Docetaxel Based Treatment for Metastatic Castration-Resistant Prostate Cancer-our Early Experience

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Abstract

Aim: Patients with metastatic prostate cancer invariably progress after primary androgen ablation and develop castration-resistant disease after a median time of 18-24 months. We report our early experience with docetaxel based chemotherapy in patients with mCRPC.

Methods: Patients with rising serum Prostate-specific antigen, increasing extent of disease on bone scans and physical findings. All eligible patients with no previous treatment with any cytotoxic drug and having a good ≥80 Karnofsky performance status were included into the study.

Results: 85 patients with metastatic prostate cancer presented with progressive disease following ADT. 67 (78.82%) patients with a mean age of 67.05 ± 2.11 accepted further treatment with docetaxel based chemotherapy, whereas the remaining 18 (21.18%) patients with mean age of 68.0 ± 4.47 years did not undergo further treatment. The overall survival was 32.61 ± 6.09 months in patients receiving docetaxel based chemotherapy as compared to 12.83 ± 2.40 months in patients in the no treatment group. The Serum PSA values at 3 and 9 months after initiation of treatment showed a downward trend in all the three risk groups receiving docetaxel.

Conclusions: In our study the overall survival was 32.61 ± 6.09 months in patients receiving docetaxel. Serum PSA decline rates at least 50% from baseline was seen in 34.32% of patients at 3 months on docetaxel treatment. Docetaxel was also effective in pain reduction, decline in serum PSA levels and improvement in health related quality of life. Our study demonstrates significant response rates to docetaxel chemotherapy but a considerable number of patients had treatment-related complications. This highlights the need for careful patient selection and optimization of chemotherapy dosing.

Keywords: Prostate cancer; Metastatic; Castration-resistant; Docetaxel

Introduction

Prostate cancer (PC) is the second most common cause of cancer and the sixth leading cause of cancer death among men worldwide. The worldwide PC burden is expected to grow to 1.7 million new cases and 499,000 new deaths by 2030 simply due to the growth and aging of the global population [1]. Prostate cancer may be curable in early stage disease, with the prostate cancer specific survival well over 90% at 15 years in some subgroups [2]. Androgen deprivation therapy (ADT) has become the primary treatment of metastatic prostate cancer. However, most men will progress on androgen-deprivation therapy and develop castration resistant prostate cancer (CRPC) [3].

Prostate cancer is a heterogeneous disease and contains cells that are androgen-sensitive and those that are androgen-resistant [4]. It is hypothesized that the androgen-resistant cells are eventually responsible for the failure of androgen deprivation therapy. CRPC is known to progress despite castrate levels of testosterone as a result of androgen receptor amplification [5,6]. It has also been suggested that reactivation of previously down-regulated androgen receptors could possibly be the mechanism of CRPC [7]. Till a decade ago there was no single agent or combination of agents that improved survival once CRPC developed. In 2004, two large randomized clinical trials showed a survival advantage of docetaxel based chemotherapy over mitoxantrone in patients with metastatic CRPC [8]. We report our early experience with docetaxel based chemotherapy in patients with metastatic CRPC.

Materials and Methods

Study period

During the study period July 2007 to March 2016 patients with metastatic prostate cancer presenting to the Urological services of the Hospital formed the study group following the...
Patients

Patients with mCRPC and aged ≥80 years having rising serum Prostate-specific antigen (PSA), increasing extent of disease on bone scans and physical findings, following androgen deprivation therapy were included into the study.

Variables

All the eligible patients underwent a detailed physical examination, routine blood and serum biochemistry tests (renal function tests, hemoglobin, blood sugar). Imaging studies included ultrasonography of the bladder and pelvis, Computed tomography scans of pelvis, MR imaging and radionuclide bone scans. All eligible patients with no previous treatment with any cytotoxic drug and having a good ≥80 Karnofsky performance status (KPS) were earmarked to receive docetaxel based chemotherapy.

