Management of patients with advanced prostate cancer: recommendations of the St Gallen Advanced Prostate Cancer Consensus Conference (APCCC) 2015


1Department of Oncology/Haematology, Kantonsspital St Gallen, St Gallen, Switzerland; 2Prostate Cancer Targeted Therapy Group and Drug Development Unit, The Royal Marsden NHS Foundation Trust and The Institute of Cancer Research, Sutton, UK; 3Department of Genitourinary Medical Oncology, MD Anderson Cancer Centre, Houston; 4Department of Genitourinary-Medical Oncology, David H. Koch Centre, The University of Texas M. D. Anderson Cancer Centre, Houston, USA; 5Department of Clinical Therapeutics, Alexandria Hospital, National and Kapodistrian University of Athens Medical School, Athens, Greece; 6Department of Cancer Medicine, Institut Gustave Roussy, University of Paris Sud, Villejuif, France; 7Department of Biostatistics and Bioinformatics, Duke University, Durham; 8Division of Human Biology, Fred Hutchinson Cancer Research Centre, Seattle; 9Tulane Cancer Centre, Tulane University, New Orleans; 10Massachusetts General Hospital Cancer Centre, Boston; 11Prostate Cancer Foundation, Santa Monica, USA; 12Research Centre for Advanced Science and Technology, The University of Tokyo, Tokyo, Japan; 13Oregon Health & Science University Knight Cancer Institute, Portland; 14Department of Medicine, Weill Cornell Medical College, New York; 15Michigan Centre for Translational Pathology, Department of Pathology; 16Department of Urology, Comprehensive Cancer Centre; 17Howard Hughes Medical Institute, University of Michigan Medical School, Ann Arbor, USA; 18Department of Oncology, Copenhagen University Hospital, Rigshospitalet, Copenhagen, Denmark; 19Monash University and Eastern Health, Eastern Health Clinical School, Box Hill, Australia; 20Cancer Research Centre, University of Warwick, Warwick, UK; 21Ludwig Boltzmann Institute for Applied Cancer Research, Kaiser Franz Josef-Spital, Vienna, Austria; 22Johns Hopkins Sidney Kimmel Cancer Center and The Brady Urological Institute, Department of Urology, Johns Hopkins University School of Medicine, Baltimore, USA; 23The Institute of Cancer Research and The Royal Marsden NHS Foundation Trust, London, UK; 24Department of Nuclear Medicine, Policlinico S. Orsola, University of Bologna, Bologna, Italy; 25Urological Sciences, Vancover Prostate Centre, University of British Columbia, Vancouver, Canada; 26Klinik und Poliklinik für Urologie, RWTH University Aachen, Aachen, Germany; 27University of Michigan Comprehensive Cancer Center, Ann Arbor, USA; 28Queen Elizabeth Hospital Birmingham, University Hospitals Birmingham, Birmingham, UK; 29Department of Radiology, Centre du Cancer et Institut de Recherche Expérimentale et Clinique (IREC), Cliniques Universitaires Saint Luc, Brussels, Belgium; 30Europa Uomo Prostate Patients, Clayhatn Ifford, UK; 31Department of Oncology-Pathology, Karolinska Institutet, Stockholm, Sweden; 32Division of Haematology and Medical Oncology, The Tisch Cancer Institute, Icahn School of Medicine at Mount Sinai, New York, USA; 33Prostate Cancer Clinical Research Unit, Spanish National Cancer Research Centre (CNIO), Madrid; 34CNIO-BIM/A Genitourinary Cancer Unit, Hospitales Universitarios Virgen de la Victoria y Regional de Málaga, Málaga; 35Centro Integral Oncológico Clara Campaña (CIOC), Madrid, Spain; 36Paul Strickland Scannor Centre, Mount Vernon Cancer Centre, Northwood; 37Prostate Cancer Targeted Therapy Group, Academic Urology Unit and Department of Diagnostic Radiology, The Royal Marsden NHS Foundation Trust and The Institute of Cancer Research, Sutton, UK; 38Institute for Precision Medicine, Meyer Cancer Center, Department of Pathology and Urology, Weill Cornell Medical College and NewYork Presbyterian, New York, USA; 39Department of Urology, Radboud University, Medical Centre, Nijmegen, The Netherlands; 40Genitourinary Oncology Service, Department of Medicine, Memorial Sloan Kettering Cancer Centre, New York; 41Department of Oncology, Assaf Harofeh Medical Centre, Tel-Aviv University, Sacker School of Medicine, Zerifin, Israel; 42Department of Urology, Carolina Urologic Research Centre, Myrtle Beach; 43Helen Diller Family Comprehensive Cancer Centre, UCSF, San Francisco, USA; 44Department of Medical Oncology, San Camillo and Forlanini Hospitals, Rome, Italy; 45Department of Urology, Toho University Sakura Medical Center, Chiba, Japan; 46Department of Medical Oncology, Dana-Farber Cancer Institute and Brigham and Women’s Hospital, Harvard Medical School, Boston, USA; 47Department of Medical Oncology and Haematology, Princess Margaret Cancer Centre, Toronto, Canada; 48Service D’Urologie, Institut de Recherche Clinique, Université Catholique de Louvain, Brussels, Belgium

