Finding the optimal treatment sequence in metastatic castration-resistant prostate cancer—a narrative review

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Abstract: Over the last two decades, there has been significant progress in the treatment of metastatic prostate cancer. Multiple treatments with diverse mechanisms of action have improved clinical outcomes for patients with metastatic castration-resistant prostate cancer (mCRPC) including taxane chemotherapy, immunotherapy, potent androgen receptor pathway inhibitors (ARPI), and radiopharmaceuticals (radium-223). As these treatments have entered standard clinical practise, clinicians have been challenged on how to optimally select and sequence them as the landmark studies establishing their efficacy had control arms with placebo or minimally effective therapy and there is a paucity of prospective trials examining treatment sequencing. More recently, the situation has been further complicated as the earlier up-front use of docetaxel and ARPI with standard gonadal testosterone inhibition has been shown to impart substantial improvements in disease control and survival for patients with castration sensitive prostate cancer. As new therapies enter into clinical practise such as the inhibitors of Poly (ADP-Ribose) Polymerase and Prostate Specific Membrane Antigen (PSMA)-targeted therapy, how to optimally select and sequence available treatments will be a continued dilemma in the absence of validated predictive biomarkers. This review will summarize the literature supporting the use of each active agent in mCRPC. We will propose a framework which will guide the selection of appropriate agents based on prior therapies, disease characteristics and biomarkers.

Keywords: Metastatic castration resistant prostate cancer; treatment; androgen receptor pathway inhibitors; taxanes; novel drugs; biomarkers

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Introduction

Prostate cancer is the most common deadly malignancy in men and resulted in an estimated 33,330 deaths for males in the United States and 78,800 deaths in the European Union in 2020 (1,2). Gonadal testosterone ablation was the earliest identified treatment for metastatic prostate cancer with proven efficacy. Its suppressive effect on metastatic prostate cancer was first demonstrated in 1941 by Huggins et al. (3), and suppression of gonadal androgen synthesis, either pharmacologically [luteinizing hormone releasing hormone (LHRH) antagonists or agonists, also referred to as androgen deprivation therapy (ADT)] or surgically remains the backbone of treatment for prostate cancer in the metastatic state to this day (4,5). Despite suppressed testosterone levels, prostate cancer will progress to a disease state, termed castration-resistant prostate cancer (CRPC) (6).

Encouragingly, the number of active treatment options for mCRPC has expanded considerably in recent years (Table 1).
Docetaxel, a microtubule-stabilizing taxane chemotherapy, was the first drug to show survival benefit in the treatment of mCRPC (10). In addition to docetaxel's direct effect on microtubule dynamics, it may also act, in part, by inhibiting androgen receptor (AR) trafficking to the nucleus. In fact, activation of the AR pathway is a hallmark of prostate cancer and is the central driver of progression, even in the castration-resistant state, in the majority of patients (18). The AR pathway inhibitors (ARPIs) are a class of oral agents which include the CYP 17A1 inhibitor abiraterone acetate, administered with prednisone (AAP), which inhibits extragonadal androgen synthesis and the direct AR inhibitors enzalutamide, apalutamide and darolutamide. This class of agents is among the most effective and well tolerated for CRPC (7-9,19-22). Cabazitaxel, is a taxane chemotherapy with proven activity in patients who have progressed both on ARPI as well as docetaxel with a demonstrated survival benefit both after one or two lines of therapy (13). Lastly, the alpha-particle emitting radiopharmaceutical Radium-223, also prolongs survival in patients with predominantly bony metastases (14). All of the aforementioned drugs have become standard of care in the treatment of mCRPC, although the best sequence in which they should be administered is unknown.

Furthermore, a number of these drugs are now approved for the treatment of up-front castration-sensitive disease, with ADT. Indeed, consecutive phase III randomized placebo-controlled trials of docetaxel (21,22) and AAP (23,24) combined with ADT demonstrated a substantial delay in disease progression and improvement in overall survival, compared with upfront ADT alone. Recent phase III randomized trials of enzalutamide (25,26) and apalutamide (27), have shown these agents to be similarly efficacious for metastatic castration-sensitive disease. This intensification of treatment earlier in the disease trajectory has undoubtedly been an important step forward, however, secondarily, the effectiveness of downstream therapy is diminished due to cross-resistance between agents and, potentially, development of more aggressive and treatment refractory disease phenotype upon progression to mCRPC.