All eligible patients received docetaxel 75 mg/m² every three weeks and oral prednisone 5 mg twice daily. Treatment was continued until disease progression, occurrence of unacceptable adverse effects or a minimum of 10 cycles of docetaxel. The primary endpoint was overall survival and the secondary endpoints were pain reduction, serum PSA decline rates of at-least 50% from baseline, quality of life and objective tumor response. The decline in serum PSA levels and overall survival was analyzed in relation to four independent baseline risk factors (i.e. Pain, visceral metastases, anemia and bone scan progression) which were further classified into group 1 (zero to one risk factor), group 2 (two risk factors) and group 3 (three to four risk factors). Pain was assessed using visual analogue scale (VAS) and the need for analgesia. The Functional Assessment of Cancer Therapy-Prostate (FACT-P) questionnaire was used to measure the health related quality of life in all these patients. Individual patient data were tabulated and summarized using descriptive statistics. Overall survival was calculated using the Kaplan-Meier method. All statistics were calculated using SPSS software package version 20.0.

Table 1

| Parameters             | Docetaxel Group n=67 | No treatment Group n=18 | p-value |
|------------------------|-----------------------|-------------------------|---------|
| Number of patients     | 67 (78.82%)           | 18 (21.18%)             | -       |
| Age (mean yrs)         | 67.05 ± 2.11          | 68.00 ± 4.47            | 0.104   |
| Pre-treatment PSA (ng/ml) | 11.05 ± 2.90      | 12.05 ± 2.62            | 0.612   |
| Risk Factors 0-1       | 26 (38.80%)           | 7 (38.80%)              | -       |
| Risk Factors 2         | 30 (44.77%)           | 7 (38.88%)              | -       |
| Risk Factors 3-4       | 11 (16.41%)           | 4 (22.22%)              | -       |
| KPS 100                | 23 (34.32%)           | 6 (33.33%)              | -       |
| KPS 90                 | 35 (52.33%)           | 9 (50%)                 | -       |
| KPS 80                 | 9 (13.43%)            | 3 (16.66%)              | -       |

The most common adverse reactions seen in the docetaxel group were anemia, febrile neutropenia, neuropathy, dyspnea, anorexia, fluid retention, asthenia, mucositis, alopecia and myalgia. In 20 (29.85%) patients on docetaxel, the chemotherapy cycles needed to be interrupted due to neutropenia and asthenia. However all the patients completed 10 cycles of docetaxel based chemotherapy. During the
therapy 15 (22.38%) patients on docetaxel therapy needed blood transfusions.

**Table 2** Decline of Sr. PSA levels following initiation of chemotherapy.

| Parameters | Docetaxel Group | No treatment Group |
|------------|-----------------|--------------------|
| Initial PSA | PSA after 3 M | PSA after 9 M | Initial PSA | PSA after 3 M | PSA after 9 M |
| Risk Factors 0-1 | 10.5 ± 1.72 | 4.5 ± 1.40 | 1.0 ± 0.51 | 9.00 ± 1.49 | 23.00 ± 3.72 | 26.00 ± 3.45 |
| Risk Factors 2 | 12.0 ± 0.83 | 8.0 ± 1.05 | 5.0 ± 1.93 | 13.00 ± 1.11 | 33.00 ± 5.12 | 38.00 ± 4.49 |
| Risk Factors 3-4 | 18.0 ± 3.05 | 14.0 ± 1.52 | 12.0 ± 1.52 | 14.50 ± 0.95 | 36.50 ± 6.37 | 47.00 ± 4.64 |
| All patients | 11.06 ± 2.96 | 6.67 ± 3.31 | 3.89 ± 3.99 | 12.06 ± 2.62 | 30.11 ± 8.10 | 34.78 ± 9.08 |

**Figure 1** Post-treatment decline in serum PSA in various risk groups.

The overall survival was 32.61 ± 6.09 months (95% confidence interval) in patients receiving docetaxel based chemotherapy as compared to 12.83 ± 2.40 months (95% Confidence Interval) in patients in the no treatment group. The overall survival (OS) was better in patients with least risk factors (Table 3 and Figure 2). During the course of treatment 13 patients in the docetaxel group needed to be treated with bone targeted radioisotope therapy using phosphorous-32 for relief of pain. The pain response rate in this group was 76.92%, PSA response was seen in 84.61% and OS was 29 months.