Received 5 May 2015; revised 26 May 2015; accepted 28 May 2015

doi:10.1093/annonc/mdv257
Published online 3 June 2015

© The Author 2015. Published by Oxford University Press on behalf of the European Society for Medical Oncology. This is an Open Access article distributed under the terms of the Creative Commons Attribution Non-Commercial License (http://creativecommons.org/licenses/by-nc/4.0/), which permits non-commercial re-use, distribution, and reproduction in any medium, provided the original work is properly cited. For commercial re-use, please contact journals.permissions@oup.com
introduction

The diagnostic and therapeutic management of men with advanced prostate cancer (APC) has been transformed in recent years. Several new treatments for men with castration-resistant disease have successfully completed phase III trials and have received regulatory approval. The trials have not only shown a significant prolongation of overall survival, but they also reported improved or preserved quality of life (QoL). The latter, however, was less rigorously assessed and documented. Importantly, the currently approved survival-prolonging treatments in metastatic castration-resistant prostate cancer (CRPC) have distinct mechanisms of action. These therapies include unique classes of agents: taxanes, docetaxel and cabazitaxel, an immunotherapeutic agent, sipuleucel-T, novel androgen receptor (AR) pathway inhibitors abiraterone acetate (abiraterone) and enzalutamide as well as a bone targeting alpha-emitting radionuclide, radium-223 chloride (radium-223) [1–7].

Large-scale, prospective randomised trials testing the optimal sequencing of these treatments have not yet been reported. Furthermore, predictive markers to facilitate the selection of patients for a specific therapy or sequence of therapies remain an unmet need. The latter is especially relevant given the increasing evidence that some castration-resistant phenotypes may be more responsive to hormonal strategies, others to cytotoxics and others to biologic approaches. Effective combination therapies may improve outcomes and are under investigation. In addition, trials examining newer agents in the castration-naive setting are now beginning to emerge, potentially supporting the upfront use of agents such as docetaxel. While addressing one aspect of the sequencing debate, such upfront use clearly then raises a whole raft of new questions about management on development of CRPC.

Novel imaging methods for evaluation of patient selection and response claiming increased sensitivity and specificity have not been adequately tested in prospective clinical trials.

In the absence of evidence, the selection of treatment is based on clinical judgement that includes the experience of the treating physician with the available agents, the status of the disease, when the patient is presenting for treatment, and potential co-morbid conditions that might preclude a particular treatment. All treatments can have side-effects, and all of the new treatments are expensive. Of note, global access to the above-mentioned CRPC therapeutics varies across the globe, and the financial burden to an individual patient, but also to the community, must be factored.

Zoledronic acid and denosumab are approved to reduce the risk of skeletal-related events (SREs) based on the results of phase III trials in men with metastatic CRPC [8–10]. Most were approved before the availability of therapies beyond taxanes. The optimal use of these osteoclast-targeted therapies (including sequencing, initiation, frequency and duration of treatment) has not been determined. Moreover, their roles, and in particular their efficacy, in the era of newly approved anticancer-treatment options for men with CRPC [many of which also reduce the risk of SREs or symptomatic skeletal-related events (SSEs)] have not been determined. Studies to address these critical knowledge gaps need to be undertaken.

Physicians may rely on national and international guidelines to base management decisions outside of clinical trials. The level of evidence on which a specific guideline is based varies considerably. Nonetheless, there still remain several topics regarding management decisions where there is either a paucity of level one evidence or conflicting evidence of results. To address this, an international expert consensus, the St Gallen Advanced Prostate Cancer Consensus Conference (APCCC), was organised with the objective of providing the recommendations of experts to complement evidence-based guidelines and to better frame the discussion between men with prostate cancer and physicians when faced with management decisions.

Importantly, several of these recommendations were derived from clinical trial data that have been obtained from trials with specific entry criteria. These criteria will differ from trial to trial. Hence, not all recommendations can be generalised and applied uncritically to every patient, rather they should be tailored to an individual and shared decision making is still required.

methods

The panel included 41 prostate cancer experts from 17 countries, covering different specialities involved in research and treatment of men with APC (Table 1).

First, the panel members agreed upon the 10 most important areas of controversy relating to the management of men with APC, and these are listed below:

(i) Management of men with castration-naive metastatic prostate cancer
(ii) Management of men with oligometastatic castration-naive prostate cancer
In a modified Delphi process (Figure 1), questions based on the above 10 sections (Figure 2) were created and in the first round sent to all panel members for input. Questions and options for answers were later revised and sent a second time to all panel members, including all inputs received shown in an anonymised fashion, so all the panellists could see every comment from their colleagues [11]. The comments from the second round were included in the third version which was then circulated. After the third round, only important changes were accepted for the fourth and final version of the questions. This process was analogous to the one used for the St Gallen International Expert Consensus on the Primary Therapy of Early Breast Cancer [12].

For all questions, if not stated otherwise, it was assumed that any drug recommended must have been approved and was readily available, no treatment contraindications existed and no clinical trial was available. In addition recommendations applied only to non-frail patients [defined as Eastern Cooperative Oncology Group (ECOG) performance status 0–2] and for patients with adenocarcinoma of the prostate (if not stated otherwise). Importantly, in an effort to address questions from an evidenced-based and was noted to be the case with a number of the questions.