Encouragingly, several novel agents for the treatment of mCRPC are anticipated to enter clinical practice in the near future. Notably, the Poly (ADP-Ribose) Polymerase (PARP) inhibitors have promising activity in tumours with genomic defects affecting double-stranded DNA repair by homologous-recombination, with olaparib (16) and rucaparib (28) now granted FDA approval and niraparib (29) which has been granted breakthrough therapy designation. In addition, $^{177}$Lutetium-PSMA-617 a Prostate Specific Membrane Antigen (PSMA) targeting radioligand labeled with β-particle emitting Lutetium$^{177}$ has shown promising activity in phase II trials (30).

This increase in the number of active agents poses a therapeutic dilemma, given the little data available to help select among the multiple possible treatment sequences. Indeed early landmark trials compared agents to now obsolete treatments such as mitoxantrone or placebo, and the majority of these proven agents have not been compared to one another in a randomized trial. A small numbers of trials have directly investigated treatments sequences and retrospective studies have provided insight into the degree of cross-resistance between treatments. Therapeutic choices are therefore guided by the side effect profile of the various agents, patient characteristics, potential biomarkers and individual clinician experience. In this review, the prospective evidence which supports current management of mCRPC will be summarized and the best evidence guiding the sequential use of therapies will be provided. Future promising agents as well as the emerging use of genomic biomarkers will also be described. We present the following article in accordance with the Narrative Review reporting checklist (available at http://dx.doi.org/10.21037/tau-20-1341).

Methods

Literature was reviewed using PubMed and the following keywords were searched: metastatic castrate resistant prostate cancer, treatment and sequence/ing. A focus was put on publication within the last 20 years (2000 to 2020), however, key work published prior to this time frame was included. The articles were selected according to their design with a preference to randomized control trials, their relevance to this review and language (English). The references from all the articles found were searched for further relevant literature.

Sequential use of ARPIs

The ARPIs as well as the taxane chemotherapy docetaxel are mainstays of first line treatment for mCRPC. However, ARPIs are favored over docetaxel for first-line treatment based on safety and side-effect profile with docetaxel typically relegated to the second line. Furthermore, many patients may not be eligible for docetaxel due to comorbidities or frailty; hence a strategy of treating...
Table 1 Active treatments for mCRPC and supporting data

| Agent | Evidence | Investigational arm [n] | Control arm [n] | Eligibility criteria | Permitted prior therapy for mCRPC | PSA RR/ radiographic RR | PFS (months) | OS (months) | Biomarkers Ref |
|-------|----------|-------------------------|-----------------|----------------------|-----------------------------------|------------------------|--------------|-------------|----------------|
| ARPI Naïve/Taxane Naïve | | | | | | | | | |
| Abiraterone COU-AA-302 | Phase III, double blind, randomized, placebo controlled trial | Placebo+ Prednisone [542] | AAP [546] | Inc: ECOG PS 0-1, No-mild symptoms (BPI-SF 0-3), Ex: Visceral mets excluded | 1st generation anti-androgens | PSA RR, 62% vs. 24% (HR 2.59, 95% CI: 2.19–3.05, P<0.001), Radiographic RR, 36 vs. 16% (HR 2.27, 95% CI: 1.59–3.25, P<0.001) | rPFS 16.5 vs. 8.3 (HR 0.53 95%CI, P<0.001) | 34.7 vs. 30.3 (HR 0.81, 95% CI: 0.7–0.93, P=0.0033) | (7) |
| Enzalutamide PREVAIL | Phase III, randomized, double blind, placebo controlled trial | Placebo (845) | Enzalutamide [872] | Inc: ECOG PS 0-1, No-mild symptoms (BPI-SF 0-3), Ex: MH/predisposition for seizures | 1st generation anti-androgens corticosteroids | PSA RR 78% vs. 3% (P<0.001), Radiographic RR 59% vs. 5% (P<0.001) | rPFS 20 vs. 5.4 (HR 0.32, 95% CI: 0.28–0.36, P<0.0001) | 35.3 vs. 31.3 (HR 0.77 95% CI: 0.67–0.88, P=0.0002) | (8,9) |
| Docetaxel TAX-327 | Phase III, randomized, open label | Mitoxantrone + prednisone [335]; [337] | Docetaxel (75 mg q3w) + prednisone [334] | Inc: KPS ≥60, Ex: Peripheral Neuropathy ≥ gr 2, Abnormal cardiac function | Docetaxel q3w vs. mitoxantrone PSA RR 45% vs. 32% (P<0.001), Radiographic response 12% vs. 7% (P=0.11) | – | docetaxel q3w vs. mitoxantrone 18.9 vs. 16.5 (HR 0.76, P=0.009) | (10) |
| | | | | | | | | | |
| Post taxanes/ARPI naïve | | | | | | | | | |
| Abiraterone COU-AA-301 | Phase III, double blind, randomized, placebo controlled trial | Placebo+ prednisone [398] | AAP [797] | Inc: Prior docetaxel ECOG PS ≤2; Ex: Uncontrolled hypertension | Docetaxel (mandatory), Other chemotherapies | PSA RR 29.1% vs. 5.5% (P<0.001), Radiographic RR 14% vs. 2.8% (P<0.001) | rPFS 5.6 vs.3.6 (HR 0.67, 95% CI: 0.55–0.78, P<0.001) | 14.8 vs. 10.9 (HR 0.66, 95% CI: 0.55–0.78, P<0.001) | (11) |
| Enzalutamide AFFIRM | Phase III, double blind, randomized, placebo controlled trial | Placebo [399] | Enzalutamide [800] | Inc: Prior docetaxel, ECOG PS ≤2 | Docetaxel (mandatory), Other chemotherapies, Up to 2 lines of chemotherapy in total corticosteroids | PSA RR 54% vs. 2% (P=0.001), Radiographic RR 29% | 18.4 vs. 13.6 (HR 0.63, 95% CI: 0.53–0.75, P<0.001) | (12) |