**Table 3** Overall survival in months.

| Parameters | Docetaxel (67) | No Treatment Group (18) | p-value |
|------------|----------------|-------------------------|---------|
| Risk Factors 0-1 | 38.25 ± 2.05 | 15.42 ± 1.61 | 0.0001 |
| Risk Factors 2 | 28.66 ± 1.65 | 11.57 ± 0.53 | 0.0001 |
| Risk Factors 3-4 | 23.00 ± 1.00 | 10.50 ± 0.57 | 0.005 |
| All Patients | 32.61 ± 6.09 | 12.83 ± 2.40 | <0.0001 |
FACT-P scores showed marked improvement in 43.28% of patients following completion of chemotherapy and was significantly higher than compared to patients not receiving any treatment (P<0.0001). The greatest benefit in the docetaxel group was in the subscale representing prostate-specific concerns (including weight loss, appetite, pain, physical comfort, and bowel and genitourinary function).

**Figure 2** Kaplan Meier overall survival function shows significant increase in the Docetaxel group.

**Discussion**

About 10-20% of men with prostate cancer present with metastatic disease, and in many others, metastases develop despite treatment with surgery or radiotherapy. In about 80 percent of men, primary androgen ablation leads to symptomatic improvement and a reduction in serum levels of prostate-specific antigen (PSA), but in all patients the disease eventually becomes refractory to hormone treatment after a median time of 18-24 months [3]. Castration-resistant prostate cancer is defined by disease progression despite androgen depletion therapy and may present as either a continuous rise in serum prostate-specific antigen levels, the progression of preexisting disease, and/or the appearance of new metastases.

In our study, 78.82% of very elderly (age ≥80 years) patients were able to complete planned treatment; however, a considerable proportion required dose alterations, highlighting the difficulties in determining optimal dosing in patients of advanced age. Similar findings have been reported in a retrospective series of 159 Japanese patients aged ≥75 years who received docetaxel in the community setting, in which 87% of patients required dose modifications [9]. Although secondary hormonal manipulations (e.g. corticosteroids, estrogens and ketoconazole) can benefit a subset of men with metastatic castrate-resistant prostate cancer (mCRPC), this benefit is usually short lived and until recently there was no evidence that secondary hormonal therapies could improve overall survival (OS) [10]. The options then include symptomatic care with narcotic analgesics, radiotherapy to dominant sites of bone pain, treatment with bone-seeking isotopes such as strontium-89, and cytotoxic chemotherapy [10,11].

Docetaxel and prednisone in combination are currently considered the standard of care for men with CRPC with detectable metastatic disease. Docetaxel is a taxane drug that induces polymerization of microtubules and phosphorylation of bcl-2 protein. Tannock et al.[12] randomized 1006 patients to one of three treatment arms: docetaxel (75 mg/m² intravenously every 3 weeks), docetaxel (30 mg/m² 5 times weekly for 5 of 6 weeks), or control therapy with mitoxantrone. The patients in our study also received oral...
prednisone 5 mg daily twice. Thus improved survival was observed with docetaxel (every 3 weeks) compared with mitoxantrone-prednisone median survival, 18.9 vs. 16.5 months; hazard ratio [HR]=0.76 (95% confidence interval [CI], 0.62-0.94), two-sided p=0.009). No overall survival benefit was observed with docetaxel given on a weekly schedule (HR=0.91, 95% CI, 0.75-1.11), two-sided p=0.36). Similar results were reported by Petrylak et al. [12] study on 666 eligible patients randomized to docetaxel and estramustine (EMP) or mitoxantrone- prednisone. In addition to dexamethasone premedication, patients in the docetaxel arm also received warfarin and/or acetylsalicylic acid (ASA) as thrombosis prophylaxis during the course of the trial.