Table 1. Panel members by country and specialty

<table>
<thead>
<tr>
<th>Name</th>
<th>Country</th>
<th>Specialty</th>
</tr>
</thead>
<tbody>
<tr>
<td>Akaza, Hideyuki</td>
<td>Japan</td>
<td>Urology</td>
</tr>
<tr>
<td>Attard, Gerhardt</td>
<td>UK</td>
<td>Medical Oncology</td>
</tr>
<tr>
<td>Beer, Tomasz M.</td>
<td>USA</td>
<td>Medical Oncology</td>
</tr>
<tr>
<td>Beltran, Himisha</td>
<td>USA</td>
<td>Medical Oncology</td>
</tr>
<tr>
<td>Chinnaiyan, Arul M.</td>
<td>USA</td>
<td>Pathology/Basic research</td>
</tr>
<tr>
<td>Daugaard, Gødske</td>
<td>Denmark</td>
<td>Medical Oncology</td>
</tr>
<tr>
<td>Davis, Ian</td>
<td>Australia</td>
<td>Medical Oncology and</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Palliative Medicine</td>
</tr>
<tr>
<td>De Bono, Johann</td>
<td>UK</td>
<td>Medical Oncology</td>
</tr>
<tr>
<td>De Santis, Maria</td>
<td>Austria</td>
<td>Medical Oncology</td>
</tr>
<tr>
<td>Drake, Charles G.</td>
<td>USA</td>
<td>Medical Oncology</td>
</tr>
<tr>
<td>Eeles, Rosalind Anne</td>
<td>UK</td>
<td>Oncogenetics and Clinical/</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Radiation Oncology</td>
</tr>
<tr>
<td>Efstathiou, Eleni</td>
<td>Greece/USA</td>
<td>Medical Oncology</td>
</tr>
<tr>
<td>Fanti, Stefano</td>
<td>Italy</td>
<td>Nuclear Medicine</td>
</tr>
<tr>
<td>Fizazi, Karim</td>
<td>France</td>
<td>Medical Oncology</td>
</tr>
<tr>
<td>Gillessen, Silke</td>
<td>Switzerland</td>
<td>Medical Oncology</td>
</tr>
<tr>
<td>Gleave, Martin E.</td>
<td>Canada</td>
<td>Urology</td>
</tr>
<tr>
<td>Halabi, Susan</td>
<td>USA</td>
<td>Statistics/Epidemiology</td>
</tr>
<tr>
<td>Heidenreich, Axel</td>
<td>Germany</td>
<td>Urology</td>
</tr>
<tr>
<td>Hussain, Maha H. A.</td>
<td>USA</td>
<td>Medical Oncology</td>
</tr>
<tr>
<td>James, Nicholas D.</td>
<td>UK</td>
<td>Clinical/Radiation Oncology</td>
</tr>
<tr>
<td>Lecouvet, Frédéric</td>
<td>Belgium</td>
<td>Radiology</td>
</tr>
<tr>
<td>Logothetis, Christopher J.</td>
<td>USA</td>
<td>Medical Oncology</td>
</tr>
<tr>
<td>Nelson, Peter</td>
<td>USA</td>
<td>Medical Oncology</td>
</tr>
<tr>
<td>Nilsson, Sten</td>
<td>Sweden</td>
<td>Medical Oncology</td>
</tr>
<tr>
<td>Oh, William K.</td>
<td>USA</td>
<td>Medical Oncology</td>
</tr>
<tr>
<td>Olmos, David</td>
<td>Spain</td>
<td>Medical Oncology</td>
</tr>
<tr>
<td>Padhani, Anwar</td>
<td>UK</td>
<td>Radiology</td>
</tr>
<tr>
<td>Parker, Chris</td>
<td>UK</td>
<td>Clinical/Radiation Oncology</td>
</tr>
<tr>
<td>Rubin, Mark A.</td>
<td>USA</td>
<td>Pathology/Basic research</td>
</tr>
<tr>
<td>Sartor, Oliver A.</td>
<td>USA</td>
<td>Medical Oncology</td>
</tr>
<tr>
<td>Schalken, Jack A.</td>
<td>Holland</td>
<td>Basic research</td>
</tr>
<tr>
<td>Scher, Howard I.</td>
<td>USA</td>
<td>Medical Oncology</td>
</tr>
<tr>
<td>Sella, Avishay</td>
<td>Israel</td>
<td>Medical Oncology</td>
</tr>
<tr>
<td>Shore, Neal</td>
<td>USA</td>
<td>Urology</td>
</tr>
<tr>
<td>Small, Eric</td>
<td>USA</td>
<td>Medical Oncology</td>
</tr>
<tr>
<td>Smith, Matthew R.</td>
<td>USA</td>
<td>Medical Oncology</td>
</tr>
<tr>
<td>Sternberg, Cora N.</td>
<td>Italy</td>
<td>Medical Oncology</td>
</tr>
<tr>
<td>Suzuki, Hiroyoshi</td>
<td>Japan</td>
<td>Urology</td>
</tr>
<tr>
<td>Sweeney, Christopher</td>
<td>USA</td>
<td>Medical Oncology</td>
</tr>
<tr>
<td>Tannock, Ian</td>
<td>Canada</td>
<td>Medical Oncology</td>
</tr>
<tr>
<td>Tombal, Bertrand</td>
<td>Belgium</td>
<td>Urology</td>
</tr>
</tbody>
</table>

Figure 1. How the Consensus Process works (modified Delphi process).
clinical utility perspective, panelists were specifically instructed not to factor in cost, reimbursement and access as factors in their deliberations, although clearly these are critical factors in decision making for the individual patient.