Table 1 (continued)
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| Agent | Evidence | Investigational arm [n] | Control arm [n] | Eligibility criteria | Permitted prior therapy for mCRPC | PSA RR/ radiographic RR | PFS (months) | OS (months) |
|-------|----------|-------------------------|-----------------|----------------------|----------------------------------|------------------------|--------------|-------------|
| Cabazitaxel TROPIC | Phase III, randomized, open label | Cabazitaxel + prednisone [377] + prednisone [378] | Mitoxantrone + prednisone | Inc: Prior docetaxel, ECOG PS ≤2, Ex: Previous mitoxantrone, Radiation to >40% of bone-marrow, LVEF <50%, Peripheral Neuropathy ≥ gr 2 | Docetaxel (mandatory), Anti-androgens | PSA RR 39.2% vs. 17.8% (P=0.0002); Radiographic RR 14.4% vs. 4.4% (P=0.0005) | PSA PFS and rPFS and symptomatic PFS 2.8 vs. 1.4 (HR 0.74, 95% CI: 0.64–0.86, P<0.0001) | 15.1 vs. 12.7 (HR 0.7, 95% CI: 0.59–0.83, P<0.0001) |
| Ra-223 ALSYMPCA | Phase III, randomized, open label | Radium 223 [614] Physicians’ choice of best standard of care [307] | Physicians’ choice of best standard of care [307] | Inc: ECOG PS ≤2; 2 or more bone mets; Prior docetaxel or ineligible/decline docetaxel. Ex: Systemic radioisotope treatment within 24 weeks; Blood transfusion or erythropoetin stimulating agent within 4 weeks; Lymphadenopathy ≥3 cm; Visceral mets | Docetaxel, Anti-androgens corticosteroids | – | – | 14.9 vs. 11.3 (HR 0.7, 95% CI: 0.58–0.83, P<0.001) |
| Post ARPI/Post Taxane | | | | | | | | |
| Cabazitaxel CARD | Phase III, randomized, open label | Cabazitaxel [129] AAP [58]/ enzalutamide [66] (ITT 126) | | Inc: Prior Docetaxel, Prior ARPI, Progression with 12 months of initiation of ARPI | Docetaxel/ARPI for mCSPC, Docetaxel (mandatory) AAP vs. 12% (P=0.004) | PSA RR 35.7% vs. 13.5% (P<0.001); Radiographic RR 37% vs. 12% (P=0.004) | rPFS 8 vs. 3.7 (HR 0.54 95% CI: 0.46–0.89, P=0.008) | 13.6 vs. 11 (HR – 0.64, 95% CI: 0.46–0.89, P=0.008) |

