A prognostic model for stratifying clinical outcomes in chemotherapy-naive metastatic castration-resistant prostate cancer patients treated with abiraterone acetate

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Abstract

Introduction: Recently, a prognostic index including six risk factors (RFs) (unfavourable Eastern Cooperative Oncology Group performance status [ECOG PS], presence of liver metastases, short response to luteinizing hormone-releasing hormone [LHRH] agonists/antagonists, low albumin, increased alkaline phosphatase [ALP] and lactate dehydrogenase [LDH]) was developed from the COU-AA-301 trial in post-chemotherapy metastatic castration-resistant prostate cancer (mCRPC) patients treated with abiraterone. Our primary objective was to evaluate this model in a cohort of chemotherapy-naive mCRPC patients receiving abiraterone. Methods: We identified 197 chemotherapy-naive patients who received abiraterone at six BC Cancer Agency centres and who had complete information on all six RFs. Study endpoints were prostate-specific antigen (PSA) response rate (RR), time to PSA progression, and overall survival (OS). PSA RR and survival outcomes were compared using \( \chi^2 \) test and log-rank test. Multivariable Cox proportional hazard analysis was performed to identify RFs independently associated with OS.

Results: Patients were classified into good (0–1 RFs), intermediate (2–3 RFs), and poor (4–6 RFs) prognostic groups (33%, 52%, and 15%, respectively). For good-, intermediate-, and poor-risk patients, PSA RR (≥50% decline) was 60% vs. 42% vs. 40% (p=0.05); median time to PSA progression was 7.3 vs. 5.3 vs. 5.0 months (p=0.02); and median OS was 29.4 vs. 13.8 vs. 8.7 months (p<0.0001).

Conclusions: The six-factor prognostic index model stratifies clinical outcomes in chemotherapy-naive mCRPC patients treated with abiraterone. Identifying patients at risk of poor outcome is important for informing clinical practice and clinical trial design.

Introduction

The therapeutic landscape for metastatic castration-resistant prostate cancer (mCRPC) has rapidly evolved in recent years, with many new agents demonstrating a benefit in overall survival (OS). One of these agents is abiraterone acetate (abiraterone), an orally available inhibitor of CYP17 that blocks adrenal and intra-tumoural androgen synthesis. Abiraterone confers an OS advantage both as first-line therapy in chemotherapy-naive patients, as well as in the post-docetaxel chemotherapy setting; however, despite its efficacy, outcomes with abiraterone are variable and not all patients derive benefit from treatment. Thus, there is an urgent need for a practical clinical tool able to stratify patients accurately into distinct prognostic categories.

In a recent post-hoc analysis of the COU-AA-301 study, a prognostic model was developed for predicting OS in post-docetaxel patients treated with abiraterone. This model comprises six risk factors (RFs) linked to poor survival: Eastern Cooperative Oncology Group performance status (ECOG PS) ≥2, liver metastases, time from initiation of androgen-deprivation therapy (ADT) to initiation of abiraterone ≤36 months, low albumin (≤4g/dl), lactate dehydrogenase (LDH) above upper limit normal (ULN) and alkaline phosphatase (ALP) above ULN. Classification of the COU-AA-301 trial population into good (0–1 RFs), intermediate (2–3 RFs), and poor (4–6 RFs) risk groups revealed OS of 21.3 months, 13.9 months (hazard ratio [HR] 2.3) and 6.1 months (HR 6.2), respectively. This model was subsequently validated in a population-based cohort of post-chemotherapy mCRPC patients treated with abiraterone. The utility of this model in chemotherapy-naive patients has also been examined in a preliminary analysis incorporating only 64 patients.
The COU-302 trial, which evaluated abiraterone in chemotherapy-naive patients with CRPC, showed a significantly longer OS than the post-chemotherapy COU-301 trial (34.7 vs 15.8 months), and the performance of our six-factor prognostic model in that setting is currently unknown. The aim of our study was to determine whether this prognostic model determines clinically relevant prognostic groups in chemotherapy-naive patients treated with abiraterone in a real-world setting across six cancer centres in British Columbia, Canada.