It was recommended that if a panelist lacked experience or was non-expert for a specific question, the option ‘unqualified to answer’ (short form ‘unqualified’) should be chosen and if a panelist felt unable to vote for a best choice for any reason or had prohibitory conflicts of interest (COI), the option ‘abstain’ should be chosen. The conference also included an explicit approach to management of COI (supplementary Appendix S1, available at Annals of Oncology online).

For the purposes of this article, the term ‘recommend’ is used to reflect the fact that the panelists considered the option as the preferred one, also in the absence of hard clinical trial data. In contrast, the use of the phrase ‘discuss the option’ was used when panelists felt that the option was a consideration, but not necessarily the preferred one.

Detailed voting records for each of the questions brought to the panel are provided in the supplementary Appendix S2, available at Annals of Oncology online. In tabulating the results, the denominator was based on the number of panel members who voted, excluding those that were ‘unqualified to answer’ but including those who chose to ‘abstain’.

If ≥75% of the panelists chose the same option, this was defined as consensus. All panelists have contributed to the editing and approved this final consensus document.

Importantly, this process was also able to emphasize areas of non-consensus where additional data acquisition might be warranted.

**management of men with castration-naïve metastatic prostate cancer**

The panel members felt that ‘castration-naïve’ is the more appropriate term instead of ‘hormone-sensitive’ or ‘castration sensitive’, as the sensitivity of the cancer to castration is not known before commencement of Androgen deprivation therapy (ADT).

**intermittent and combined ADT.** ADT by means of orchiectomy, GnRH agonists or antagonists is the standard systemic treatment of metastatic prostate cancer [13–18].

The data supporting equivalence of GnRH analogues and orchiectomy were primarily established by studies using a testosterone suppression end point rather than appropriately sized and powered clinical trials with an end point of clinical efficacy. The majority of patients in economically well-developed countries today are treated with medical castration whereas, in developing countries, surgical castration is a more commonly used option for ADT [19]. Although the majority of the patients experience a profound prostate-specific antigen (PSA) decline with ADT, the median failure-free survival is ∼1 year, yet with a wide range. (11.2 months; interquartile range: 5.1–28.8) [20].

The concept of using ADT intermittently instead of continuously was developed as a consequence of several different hypotheses. These included the theory, based on experimental models [21–23], that intermittent use of ADT might be associated with a delay in development of castration resistance and the expectation that it would lead to less toxicity resulting in improvements in QoL and reduction in costs of treatment during the off phase.

Intermittent ADT (iADT) versus continuous ADT in men with metastatic prostate cancer has been evaluated in several trials, but only two trials have included solely metastatic patients and used the end point of overall survival. One of these trials was small (n = 173) whereas the SWOG 9346 trial randomised more than 1500 patients with initial PSA decline on ADT [24, 25]. The results of the latter trial failed to provide clear evidence for non-inferiority of iADT compared with continuous ADT [hazard ratio (HR) for death 1.1, 95% confidence interval (CI) 0.99–1.23]. The median overall survival (OS) was longer in the continuous ADT arm (5.8 versus 5.1 years) compared with the iADT arm. The findings serve to reject the theory of delay in castration resistance via iADT.

A recent systematic review by Niraula et al. has summarised the results of 9 studies with 5508 patients [24]. This review included men with various stages of disease, including those starting treatment of rising PSA after local treatment and trials with different end points. The authors concluded that there is evidence to recommend use of iADT (combined HR for OS = 1.02).
However, meta-analysis does not replace data from large prospective trials. Therefore, the value of iADT in men with metastatic castration-naïve prostate cancer is still controversial since the only study powered and designed to assess non-inferiority, the SWOG 9346 trial, failed to show non-inferiority of iADT compared with continuous ADT. In patients with metastatic prostate cancer achieving an adequate PSA decline (confirmed PSA fall below 4 ng/ml after ~6 months of ADT), 71% of the panellists recommended intermittent instead of continuous ADT only for a minority of selected patients.

In a clear consensus, 94% of the panellists would discuss the option of intermittent ADT in metastatic patients, 54% in the majority of patients and 40% in a minority of selected patients who achieved an adequate PSA decline.

For physicians, the interpretation of non-inferiority trials in general and non-positive non-inferiority trials in particular may be challenging especially when applied to the clinical setting and patient management. Due to the fact that residual androgen production by the adrenals may stimulate prostate cancer growth [25, 26] another attempt to improve the results of ADT treatment alone is combined (or maximal) androgen blockade (CAB), using a permanent combination of ADT and an earlier generation of AR antagonist such as bicalutamide or flutamide. Several phase III trials have evaluated the utility of front-line CAB (generally compared with the late addition of an AR antagonist to gonadal androgen suppression). Of note, most trials did not use the AR antagonist bicalutamide.