Table 1 (continued)
| Agent  | Evidence          | Investigational arm [n] | Control arm [n] | Eligibility criteria | Permitted prior therapy for mCRPC | PSA RR/ radiographic RR | PFS (months) | OS (months) | Biomarkers Ref |
|--------|-------------------|--------------------------|----------------|----------------------|-----------------------------------|------------------------|--------------|-------------|----------------|
| Olaparib PROfound | Phase III, randomized, open label | Olaparib [162] | AAP/ enzalutamide [83] | Inc: Prior enzalutamide/AAP ARPI for nmCRPC/ mCRPC/ mCSPC, Taxane chemotherapy | Cohort A (BRCA1, BRCA2, ATM), PSA RR 43% vs. 8%, radiographic RR 33% vs. 2% (P<0.001). Overall population (cohort A+B), PSA RR 30% vs. 10%; PSA RR 22% vs. 4% | Cohort A (BRCA1, BRCA2, ATM), rPFS 7.4 vs. 3.6 (HR 0.34, 95% CI: 0.25–0.47, P<0.001). Overall population (cohort A+B), rPFS 5.8 vs. 3.5 (HR 0.49, 95% CI: 0.38–0.63, P<0.001) | Cohort A (BRCA1, BRCA2, ATM), OS 19.1 vs. 14.7 (HR 0.69, 95% CI: 0.5–0.97, P=0.02). Cohort B (12 other HRR), OS 14.1 vs. 11.5 (HR 0.96, 95% CI: 0.63–1.49), Overall population (cohort A+B), OS 17.3 vs. 14 (HR 0.79, 95% CI: 0.61–1.03) | BRCA1, BRCA2, ATM, BRIP1, CDK12, CHEK1, CHEK2, PALB2, RAD51B, RAD51C, RAD51D, RAD54L |

Primary outcomes are in bold. n, number; PSA, prostate specific antigen; RR, response rate; PFS, progression free survival; OS, overall survival; ref, reference; ARPI, androgen receptor pathway inhibitors; AAP, abiraterone acetate and prednisone; inc, inclusion criteria; ex, exclusion criteria; rPFS, radiographic progression free survival; HR, hazard ratio; mets, metastases; LVEF, left ventricular ejection fraction; nmCRPC, non metastatic castrate resistant prostate cancer.
Sequencing of ARPI and chemotherapy

Due to the absence of direct comparative randomized data, the choice of initial therapy between ARPI or a taxane poses a challenge. In general terms, based on current practice guidelines (42), the majority of eligible patients will receive an ARPI and a taxane in either order. An important consideration is the diminished activity of agents in the second line, highlighting the importance of determining the ideal initial therapy. A degree of cross-resistance between docetaxel and ARPIs may exist due to a shared inhibitory effect on the AR (43-45). Retrospective clinical data indicates continued activity of docetaxel post-ARPIs with a PSA decline ≥50% of 26–35% (46-48) and phase III studies have shown a survival benefit of AAP and enzalutamide in the post-docetaxel setting (11,12). Hence, this partial cross resistance should not preclude docetaxel in subsequent lines of treatment following ARPIs or the opposite sequencing. An important consideration is that the main phase III trials of AAP and enzalutamide accrued patients with absent to mild pain, inherently a better-prognosis population, and extrapolation of this data to higher risk patients should be done with caution (7,11,12,49). Certain clinical factors, including poor response to prior ADT and visceral metastasis, are associated with worse outcomes on ARPI and prognostic indices have been developed which identify subgroups of patients with particularly poor prognosis when treated with AAP (50) or enzalutamide (51,52). Conversely, combination chemotherapy has shown high response rates in patients with a particular poor prognosis disease (53). Ultimately; it is important to define the optimal initial treatment choice in poor prognosis patients of which many will not reach next line of therapy. A recent randomized phase II cross-over trial (OZM-054) (54) compared a sequence of cabazitaxel and ARPIs (AAP or enzalutamide at investigator’s discretion) vs. the opposite sequence in patients with poor prognosis mCRPC that were ARPI-