Methods

Patient population

The BC Cancer Agency (BCCA) consists of six distinct centres and coordinates cancer care delivered throughout British Columbia, Canada. The Cancer Registry at BCCA was reviewed for chemotherapy-naive mCRPC patients who started abiraterone between July 2009 and October 2014. Patient demographics, prior treatments, clinicopathological characteristics, and outcomes on abiraterone were documented from medical records of each patient. Only patients with available data for all six RFs in the prognostic index were included in this study. Research ethics board approval was obtained prior to commencing this study.

Outcome measures

Patients were classified into good (0–1 RFs), intermediate (2–3 RFs) and poor (4–6 RFs) risk groups. The following endpoints were determined for each patient: confirmed prostate-specific antigen (PSA) response rate (PSA decline ≥50% from baseline maintained for ≥3 weeks), time to PSA progression (Prostate Cancer Working Group 2 [PCWG2] criteria), time on treatment (time from initiation of abiraterone to discontinuation for any reason), and OS (time from initiation of abiraterone to death of any cause or censoring on November 1, 2016). Reasons for discontinuation of abiraterone were recorded as follows: radiographic (PCWG2 criteria), biochemical (PCWG2 criteria), or clinical (worsening disease-related symptoms requiring a change in anti-neoplastic therapy or a decrease in ECOG PS of ≥2 levels).

Development of the prognostic model

Construction of the six-factor prognostic model has been previously described. In brief, the following steps were involved: 1) key clinicopathological factors were identified and dichotomised for high/low values as necessary; 2) association between baseline clinicopathological factors and OS was investigated using a univariate Cox proportional hazards model; 3) factors that were significant on univariate analysis were incorporated into a multivariate Cox proportional hazards regression model (stepwise procedure); 4) factors that were significant on multivariate analysis were incorporated into the final model, which was then subjected to validation by a bootstrapping approach; 5) the C-index was used to determine accuracy of the model, which comprised six separate RFs; and 6) patients were then classified into risk groups based on the number of baseline RFs with median OS calculated for each risk group.

Statistics

Univariate analysis examining association between prognostic group and PSA response was performed using X² test. Survival outcomes were estimated using the Kaplan-Meier method. Log-rank test was performed to assess survival differences between groups. Multivariable analysis using Cox proportional hazards model was performed to identify RFs independently associated with OS. Statistical analysis was performed using SPSS® v.14.0 software. To determine the model’s accuracy, the C-index was calculated for time on treatment, time to PSA progression, and OS.

Results

Patient population

Two hundred and forty-six chemotherapy-naive patients, who received abiraterone from July 2009 (when abiraterone became available) until October 2014, were identified. One hundred and ninety-seven patients were included for this analysis, as they had available data for each of the six RFs comprising the six-factor prognostic model. Patient characteristics at initiation of abiraterone are listed in Table 1. The median age at start of abiraterone was 80 years (interquartile range 71–84), 38% of patients had ECOG PS of 2 or higher, and 3% had liver metastasis. Overall, 33% (65/197), 52% (102/197), and 15% (30/197) of patients were classified as good, intermediate and poor prognosis, respectively, as per the prognostic index. Reasons for abiraterone discontinuation (more than one could apply) were clinical progression (36%), PSA progression (67%), radiological progression (33%), and toxicity (8%), while 5.5% (11/197) of patients were still on abiraterone as of November 1, 2016. Post-abiraterone systemic treatment was administered to 54% of patients (100/186).

PSA response

Confirmed PSA declines ≥90%, ≥50%, and ≥30% were seen in 13% (26/197), 48% (94/197), and 51% (100/197)
Stratifying clinical outcomes in mCRPC