Three separate meta-analyses based on the results of these trials have concluded that there is a 3–5% overall survival advantage of CAB versus ADT alone that is statistically significant when less effective steroidal AR antagonists such as cyproterone acetate are excluded from the analysis [27–29]. Of note, one Japanese trial was positive, testing CAB with bicalutamide, raising the possibility that Asian patients may benefit more than other patients from this treatment [30, 31].

Half of the panel did not recommend CAB whereas 35% recommended it in a minority of selected patients and 15% recommended it in the majority of patients.

Considerations influencing the use of CAB include ethnicity and added toxicity. Data about CAB using a combination of GnRH analogues with newer, more potent AR pathway inhibitors are not yet available.

docetaxel in men with castration-naïve metastatic prostate cancer. ADT alone versus ADT plus docetaxel in patients with metastatic castration-naïve prostate cancer has been tested in two randomised phase III trials, which both reported improved progression-free survival but provided results which differ in respect to overall survival benefit. The French GETUG-15 trial (n = 385) reported no OS benefit [32, 33] whereas, in the US ECOG E3805 (CHAARTED) trial (n = 790), a clinically and statistically significant improvement in OS was demonstrated [34]. Both trials included a high proportion of men (>70%) who presented with de novo metastatic prostate cancer, and this patient population may not be representative of men who develop metastatic disease at some time after diagnosis of localised cancer.

The different OS results between the trials may be due to sample size, a lower percentage of patients with higher tumour volume patients in GETUG-15 (47% versus 65% in CHAARTED), differential use of subsequent life-prolonging treatments (including docetaxel) and/or geographic differences (e.g. prevalence of PSA screening).

Sixty-one percent of the panellists accepted the high-volume definition as used in CHAARTED [visceral (lung or liver) and/or 4 bone metastases, at least one beyond pelvis and vertebral column] for use in daily clinical practice, whereas 11% recommended the high-volume definition developed by SWOG [visceral (lung or liver) and/or any appendicular skeletal involvement] [35], and 14% recommended the definition of Glass [diffuse bone disease (chest, head and/or extremities) and/or visceral organ (lung or liver) involvement] [36].

Half of the panel recommended docetaxel with ADT in castration-naïve M1 patients with high-volume disease in the majority of patients, 39% in a minority. Eleven percent did not recommend docetaxel with ADT in these patients at all. In contrast, in patients with low-volume disease, 74% of the panellists did not recommend routine use of docetaxel with ADT (Figure 3A).

Of note at the time of the consensus meeting (March 2015), the voting above were based on the published data of GETUG15 and ASCO 2014 meeting presentation of CHAARTED, knowing the STAMPEDE study would read out in 2015 after the consensus meeting. As such, the consensus voting do not reflect the results of the M1 arm of the STAMPEDE study. The subsequent consensus meeting planned for 2017 will make use of the peer reviewed published data from all three studies and will aim to address issues such as patient profile, performance status, metastatic load, subsequent therapies and other related factors.

An abstract of data of four arms of the STAMPEDE trial (n = 2962 men with high-risk locally advanced or metastatic prostate cancer) was presented at ASCO 2015. Median OS was 67 months in the standard of care (ADT) arm compared with 77 months in the arm in which docetaxel was added to ADT (HR 0.76, 95% CI 0.63–0.91) [37].

osteoclast-targeted therapy in men with M1 castration-naive prostate cancer. A randomised phase III trial (CALGB 90 202; n = 645) with the bisphosphonate zoledronic acid was conducted in the castration-naïve metastatic bone setting [38]. Compared with placebo, zoledronic acid did not improve time to first SRE, the primary study end point, or overall survival. Denosumab has not been tested for reducing risk of SREs in castration-naïve prostate cancer.

There was consensus among panellists that patients with castration-naive prostate cancer and bone metastases should not receive zoledronic acid (81% of the panel) or denosumab (79% of the panel) for reducing risk of SREs (Figure 4A).

Unfortunately, the approval indications for these drugs oftentimes are not defined clearly for the castration-resistant setting and, hence, may lead to their overuse and increase the incidence of their toxicities including osteonecrosis of the jaw (and potentially life threatening) hypocalcaemia and/or hypophosphataemia.

Notably, denosumab and bisphosphonates (at lower dose or schedule than for reducing risk of SREs) have an established role for the treatment/prevention of osteoporosis/osteoporotic fractures. The panel did not review the use of osteoclast-targeted therapy for treatment/prevention of osteoporosis and osteoporotic fractures.

oligometastatic castration-naïve prostate cancer

As in other tumour types, there is growing evidence that prostate cancer patients diagnosed with a limited number of metastases (oligometastatic) may have a better prognosis compared with those with extensive metastatic recurrence [39]. A recent meta-analysis of 15 single-arm case series in patients with oligometastatic disease concluded that local treatment of metastases in this setting may be promising but that the low level of evidence does not allow its recommendation as standard of care [40].

There was consensus (85% of the panel) that the presence of ≤3 synchronous metastases (bone and/or lymph nodes) is the most meaningful definition of oligometastatic prostate cancer.