with this hypothesis, a prospective study assessing patient reported outcomes in patients treated with AAP or enzalutamide, per investigator’s choice, demonstrated fatigue and cognitive scores were inferior for enzalutamide (40,41). Although selecting a sequence of ARPIs may be suitable for some patients, particularly those not suitable for treatment with chemotherapy, in the absence of a survival advantage or biomarkers predicting a response to a second APRI, alternating to another class of agents is preferable in most cases.
naive. Poor prognosis was defined as liver metastases, early CRPC (<12 months from ADT start to castrate resistance), and/or presence of ≥4 of 6 poor prognostic criteria (LDH > upper limit of normal (ULN), ECOG PS 2, presence of liver metastases, albumin ≤4 g/dL, Alkaline phosphatase > ULN, time from start of ADT to initiation of treatment for mCRPC <36 months) (50). The primary outcome of the trial was clinical benefit rate (CBR) defined as PSA decline ≥50%, measurable disease response and/or stable disease >12 weeks. Study accrual was terminated early in part due to slow enrollment. Ninety-five patients were enrolled and of these, approximately 50% received prior docetaxel, for either mCSPC (29% and 24% in the cabazitaxel and ARPI cohorts, respectively) or mCRPC (24% and 30%). CBR in first line treatment was greater in the cabazitaxel first arm than the ARPI arm (88.4% vs. 70%; P=0.043). All other measures including CBR in second line did not differ significantly. Of note, median overall survival was numerically longer in the cabazitaxel first arm (37 months, 95% CI: 18.9–NR) vs. the ARPIs (15.5 months, 95% CI: 12.4–NR). Although this trial cannot definitively determine whether chemotherapy or ARPI should be used up-front, the results support the view that poor prognosis disease phenotypes remain chemotherapy-responsive and may be better managed with upfront chemotherapy.

The question of optimal treatment in the third line post-docetaxel, post-ARPI setting was addressed in the CARD trial (15). In this phase III, randomized, open label trial, patients who previously received both docetaxel and one ARPI and whose disease progressed during the initial 12 months of treatment with the ARPI, were randomized in a 1:1 ratio to cabazitaxel at a dose of 25 mg/m² every 3 weeks with standard supportive measures or ARPI not previously administered (patients who previously received AAP were randomized to receive enzalutamide and vice versa). All measures, including the primary outcome, median radiographic progression free survival (8 vs. 3.7 months; HR 0.54, CI: 0.4–0.73, P<0.001), median overall survival (13.6 vs. 11 months HR 0.64, CI: 0.46–0.89, P=0.008), PSA decline ≥50% (37.5% vs. 13.5%; P<0.001) and confirmed pain response (45% vs. 19.3%) were superior in the cabazitaxel arm compared to the ARPI arm. In a post-hoc analysis, cabazitaxel maintained its superiority regardless of type of ARPI and the timing of treatment with ARPI in relation to docetaxel. Of note, the rate of serious adverse events was similar in both arms (~39%), however, adverse events leading to treatment discontinuation were more common in the cabazitaxel arm (19.8% vs. 8.9%).

These results have established the superiority of cabazitaxel over a second ARPI. The ARPI arm may have performed poorly given that short lived response to first ARPI correlate with poor response to subsequent ARPI (55), and therefore results may not be generalizable to patients with long-term response to ARPI. Nevertheless, cabazitaxel is the first treatment to demonstrate a survival benefit in third line for mCRPC, an important finding which has solidified the role of cabazitaxel in this advanced setting.

**How does this data apply to agents that have moved to earlier disease setting?**

It is now standard practice to use either docetaxel, AAP, enzalutamide or apalutamide for the majority of patients in the setting of mCSPC. Furthermore, enzalutamide, apalutamide and darolutamide, novel AR inhibitors, are now indicated for use for patients who transition through a non-metastatic castration-resistant disease state (19,20,56). Thus, many patients will have been exposed to at least one ARPI or docetaxel at the emergence of mCRPC. Analyses of time to second progression (PFS2) from the phase III SPARTAN and TITAN trials of apalutamide + ADT vs. placebo + ADT demonstrated that the benefit of early treatment with ARPI extends beyond subsequent treatments, irrespective of whether taxane or ARPI is used at progression. (19,27,57). However, these analyses did not directly compare the use of taxane vs. second ARPI post apalutamide. In practice, it is assumed that cross-resistance between agents operates in the same manner when the first is received in the mCSPC setting or for mCRPC. However, prospective data are lacking, as to whether response to subsequent therapy is identical in these 2 settings, or may differ either mechanistically or in terms of rates of response. The considerations taken in determining subsequent treatment for patients who have progressed post docetaxel/ARPIs for mCSPC are similar to those taken in prescribing second line and beyond in the pre docetaxel/ARPIs for mCSPC era. For those patients who were treated with ARPIs in the mCSPC setting, concerns of cross-resistance should guide the clinicians to favor other classes of drugs. Similar considerations could be adopted for patients who received docetaxel in the mCSPC state, however, reintroduction of docetaxel in the mCRPC setting may also be an option. This approach was commonly utilized prior to emergence of alternative potent agents for mCRPC and limited, primarily retrospective data demonstrated a PSA decline ≥50% in 25–50% of patients re-introduced to docetaxel.
(58–60). These studies showed a correlation between docetaxel free interval until progression and efficacy of docetaxel re-introduction (61). A retrospective analysis (62) of the GETUG-AFU15 phase 3 trial comparing docetaxel and ADT to ADT alone for mCSPC examined the efficacy of docetaxel re-introduction for mCRPC following docetaxel treatment in the mCSPC setting. In this trial, patients who received docetaxel for mCSPC showed a more modest response when re-introduced to docetaxel in the mCRPC setting in the first line with a PSA decline ≥50% of 20% (4/20; 95% CI: 6–44). Additional prospective data is required to determine the utility of docetaxel re-introduction as well as the appropriate sequencing of treatment after exposure to ARPIs in the mCSPC setting. Until such data is available, clinicians may be cautiously guided by sequencing trials established in the mCRPC setting as well as patient and physician preferences.