For the overall cohort, median time to PSA progression, median time on treatment, and median OS was 6.5 months (95% confidence interval [CI] 5.6–8.0), 7.4 months (95% CI 6.0–8.5), and 15.7 months (95% CI 12.9–18.9), respectively. As shown in Table 2, time to PSA progression and time on treatment were significantly longer in the good prognosis group vs. the intermediate- and poor-risk groups (p=0.02 and p<0.0001, respectively; log-rank). Hazard ratios for each group are shown in Table 3. OS was significantly better in the good prognosis group compared with the intermediate and poor prognostics groups (29.4 vs. 13.8 vs. 8.7 months, respectively; p<0.0001). Kaplan-Meier curves are shown in Fig. 1. On multivariable analysis incorporating the six RFs from the prognostic index, ECOG PS (p<0.0001), liver metastases (p= 0.0008), time from initiation of ADT to abiraterone ≤36 months (p= 0.02), and serum ALP (p<0.0001) were confirmed as independent prognostic factors for OS, whereas serum albumin and serum LDH did not meet statistical significance (Table 4). The predictive accuracy of our model as measured by the C-index was 0.79, 0.68, and 0.66 for OS, time on treatment, and time to PSA progression, respectively.

Discussion

Although abiraterone has proven efficacy in chemotherapy-naive patients with mCRPC, treatment outcomes are variable and difficult to predict at an individual patient level. Recently, a prognostic model for OS in patients receiving abiraterone after prior docetaxel chemotherapy was developed based on data from the COU-AA-301 phase 3 trial, and subsequently validated in a population-based cohort. In the present study, we confirm the performance and discriminatory power of this model in a large, unselected and sequential cohort of chemotherapy-naive mCRPC patients treated with abiraterone. Our real-world population differed significantly from patients in the COU-AA-302 study, which established the efficacy of abiraterone in chemotherapy-naive patients. We included patients with poor performance status, significant pain symptoms, and visceral metastasis, all of which were exclusion criteria for the COU-AA-302 study. This is reflected in the difference between the median OS of our cohort (15.7 months) compared to the median OS observed in the COU-AA-302 study (34.7 months). In addition, a lower proportion of our patients had low-risk vs. intermediate- or high-risk prognostic scores (33% vs. 67%, respectively), compared to the COU-AA-301 study (46% vs. 53%). This also further emphasizes the applicability and generalizability of the prognostic model to the broader patient population treated with abiraterone.

We observed a clear and statistically significant difference in OS between good-, intermediate-, and poor-risk patients and confirmed that presence of liver metastases, time from initiation of ADT to initiation of abiraterone ≤36 months, serum ALP ≥ ULN, and ECOG PS ≥2 are independent prognostic factors. Interestingly, serum LDH and serum albumin showed no independent prognostic value for OS, although this is likely due to the limited sample size of our study. These results contrast with those of an external validation
of the prognostic index performed by Ravi et al at Royal Marsden.\textsuperscript{(6)} In their cohort of 64 chemotherapy-naïve patients, low albumin was the only independent factor predicting OS in multivariable analysis; however, this was a relatively small validation cohort that included only one chemotherapy-naïve patient with poor prognosis disease and differed from our cohort in that no chemotherapy-naïve patients with ECOG PS ≥2 were included (these patients comprised 38% of our cohort). The different cohort characteristics and larger sample size may also account for the better predictive power of the model in our study, as Ravi et al observed a relatively modest OS difference of 10 months between good- and intermediate-/poor-risk patients.

There is strong evidence from other studies supporting the prognostic value of the clinical factors included in our model. A prognostic nomogram was developed and validated by Halabi et al from two randomized controlled trials of first-line docetaxel for mCRPC.\textsuperscript{(12)} Their analysis also showed that ECOG PS, site of metastasis, LDH, albumin, hemoglobin, and ALP predict OS on multivariable analysis, in addition to opioid analgesia use and PSA; however, they did not test the prognostic significance of duration of primary ADT when building their model, and its performance in the setting of first-line abiraterone has not been verified. In a study of 161 patients treated with abiraterone, McKay et al demonstrated that duration of primary ADT >12 months and no prior docetaxel chemotherapy were independently associated with a longer time on abiraterone.\textsuperscript{(11)} In addition, a recent meta-analysis evaluating the impact of site of metastasis on OS in men with CRPC showed that liver metastasis predicted worse OS compared to bone metastasis and lymph node only metastasis (13.5 vs. 21.3 vs. 31.6 months, respectively).\textsuperscript{(14)}