The panel addressed whether local treatment of both the primary and all evident metastases was appropriate in patients with oligometastatic disease. Sixty-two percent of the panel did not recommend using this approach instead of systemic treatment (ADT) and 38% recommended it only in a minority of selected patients. When this local therapy was considered in the context of additional short-term ADT, 62% of the panel recommended this treatment in a minority of selected patients and 27% of the panel recommended this treatment in the majority of patients.

Similarly, in the case of relapse with oligometastatic disease after radical local treatment, 58% of the panel did not recommend local treatment of all metastases instead of systemic treatment (ADT), but 39% of the panel would...
consider it in a minority of selected patients. In the context of additional short-term ADT, 27% of the panel did not recommend local treatment of all metastases, 46% of the panel recommended it in a minority of selected patients and 27% of the panel recommended this treatment in the majority of patients.

The panel recognised that the use of bone scintigraphy and computed tomography (CT), compared with newer magnetic resonance imaging (MRI) and positron emission tomography (PET)/CT imaging modalities, may result in an underestimation of lesion number. This in turn makes recommendations for a specific therapeutic approach of the oligometastatic state even more difficult [41–43].

definition of castration resistance

Over time, castration resistance has been defined in multiple ways, and with the approval of novel agents for the treatment of CRPC it is important to clarify the definition.

There was clear consensus (94% of the panel) that testosterone levels need to be measured and a specific value is required to designate a patient castration-resistant. As a consensus, 82% of the panel recommended a testosterone level <50 ng/dl (<1.7 nmol/l) as an appropriate cut-off value in daily clinical practice, which is in line with the US FDA definition.
C

<table>
<thead>
<tr>
<th>Metastatic CRPC Second-Line</th>
<th>Metastatic CRPC Third-Line</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Prospective phase III trials (post-docetaxel) 2nd line:</strong></td>
<td><strong>No prospective phase III trials for 2nd line after abiraterone, enzalutamide, radium-223 or sipuleucel-T. Options for patients with good PS:</strong></td>
</tr>
<tr>
<td>- Abiraterone</td>
<td>- Abiraterone</td>
</tr>
<tr>
<td>- Cabazitaxel</td>
<td>- Cabazitaxel</td>
</tr>
<tr>
<td>- Enzalutamide</td>
<td>- Docetaxel</td>
</tr>
<tr>
<td>- Radium-223*</td>
<td>- Enzalutamide</td>
</tr>
</tbody>
</table>

I: Do you recommend second-line treatment with abiraterone or enzalutamide in otherwise healthy patients judged to have primary (inmate) resistant disease (no PSA decline, no radiological improvement, no clinical benefit) to first-line abiraterone or enzalutamide? 

K: Do you recommend second-line treatment with abiraterone or enzalutamide in otherwise healthy patients with secondary (acquired) resistance (initial response followed by progression) to first-line abiraterone or enzalutamide 

L: Do you recommend second-line treatment with cabazitaxel in otherwise healthy patients after first-line docetaxel (prior to abiraterone/enzalutamide/radium-223)?

Consider clinical trial participation

*Bone metastases and symptomatic, no visceral or bulky lymph node metastases, not fit, unwilling to have no access to chemotherapy or post-chemotherapy

**Low tumour volume, no visceral metastases

***no visceral metastases

Figure 3. Continued

Again there was clear consensus (94% of the panel) that a confirmed (by a second value three or more weeks later) rising PSA on ADT in the presence of suppressed testosterone is sufficient for the characterisation of a castration-resistant state in clinical practice.

In case of a rising PSA on ADT, and if testosterone level is not adequately suppressed, luteinizing hormone (LH) can be measured. In case of non-suppressed LH, correct administration of the GnRH analogue should be verified.

If testosterone is not sufficiently suppressed in the presence of suppressed LH, the panel considered several next management options including bilateral orchiectomy (22%), change to alternative GnRH agonist (22%), change to GnRH antagonist (44%) or addition of an AR antagonist (9%).

**management of men with non-metastatic (M0) CRPC**

The definition of M0 prostate cancer (rising PSA on ADT and no documented metastatic disease with radiographic imaging) is dependent upon the imaging technology chosen. In current clinical practice, the ideal combination of imaging methods to define the M0 state is unclear, as is when to use it. M0 is arguably an artificial disease stage designation, as there is a high likelihood that systemic micro-metastases are missed by commonly used imaging tools (CT and bone scintigraphy). In the absence of data from positive prospective clinical trials concerning overall survival, it is unclear what treatment options should be recommended if no metastases are found radiographically and the PSA continues to rise.

Of note, the randomised trials conducted in this setting with bone-targeted therapies with the objective of delaying the onset of bone metastases were either negative or not convincingly positive [8, 44, 45]. In the placebo arm of the trial testing denosumab in this setting, time to first bone metastasis was 40.8 months in the overall population, and 26 and 18.5 months in the patients with a PSA doubling time (PSA-DT) ≤10 and ≤4 months, respectively [46].

There was clear consensus (91% of the panel) that a PSA-based trigger (level and/or kinetics) should be used for restaging asymptomatic patients with rising PSA on ADT and no known metastases.

The earlier detection of metastases with newer imaging methods (i.e. techniques other than CT and planar bone scintigraphy) and consequent earlier initiation of treatment has not been shown to be associated with a patient benefit.