**New therapies in mCRPC**

The landscape of treatment for mCRPC is continuously evolving with a number of trials examining the efficacy of new classes of drugs ongoing and recently reported. The discovery in the past decade that 20–30% of patients with mCRPC harbour underlying genomic defect in genes involved in DNA damage repair (DDR) (63,64), prompted the investigation of PARP inhibitors as a novel therapeutic option. Phase I/II (65,66) studies demonstrated that the identification of a germline or somatic defect in homologous recombination repair (HRR) genes, a subset of genes involved in DDR, particularly BRCA2 alterations, are predictive for a response to the PARP inhibitor, olaparib. The PROfound (16), a phase III, randomized, open label study enrolled patients with mCRPC with a somatic or germline defect in genes involved in DDR with prior exposure to ARPIs, enzalutamide or AAP. Patients could have also received prior taxane chemotherapy. Patients in cohort A of the trial, harbour a BRCA1, BRCA2 or ATM defect, while those in cohort B had one of 12 other HRR defects (BRIP1, BARD1, CDK12, CHEK1, CHEK2, FANCL, PALB2, PPP2R2A, RAD51B, RAD51C, RAD51D, RAD54L). Both cohorts were randomized in a 2:1 ratio to receive either olaparib or an ARPI (AAP or enzalutamide per investigator’s choice). Cross-over at progression was permitted. Radiographic progression-free survival, the primary outcome of the trial, was superior for olaparib in cohort A (median 7.4 vs. 3.6 months, HR 0.34; 95% CI: 0.25 to 0.47; P<0.001) and in the overall study population (median 5.8 vs. 3.5 months, HR 0.49; 95% CI: 0.38 to 0.63; P<0.001). Grade 3 or higher adverse events were more common in the olaparib arm (51% vs. 38%) The most common of which were anemia (39%), nausea (36%) and fatigue/asthenia (32%). In a final analysis, overall survival in cohort A, a key secondary end point, was significantly longer in the olaparib arm than in the control arm (19.1 vs. 14.7 months, HR 0.69; 95% CI: 0.5–0.97; P=0.02). Cohort A derived the greatest survival benefit from olaparib, particularly patients with BRCA2 alterations, while there was no significant survival benefit in cohort B or the entire cohort, even after adjusting for crossover (17). This distinction between cohorts could be driven by gene specific response, however, the study was not powered to test such hypothesis. The results of this study established the efficacy of olaparib in the treatment of patients with mCRPC who harbor germline or somatic mutations in HRR, particularly for BRCA1, BRCA2 and ATM alterations. However, the trial could not determine the efficacy of olaparib in lower prevalence HRR genes which were grouped in cohort B, and further data will be required to elucidate the spectrum of alterations conferring sensitivity to PARP inhibitors. Interestingly, patients in this trial who were initially randomized to the control arm and then crossed over to the olaparib arm had a shorter lived radiographic progression free survival on olaparib when compared to those randomised to the olaparib arm (4.8 vs. 7.6 months), suggesting that olaparib treatment at an earlier line of treatment, in particular prior to a second line of ARPI, may fare better. In addition, patients with BRCA1/2 alterations who were taxane-naïve appeared to have greater survival benefit with olaparib (HR 0.3, 0.1–0.78) than those patients that were taxane-experienced (HR 0.64, 0.39–1.08). This suggests that olaparib should be preferentially used prior to taxane chemotherapy in this subgroup of patients rather than a second ARPI, irrespective of the sequence. The control arm of the PROFound trial also provides further evidence of the limited clinical activity of switching ARPI in patients already exposed to prior ARPI, as also demonstrated in the CAbD and abi-enza trials. Further investigation is required to fully define the optimal sequencing of olaparib within the treatment paradigm of mCRPC harboring HRR mutations. Additional studies examining the efficacy of other PARP inhibitors in this late stage mCSPC setting (67–69) and the utility of PARP inhibitors in combination with other active agents in earlier lines of treatment are ongoing.