In addition to OS, we observed that the six-factor prognostic index model is predictive of time to PSA progression, PSA response, and time on treatment in chemotherapy-naïve patients treated with abiraterone. The use of PSA parameters as a surrogate endpoint for OS in CRPC has generated considerable discussion and controversy; however recently, Xu et al constructed a biomarker survival modelling framework to explore the relationship between PSA kinetics (including time to PSA progression and PSA response) and OS in mCRPC patients following administration of abiraterone.\textsuperscript{(15)} In their analysis, which was based on data from the COU-AA-301 and COU-AA-302 trials, PSA kinetics were highly associated with OS in both chemotherapy-experienced patients and chemotherapy-naïve patients. The authors concluded that PSA kinetics should be considered as surrogate endpoints of clinical benefit in abiraterone-treated patients regardless of chemotherapy treatment.\textsuperscript{(15)} Our observation that the prognostic model predicted for both PSA response and OS in the present cohort is in accordance with this data. Our model was also predictive for time on treatment, a useful surrogate in the real-world setting for the duration of clinical benefit on treatment and an important endpoint, as highlighted in the updated recommendations on trial design and objectives from the PCWG 3.\textsuperscript{(16)} These findings demonstrate that our model is predictive of outcomes on abiraterone and thus may be valuable in selecting patients likely to benefit

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Good prognosis (0–1 RF)</th>
<th>Intermediate prognosis (2–3 RF)</th>
<th>Poor prognosis (4–6 RF)</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>PSA decline\textsuperscript{a}</td>
<td>Decline ≥90, n (%)</td>
<td>14 (22)</td>
<td>11 (11)</td>
<td>1 (3)</td>
</tr>
<tr>
<td></td>
<td>Decline ≥50, n (%)</td>
<td>39 (60)</td>
<td>43 (42)</td>
<td>12 (40)</td>
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<tr>
<td></td>
<td>Decline ≥30, n (%)</td>
<td>40 (62)</td>
<td>46 (45)</td>
<td>14 (47)</td>
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<tr>
<td>Time on treatment</td>
<td>Median (month, 95% CI)</td>
<td>11.6 (9.7–17.2)</td>
<td>5.8 (4.8–7.8)</td>
<td>5.3 (2.9–6.9)</td>
</tr>
<tr>
<td>Time to PSA progression\textsuperscript{b}</td>
<td>Median (months, 95% CI)</td>
<td>7.3 (5.7–9.4)</td>
<td>5.3 (4.6–6.5)</td>
<td>5.0 (2.8–6.4)</td>
</tr>
<tr>
<td>Overall survival\textsuperscript{c}</td>
<td>Median (months, 95% CI)</td>
<td>29.4 (22.6–38.7)</td>
<td>13.8 (11.4–16.1)</td>
<td>8.7 (5.8–11.9)</td>
</tr>
</tbody>
</table>

\textsuperscript{a} PSA decline confirmed ≥3 weeks later; \textsuperscript{b} PCWG2 criteria; \textsuperscript{c} from time of commencing abiraterone. CI: confidence interval; PSA: prostate-specific antigen; RF: risk factors (including: ECOG performance status ≤2; liver metastases; time on androgen-deprivation therapy to initiation of abiraterone ≥36 months; low albumin <4 g/dl; high lactate dehydrogenase >upper limit normal); and high alkaline phosphatase (upper limit normal).
stratifying clinical outcomes in mCRPC from first-line abiraterone, and identifying those for whom alternate treatments could be considered, such as first-line chemotherapy or clinical trials.

A key strength of the six-factor prognostic index model is that it is a pragmatic tool for risk stratification using easily available (and inexpensive) clinical parameters. Nevertheless, integration of this model with emerging molecular biomarkers, including androgen receptor (AR) splice variants, circulating tumour DNA (ctDNA), and circulating tumour cells (CTCs), will be important. AR splice variant 7 (ARv7) detection in CTCs was recently proposed as a mechanism driving primary resistance to enzalutamide, and high levels of full-length AR mRNA and presence of ARv7 have been shown to correlate with PSA progression-free survival and OS on abiraterone or enzalutamide. Recent evidence has revealed that structural variants of the AR gene are associated with the presence of AR splice variants, and may also be important drivers of treatment resistance. In addition, AR gene aberrations (copy number change and/or mutations) in pre-treatment ctDNA have been linked to adverse outcomes in mCRPC patients commencing abiraterone and enzalutamide. A biomarker panel incorporating CTC enumeration and LDH was also recently shown to predict OS in a post-hoc analysis of the COU-AA-301 trial.