Several investigators have reported that a PSA decline of ≥30% within 3 months of treatment with docetaxel or mitoxantrone has the highest degree of surrogacy for OS. Analyzing their series reported on four independent baseline factors (i.e. pain, visceral metastases, anemia and bone scan progression) that predicted PSA decline of ≥30% within 3 months of treatment with chemotherapy. On the basis of this finding, we developed three risk groups with a median OS of 25.7 months (0-1 risk factors), 18.7 months (2 risk factors) and 12.8 months (3-4 risk factors) [12-15]. Similarly, our results showed Serum PSA decline rates at least 50% from baseline was seen in 34.32% of patients at 3 months on docetaxel treatment. In our study the overall survival was 32.61 ± 6.09 months; hazard ratio [HR]=0.76 (95% CI, 0.62-0.94), two-sided p=0.009). No overall survival benefit was seen in patients receiving docetaxel. Serum PSA decline rates at least 50% from baseline was seen in 34.32% of patients at 3 months on docetaxel treatment. Docetaxel was also effective in pain reduction, decline in serum PSA levels and improvement in health related quality of life. In our study over 59.70% of the patients on docetaxel therapy had minor side effects.

However, as of today there are no tools to predict the benefit of chemotherapy in an individual patient with mCRPC. Further, In a post docetaxel setting, the number of progression factors (PSA, pain, and tumor size), duration of therapy and whether progression occurred during or after the treatment with docetaxel independently predicted post-progression survival in men with mCRPC [16]. These predictors of survival may be useful for prognostication as well as for stratification in and interpretation of clinical trials and sample size planning. Chemotherapy can reduce serum PSA levels in patients with mCRPC and known to relieve pain in some patients, but tolerability is of concern, particularly since most patients are elderly and many have other medical problems. Both the TAX 327 and SWOG 99-16 studies have shown improved survival for patients receiving 3 weekly docetaxel over mitoxantrone, with acceptable toxicity rates. In TAX 327 trial, it was noted that patients in the docetaxel arm were more likely to have at least one serious adverse event (26% vs. 20%) than the mitoxantrone arm. In SWOG 99-16 trial, 16% of the patients in the docetaxel group and 10% in the mitoxantrone group were withdrawn from the study as a result of adverse events. Grade 3 and 4 febrile neutropenia, cardiovascular events, and neurologic events were significantly higher in the docetaxel group [12,13]. In our study too over 59.70% of the patients on docetaxel therapy had minor side effects, 29.85% patients needed interruption of the therapy due to neutropenia and severe asthenia. However all the patients completed 10 cycles of docetaxel based chemotherapy and 22.38% of patients needed blood transfusions. Several docetaxel combinations have been evaluated in phase studies for mCRPC, including combinations with tyrosine kinase inhibitors, angiogenesis agents and immunologic agents [17].

Conclusions

In conclusion, Docetaxel remains the mainstay chemotherapeutic option in the management of mCRPC. The selected fit elderly patients appear to tolerate docetaxel and develop clinical benefit from treatment, but optimal dosing remains unclear, as pharmacokinetic changes with aging could potentially alter drug exposure and metabolism. Novel hormonal agents such as abiraterone and enzalutamide are striking substitutes in elderly patients due to lower rates of myelosuppression, though optimal treatment sequencing is not yet known. Comprehensive geriatric evaluation is likely to be helpful in determining patients at risk of increased toxicity, but is time-consuming and challenging to implement in a busy oncology practice. However, our study demonstrates significant response rates to docetaxel chemotherapy but that a considerable number of patients had treatment-related complications. This highlights the need for careful patient selection and optimization of chemotherapy dosing.