There was consensus (77% of the panel) that in daily clinical practice a negative CT (thorax and abdomen and pelvis) and a negative bone scintigraphy are sufficient for diagnosis of M0 disease.

With regards to the total PSA cut-off to initiate imaging, the entire panel recommended a PSA below 20 as cut-off, almost equally divided between a PSA of between 2 and 10 (54%) and a PSA between 10 and 20 (46%). The preferred absolute value is influenced by the prior local therapy (radical prostaticectomy versus radiation therapy) and PSA-DT. For PSA-DT as a trigger for imaging, 74% of the panel recommended a PSA-DT of ≤6 months, and 9% recommended a PSA-DT of ≤3 months.

A risk-adapted strategy, taking PSA level and kinetics as well as patient preference and characteristics into consideration should be adopted.

Suggestions for when to initiate and repeat imaging in M0 CRPC patients have been recently published by the Prostate Cancer Radiographic Assessments for Detection of Advanced Recurrence Group [47].

As a trigger to initiate systemic treatment of M0 CRPC, about half of the panel (52%) recommended a combination of PSA-DT and absolute PSA value whereas 30% of the panel did not recommend treatment of patients with M0 CRPC outside of clinical trials at all, irrespective of PSA level and kinetics.
However, most of the panellists acknowledged that withholding additional treatment in a patient who knows that his PSA is rising on ADT can be challenging even without supporting data that any therapy in this stage impacts overall survival.

The treatment option for such men with M0 CRPC, outside of clinical trials, chosen by 84% of the panellists was one of the endocrine manipulations without proven OS benefit (for definition see management of men with non-metastatic (M0) CRPC section; Figure 3A).
There are no data about the benefits of abiraterone or enzalutamide in this situation, but three large randomised phase III clinical trials are ongoing and enrolment of patients with M0 CRPC in such trials is encouraged.

value of endocrine manipulations without proven survival benefit in men with metastatic CRPC

In the era before the newer AR pathway inhibitors, abiraterone and enzalutamide, were shown to improve overall survival, several drugs were used as secondary hormonal manipulation in men progressing on ADT. These drugs are considered as ‘endocrine manipulations without proven OS benefit’.

The drugs most frequently used are AR antagonists (non-steroidal including bicalutamide, flutamide, nilutamide and steroidal cyproterone acetate), oestrogens and estramustine phosphate, ketoconazole and corticosteroids (dexamethasone and prednisone/prednisolone). These agents have been tested in numerous small short-term phase II trials, demonstrating biochemical and/or clinical responses. One phase III trial (n = 260) tested AR antagonist withdrawal and ketoconazole plus hydrocortisone compared with AR antagonist withdrawal alone. The addition of ketoconazole resulted in an improvement in the response rate (PSA and objective), but not OS, although there was considerable crossover [48].

In the absence of large randomised phase III trials, the effect of these drugs on OS remains unknown. The advantages of the above-mentioned drugs are their relatively low cost, widespread access and, for some, their rather favourable safety profile.

For a patient who is not considered a candidate for chemotherapy, 52% of the panel members felt that in the era of AR pathway inhibitors with proven overall survival benefit (abiraterone and enzalutamide), these older agents are not appropriate treatments, if abiraterone and enzalutamide are available. Nevertheless, 32% would use them in a minority of selected patients and 16% in the majority of patients.

If abiraterone and enzalutamide are not available, all panel members considered it appropriate to use these agents (endocrine manipulations without proven OS benefit), 88% in the majority of patients, the remaining panel members in a minority of selected patients.

Even in countries where abiraterone and enzalutamide are officially available, not all patients have access to these agents, either because the ‘out of pocket cost’ that has to be paid by the patient is too high (e.g. United States) or because the drugs are not reimbursed in some countries.

The preferred first treatment option among these alternate endocrine agents was an AR antagonist such as bicalutamide for 63% of the panel members, while another 25% of panel members would use dexamethasone (of note, 0% would use prednisone) in this setting. Six percent of the panel recommended ketoconazole in this situation; however, availability of this agent is limited in many countries.

treatment choice and sequencing for men with metastatic CRPC

To date, six therapies have been shown to prolong survival in men with metastatic CRPC. After docetaxel became the first approved therapy to prolong survival for men with metastatic CRPC in 2004, two registration-driven (i.e. not based on disease biology) treatment ‘spaces’ for patients with CRPC evolved, defined by application of chemotherapy: pre-docetaxel (chemotherapy-naive) and post-docetaxel. Abiraterone and enzalutamide have both been tested and shown to prolong overall survival in large phase III trials in both the pre- and post-docetaxel states [3, 4, 6, 7]. Sipuleucel-T was tested predominantly (~85%) in chemotherapy-naive men with CRPC [2]. Cabazitaxel was exclusively tested in patients progressing on or after docetaxel [1]. The phase III trial with radium-223 included post-docetaxel patients (57%) or patients who were judged as unfit for chemotherapy, those who declined docetaxel, or had no access to chemotherapy [5]. The trials of abiraterone and enzalutamide excluded patients treated with the other novel AR pathway inhibitor, and abiraterone and enzalutamide were either not available or only available as part of clinical trials when the trials of docetaxel, sipuleucel-T, radium-223 and cabazitaxel were conducted. Evidence from a number of small retrospective cohort studies suggests limited activity from whichever of abiraterone and enzalutamide are used second in a sequential fashion [49–53]. Docetaxel after one or both of newer AR pathway inhibitors may have less activity than in the pivotal trials [54–56]. Cabazitaxel used in the third-line setting post-docetaxel and after one or two lines of newer AR pathway inhibitors abiraterone or enzalutamide seems to have similar anti-tumour activity when compared with that seen in the phase III trial [57–59]. It is of note that all of these studies were not only retrospective but had small patient numbers and heterogeneous patient populations and largely represented mono- or oligocenter experiences. These investigations can be considered hypothesis generating for the development of properly powered randomised, prospective trials.

