stewart F, Lam TB, Canfield SE, et al. Local treatments for metastases of renal cell carcinoma: a systematic review. Lancet Oncol. 2014;15(12):e549-61.

9. Bristow RE, Tomacruz RS, Armstrong DK, Trimble EL, Montz FJ. Survival effect of maximal cytoreductive surgery for advanced ovarian carcinoma during the platinum era: a meta-analysis. J Clin Oncol. 2002;20(5):1248-59.

10. Glehen O, Mohamed F, Gilly FN. Peritoneal carcinomatosis from digestive tract cancer: new management by cytoreductive surgery and intraperitoneal chemohyperthermia. Lancet Oncol. 2004;5(4):219-28.

11. Ost P, Reynders D, Decaestecker K, Fonteyne V, Lumen N, De Bruycker A, et al. Surveillance or Metastasis-Directed Therapy for Oligometastatic Prostate Cancer Recurrence: A Prospective, Randomized, Multicenter Phase II Trial. J Clin Oncol. 2018;36(5):446-53.

12. Radwan N, Phillips R, Ross A, Rowe SP, Gorin MA, Antonarakis ES, et al. A phase II randomized trial of Observation versus stereotactic ablative Radiation for OLigometastatic prostate CancEr (ORIOLE). BMC Cancer. 2017;17(1):453.

13. Siva S, Bressel M, Murphy DG, Shaw M, Chander S, Violet J, et al. Stereotactic Abative Body Radiotherapy (SABR) for Oligometastatic Prostate Cancer: A Prospective Clinical Trial. Eur Urol. 2018;74(4):455-62.

14. Wei XX, Ko EC, Ryan CJ. Treatment strategies in low-volume metastatic castration-resistant prostate cancer. Curr Opin Urol. 2017;27(6):596-603.

15. Chi KN, Kheoh T, Ryan CJ, Molina A, Bellmunt J, Vogelzang NJ, et al. A prognostic index model for predicting overall survival in patients with metastatic castration-resistant prostate cancer treated with abiraterone acetate after docetaxel. Ann Oncol. 2016;27(3):454-60.

16. Khalaf DJ, Avilés CM, Azad AA, Sunderland K, Todenhöfer T, Eigl BJ, et al. A prognostic model for stratifying clinical outcomes in chemotherapy-naive metastatic castration-resistant prostate cancer patients treated with abiraterone acetate. Can Urol Assoc J. 2018;12(2):E47-e52.

17. Azad A, Lester R, Leibowitz-Amit R, Joshua AM, Heng DYC, Eigl BJ, et al. Population-based analysis of a novel prognostic model for metastatic castration-resistant prostate cancer (mCRPC) patients (pts) treated with abiraterone acetate (AA). J Clin Oncol. 2014;32(4_suppl):29-.

18. Connor MJ, Shah TT, Horan G, Bevan CL, Winkler M, Ahmed HU. Cytoreductive treatment strategies for de novo metastatic prostate cancer. Nat Rev Clin Oncol. 2019.

19. Livi L, Detti B, Francolini G, Terziani F, Triggiani L, D'Angelillo RM, et al. Combining abiraterone and radiotherapy in metastatic castration-resistant prostate cancer: a review of current evidence. Tumori. 2019;105(4):277-81.

20. Spratt DE, Evans MJ, Davis BJ, Doran MG, Lee MX, Shah N, et al. Androgen Receptor Upregulation Mediates Radioresistance after Ionizing Radiation. Cancer Res. 2015;75(22):4688-96.
21. Polkinghorn WR, Parker JS, Lee MX, Kass EM, Spratt DE, Iaquinta PJ, et al. Androgen receptor signaling regulates DNA repair in prostate cancers. Cancer Discov. 2013;3(11):1245-53.

22. Detti B, D'Angelillo RM, Ingrosso G, Olmetto E, Francolini G, Triggiani L, et al. Combining Abiraterone and Radiotherapy in Prostate Cancer Patients Who Progressed During Abiraterone Therapy. Anticancer Res. 2017;37(7):3717-22.

23. Yildirim BA, Onal C, Kose F, Oymak E, Sedef AM, Besen AA, et al. Outcome of loco-regional radiotherapy in metastatic castration-resistant prostate cancer patients treated with abiraterone acetate. Strahlenther Onkol. 2019;195(10):872-81.

24. Valeriani M, Marinelli L, Macrini S, Reverberi C, Aschelter AM, De Sanctis V, et al. Radiotherapy in metastatic castration resistant prostate cancer patients with oligo-progression during abiraterone-enzalutamide treatment: a mono-institutional experience. Radiat Oncol. 2019;14(1):205.

25. Giri D, Ozen M, Ittmann M. Interleukin-6 is an autocrine growth factor in human prostate cancer. Am J Pathol. 2001;159(6):2159-65.

26. Sehgal I, Powers S, Huntley B, Powis G, Pittelkow M, Maihle NJ. Neurotensin is an autocrine trophic factor stimulated by androgen withdrawal in human prostate cancer. Proc Natl Acad Sci U S A. 1994;91(11):4673-7.

27. Brooks ED, Chang JY. Time to abandon single-site irradiation for inducing abscopal effects. Nat Rev Clin Oncol. 2019;16(2):123-35.

28. Berghen C, Joniau S, Ost P, Poels K, Everaerts W, Decaestecker K, et al. Progression-directed Therapy for Oligoprogession in Castration-refractory Prostate Cancer. Eur Urol Oncol. 2019.

29. Guckenberger M, Lievens Y, Bouma AB, Collette L, Dekker A, deSouza NM, et al. Characterisation and classification of oligometastatic disease: a European Society for Radiotherapy and Oncology and European Organisation for Research and Treatment of Cancer consensus recommendation. Lancet Oncol. 2020;21(1):e18-e28.

**Figures**
Figure 1

Study flow diagram.
Figure 2

(A) Overall survival for patients with mCRPC treated with and without cytoreductive radiotherapy. (B) Overall survival of patients treated with cytoreductive RT before abiraterone failure (AbiRT), after abiraterone failure together with other secondary hormone therapies (cRT+non-Abi) and no cytoreductive RT (non-cRT).

Figure 3

Overall survival for mCRPC patients treated with (AbiRT) and without (non-AbiRT) cytoreductive RT before abiraterone failure before (A) and after (B) propensity score matching.
Figure 4

Forest plot of the association between cytoreductive radiotherapy before abiraterone failure and overall survival by subgroup. HR, hazard ratio; 95% CI, 95% confidence interval.