References

1. Ferlay J, Shin HR, Bray F (2010) GLOBOCAN 2008, Cancer Incidence and Mortality Worldwide: IARC Cancer Base No. 10. International Agency for Research on Cancer, Lyon, France.
2. Han M, Partin AW, Pound CR, Epstein JI, Walsh PC (2001) Long term biochemical disease free and cancer specific survival following anatomic radical retropubic prostatectomy. The 15 year John Hopkins experience. Urol Clin North Am 28: 555-565.
3. Eisenberger MA, Blumenstein BA, Crawford ED (1998) Bilateral orchectomy with or without flutamide for metastatic prostate cancer. N Engl J Med 339: 1036-1042.
4. Isaacs JT, Heston WD, Weissman RM, Coffey DS (1978) Animal models of the hormone-sensitive and insensitive prostate adenocarcinomas, Dunming R-3327-H, R-3327-H, and R-3327-AT. Cancer Res 38: 4353-4359.
5. Edwards J, Krishna NS, Grigor KM, Bartlett JM (2003) Androgen receptor gene amplification and protein expression in hormone refractory prostate cancer. Br J Cancer 89: 552-556.
6. Mellinghoff IK, Vivanco I, Kwon A, Tran C, Wongvipat J, et al. (2004) HER2/neu kinase-dependent modulation of androgen receptor function through effects on DNA binding and stability. Cancer Cell 6: 517-527.
7. Mousses S, Wagner U, Chen Y (2001) Failure of hormone therapy in prostate cancer involve systematic restoration of androgen responsive genes and activation of rapamycin sensitive signaling. Oncogene 20: 6718-6723.
8. Adesunloye BA, Dahut WL (2011) Current state of docetaxel based therapy in castration resistant prostate cancer. In Gulley JL (Ed) Prostate Cancer, Demos Medical, New York pp: 539-552.
9. Italiano A, Ortholan C, Oudard S, Pouessel D, Gravis G, et al. (2009) Docetaxel-based chemotherapy in elderly patients (age 75 and older) with castration-resistant prostate cancer. Eur Urol 55: 1368-1375.

10. Ryan CJ, Small EJ (2003) Role of secondary hormonal therapy in the management of recurrent prostate cancer. Urology 62: 87-94.

11. Saad F, Hotte SJ (2010) Guidelines for the management of Castrate Resistant Prostate Cancer. Can Urol Asso J 4: 380-384.

12. Tannock IF, de Wit R, Berry WR, Horti J, Pluzanska A, et al. (2004) Docetaxel plus prednisone or mitoxantrone plus prednisone for advanced prostate cancer. N Engl J Med 351: 1502-1512.

13. Petrylak DP, Tangen CM, Hussain MHA, Lara PN, Jones JA, et al. (2004) Docetaxel and estramustine compared with mitoxantrone and prednisone for advanced refractory prostate cancer. N Engl J Med 351: 1513-1520.

14. Petrylak DP, Ankerst DP, Jiang CS (2006) Evaluation of prostate specific antigen declines for surrogacy in patients treated on SWOG 99-16. J Natl Cancer Inst 98: 516-521.

15. Armstrong AJ, Garrett-Mayer E, Ou Yang YC (2007) Prostate specific antigen and pain surrogacy analysis in metastatic hormone refractory prostate cancer. J Clin Oncol 25: 3965-3970.

16. Armstrong AJ, Garrett-Mayer E, de Wit R (2010) Prediction of survival following first line chemotherapy in men with castration-resistant metastatic prostate cancer. Clin Cancer Res 16: 203-211.

17. Galsky MD, Vogelzang NJ (2010) Docetaxel based combination therapy for castration resistant prostate cancer. Annals of Oncol 21: 2135-2144.