Another promising therapeutic is $^{177}$Lu-PSMA-617, a prostate-specific membrane antigen (PSMA)-targeting agent bound to beta-particle emitting radioisotope
markers are investigated in ongoing studies.

AR splice variant 7 (AR-V7) is one of the well-established resistance mechanisms to treatments targeting AR in mCRPC. This variant of AR lacks the ligand binding domain and is constitutively activated. AR-V7 may be found in approximately 10–30% of men with mCRPC. There are 2 validated methods to detect AR-V7 either via nuclear protein detection (Epic) or mRNA detection in circulating tumor cells (AdnaTest) (75). Studies comparing the response to ARPIs in AR-V7(+) and AR-V7(-) mCRPC patients suggest improved response among men with AR-V7(-) mCRPC (76–78). In a study utilizing AR-V7 Epic nuclear protein detection Sharp et al. (77) demonstrated that AR-V7 is rarely expressed in primary prostate cancer. However, its prevalence increases dramatically following treatment with ADT, rising even further following treatment with abiraterone or enzalutamide, suggesting that the variant is developed under the selective pressure of AR inhibition. In this retrospective study, AR-V7 (+), docetaxel naïve patients were shown to have inferior PSA response (54% vs. 100%; P=0.03) and overall survival (25.2 vs. 74.3 months, HR 0.23, 0.07–0.79, P=0.02) in response to ARPIs when compared to AR-V7(-) docetaxel naïve patients, this is in contrast to the response to docetaxel in which case no significant differences were seen between AR-V7(+) and AR-V7(-) patients. Similarly, in the PROPHECY trial (78), AR-V7(+) tumors were associated with a shorter overall survival and progression free survival in response to ARPIs, when compared to AR-V7(-) tumors. The whole of this data suggest a potential, yet limited, role for AR-V7 as a negative predictive biomarker for the response to ARPIs in view of its low prevalence in the pre-ARPI settings. Prospective AR-V7 biomarker driven randomized clinical trials are needed to define the role of this AR splicing variant isoform in treatment selection.

Plasma circulating tumour DNA (ctDNA) has been extensively researched as a minimally invasive tool to profile mCRPC. Data from a number of groups has shown that ctDNA in mCRPC is abundant and detectable in a majority of patients, it correlates with overall tumour burden and provides a more objectively quantifiable, homogenous marker than disease burden (79-81). Within the aforementioned ABI-ENZA trial, patients provided serum for ctDNA analysis, allowing correlation of findings with outcomes, and determination of putative prognostic biomarkers in the setting of 1st line ARPI for mCRPC (79). In this analysis, the proportion of ctDNA of the plasma circulating free DNA (ctDNA%) was examined.