The present study has various limitations. These include its retrospective design, being limited to a single province in Canada, and the relatively small sample size. We could not assess radiological response to treatment since imaging was not consistently performed. Data on radiographic progression-free survival were not analyzed due to wide variation in followup, including timing of imaging.

**Conclusion**

We observe that the six-factor prognostic index model provides reliable risk stratification for chemotherapy-naive patients receiving abiraterone by predicting PSA response, time to PSA progression, time on treatment, and OS. ECOG PS, liver metastases, time from ADT to initiation of abiraterone ≤36 months, and serum ALP were confirmed as independent risk factors for poor OS in the pre-chemotherapy setting. Due to its predictive capability, we suggest that incorporating the prognostic model in clinical practice and future trials assessing the use of novel hormonal agents and cytotoxics will allow improved risk stratification and optimized treatment selection.

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>HR</th>
<th>95% CI</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time on ADT</td>
<td>1.5</td>
<td>1.1–2.1</td>
<td>0.02</td>
</tr>
<tr>
<td>ECOG PS</td>
<td>2.2</td>
<td>1.6–3.1</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Serum LDH</td>
<td>1.2</td>
<td>0.8–1.7</td>
<td>0.39</td>
</tr>
<tr>
<td>Serum ALP</td>
<td>2.1</td>
<td>1.5–3.0</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Liver metastases</td>
<td>5.0</td>
<td>1.9–12.7</td>
<td>0.0008</td>
</tr>
<tr>
<td>Serum albumin</td>
<td>1.0</td>
<td>0.7–1.4</td>
<td>0.88</td>
</tr>
</tbody>
</table>

ADT: androgen-deprivation therapy; ALP: alkaline phosphatase; CI: confidence interval; ECOG PS: Eastern Cooperative Group Performance Status; HR: hazard ratio; LDH: lactate dehydrogenase; mCRPC: metastatic castration-resistant prostate cancer.

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**Fig. 1.** Kaplan-Meier curves for: (A) overall survival; (B) time on treatment; and (C) prostate-specific antigen (PSA) progression for good, intermediate, and poor prognosis groups.
Competing interests: Dr. Azad has been an advisor for Astellas, AstraZeneca, Novartis, Sanofi, and Tolmar; a speaker for Janssen; has received grants/honoraria from Astellas; and has participated in clinical trials supported by AstaZeneca, GSK, MedImmune, Medivation, Merck, Sanofi, and Serono. Dr. Todenhöfer has received speaker honoraria from Astellas and Janssen. Dr. Egi has been an advisor for AstraZeneca, Novartis, and Roche; has received grants from Roche; and has participated in clinical trials supported by AstraZeneca, Novartis, Pfizer, and Roche. Dr. Finch has received speaker honoraria from and has participated in clinical trials supported by Astellas, AstraZeneca, Janssen, Pfizer, and Roche. Dr. Le has received speaker honoraria from Bayer; and was involved in the BC Cancer Agency Canadian Cancer Trials Group. Dr. Kollmannsberger has been an advisor for Astellas, BMS, Novartis, and Pfizer; has received speaker honoraria from Astellas, BMS, Novartis, Pfizer, and Sanofi; and has participated in clinical trials supported by Aetelas, BMS, Janssen, Novartis, and Pfizer. Dr. Chi has received grants/honoraria from Astellas, Bayer, Essai, Janssen, and Sanofi; and has participated in clinical trials supported by Astellas, AstraZeneca, Bayer, Essai, Lilly, Janssen, Merci, and Sanofi. The remaining authors report no competing personal or financial interests related to this work.

This paper has been peer-reviewed.

References


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