Biomarkers

The sequencing trials detailed above provide some indication as to how to best approach treatment choice for mCRPC. However, the endeavor to prospectively define optimal sequencing of treatment for mCRPC by comparing all potential agents, at all lines of treatment, particularly, as new agents emerge in the continuously evolving treatment landscape of mCRPC, is impractical. Predictive biomarkers offer a different approach to treatment tailoring by predicting response to different agents according to patients’ specific characteristics. Such is the case for HRR mutations and treatment with PARP inhibitors as well as 68Ga-PSMA-11 PET/CT avidity and treatment with 177Lu-PSMA-617 in mCRPC patients. Both HRR mutations and PET avidity serve as positive predictive marker for their respective treatments. Other potential predictive bio-
and the cohort was subdivided into high ctDNA% (>30%), low ctDNA% (2–30%) and undetectable ctDNA% (<2%). Higher ctDNA % correlated with worse time to progression and OS, while a ctDNA% >30% (compared with undetectable ctDNA), correlated most strongly with overall survival, independently of other clinical prognostic factors (HR 12.92, 95% CI: 5.68–29.4; P<0.001). Similar findings were shown in the OZM-054 trial, detailed above (54). Of the 95 patients enrolled in the trial, 76 (80%) had ctDNA% >2%. The most significant finding of this trial was that High baseline ctDNA% correlated with both a poorer progression free survival (HR 6.58, 95% CI: 2.95–14.69, P<0.001) and overall survival (HR 25.43, 95% CI: 3.36–190.99, P=0.002) when compared to undetectable ctDNA%, as was low ctDNA% when compared to undetectable ctDNA% (HR 3.05, 95% CI: 1.52–6.14, P=0.002 and HR 14.04, 95% CI: 1.89–104.1, P=0.01, respectively). This was maintained in a multivariate analysis, demonstrating that ctDNA% is prognostic, independent of known clinical prognostic factors including the presence of visceral metastasis, elevated lactate dehydrogenase (LDH) and elevated alkaline phosphatase (ALP). A phase II trial (NCT04015622) (82) designed to elucidate the predictive role and clinical utility of ctDNA% will compare ctDNA% guided treatment to clinician choice of treatment. Patients randomized to the biomarker guided arm with ctDNA% >2 will receive docetaxel, while patients with ctDNA% <2 will receive enzalutamide. Patients in the clinician guided arm will receive enzalutamide or docetaxel according to clinician choice. Additional findings of both the ABI-ENZA and OZM-054 studies were that somatic and/or germline gene alterations in TP53, BRCA2, ATM, RB1, AR and the PI3K pathway also correlated with worse outcomes on ARPI. Numerous studies have now demonstrated the deleterious prognostic impact of tumour suppressor loss, particularly TP53 and RB1, both in the setting of metastatic castration-sensitive disease and mCRPC (83–86). Although BRCA2 mutations have been associated with a higher rate of disease progression and metastasis for localized disease, some studies have shown conflicting results with regards to the prognostic and predictive impact of BRCA2 in mCRPC (87-90). Collectively, these findings suggest that ctDNA% as well as detection of germline and somatic genomic alterations may provide prognostic, and possibly predictive data. This concept requires prospective evaluation.

Conclusions

In the past decade the treatment paradigm of advanced prostate cancer has been substantially modified, initially by introduction of new active agents in the treatment of this disease state, followed by shift of treatment with docetaxel and ARPIs to the mCSPC setting and, more recently, the addition of ‘targeted therapy’ with olaparib for patients harboring HRR aberrations. This progress in the treatment of metastatic prostate cancer has had a positive impact on disease control and patients’ survival. However, while all of the aforementioned therapies have well established efficacy, data is lacking regarding the optimal sequencing of these drugs. In the future, new drugs such as 177Lu-PSMA-617, may continue to re-define standard of care in the mCRPC setting and will further challenge clinical decision making around selecting and sequencing therapies.

Recent head-to-head comparator studies have shed some light on the optimal sequencing of AAP, enzalutamide and taxane chemotherapy in the first, second and third line of treatment. The results of the ABI-ENZA trial indicate that the sequencing of AAP followed by enzalutamide should be favoured over the opposite sequencing due to superior second line efficacy of the former sequence. In the CARD trial cabazitaxel was superior to ARPI among those patients with short lived response to ARPI in prior line and had been treated with prior docetaxel. Currently, this data along with considerations of the side effect profile of the various agents, patient characteristics and clinician’s and patient’s preferences provide limited guidance for clinician in treatment decision. Furthermore, enrollment in a clinical trial, if eligible, should always be considered.

As the treatment landscape for advanced prostate cancer continues to evolve, comparator trials aimed at defining the optimal sequence of agents will prove impractical. Alternatively, biomarkers offer a practical potential method to guide treatment choice, particularly with the advent in ctDNA analysis. Somatic and germline mutations have been shown to be a predictive biomarker for the response to treatment with PARP inhibitors in men with mCRPC. AR-V7 status, AR gene amplification and aberrations, TP53 and RB1 defects and ctDNA% have all been suggested to have a prognostic as well as possible predictive value. Nonetheless, standardization of these tests as well as prospective validation are required in order to determine their clinical utility.
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Footnote

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