Prospective phase III trial data of novel agents in the second line are only available in men with CRPC who had been treated with first-line docetaxel.

In daily practice, clinicians often face the difficult task of choosing among treatment options with different mechanisms of action, administration and toxicity profiles. Importantly, these agents have not been compared with one another prospectively. Optimal sequential use of agents with potential for survival prolongation as well as the optimal time point to initiate treatment remains uncertain.

Physicians also are challenged by the problem that the average man with CRPC might not have fulfilled the selection criteria for all of the registration trials. Also patients enrolled in clinical trials were monitored closely with bone scintigraphy and CT scans. Therefore, this raises an important question on whether the trial results may be generalised and extrapolated for a specific patient.

Sixty-three percent of the panel recommended that in patients with metastatic CRPC progressing by PSA without evidence of radiographic progression and in the absence of symptoms and imminent complications, agents with potential for survival prolongation should be initiated within 4–8 weeks. Conversely, 38% of the panel felt that in such patients treatment can be postponed in the presence of adequate disease monitoring.

The most meaningful definition of asymptomatic/mildly symptomatic (related to pain) men with metastatic CRPC was considered by 71% of the panel to be ‘No pain medication or only PRN (as needed) pain medication’. Fatigue and loss of appetite were mentioned as other important symptoms of the disease.

AR pathway inhibitors. There was consensus (88% of the panel) that abiraterone or enzalutamide are recommended as first-line therapy for otherwise healthy, asymptomatic or minimally symptomatic men with CRPC in addition to ADT (Figure 3B).

Both pivotal trials of abiraterone (COU-302) and enzalutamide (PREVAIL) in the pre-chemotherapy setting have only included asymptomatic or minimally symptomatic patients [defined as asymptomatic (score 0–1) or mildly symptomatic (score 2–3) on question 3 of the brief pain inventory short form] [6, 7]. Prior treatment with ketoconazole was not allowed for both trials.

There was consensus (77% of the panel) that it is appropriate to extrapolate the results of the COU-302 and PREVAIL trials to certain symptomatic chemotherapy-naive men with CRPC; however, 23% did not support this extrapolation. It is important to recognise that symptoms are not the only clinical factor to take into consideration when making a treatment choice (see section on predictive markers and clinically important factors for decision making in daily clinical practice).

In contrast to the PREVAIL trial, the COU-302 trial excluded patients with visceral (lung and liver) metastases.
There was consensus (88% of the panel) that it was appropriate to extrapolate the results of COU-302 to certain chemotherapy-naïve patients with visceral metastases. This may be based on the fact that abiraterone in the post-chemotherapy setting had activity in patients with visceral metastases; 12% of the panel felt that this extrapolation was not appropriate.

With regard to the preferred first-line AR pathway inhibitor, the panel was almost equally divided between abiraterone (39%), enzalutamide (27%) or either one of the two (33%).

Since there are no head-to-head trials comparing these two agents, individual patient factors such as co-medication (several drug interactions have been described for both abiraterone and enzalutamide), comorbidities as well as patient preference concerning the expected side-effects are important when making a shared choice for either of the AR pathway inhibitors (e.g. abiraterone: low potassium, fluid retention, impaired liver function and cardiac side-effects; enzalutamide: fatigue/asthenia, risk of seizures, QTc prolongation).

Based on the available retrospective data that indicate impaired activity of the AR pathway inhibitors when used sequentially, the panel did not recommend (55%) or recommended only in a minority of selected patients (42%) second-line treatment with abiraterone or enzalutamide in patients judged to have primary (innate) resistant disease (no PSA decline, no radiological soft-tissue response, no clinical benefit) to first-line enzalutamide or abiraterone (Figure 3B).

In the case of acquired resistance (initial response followed by progression) to first-line abiraterone or enzalutamide, 24% of the panel did not recommend, and 53% recommended only in a minority of selected patients, the other AR pathway inhibitor as immediate next-line treatment.

docetaxel. The panel did not (49%), or only in a minority of selected patients (42%), recommend docetaxel chemotherapy as first-line therapy for otherwise healthy asymptomatic/minimally symptomatic men with CRPC. In symptomatic patients, the panel was divided, with 41% recommending docetaxel as first-line treatment in the majority and 50% in a minority of selected patients (Figure 3B).

When an otherwise healthy symptomatic patient had a short response (<12 months) to primary ADT, the proportion changed to 56% of the panel recommending docetaxel in the majority of patients and 41% in a minority of selected patients. In contrast, in patients with a short response to primary ADT, but who are asymptomatic or minimally symptomatic, the panel did not (21%), or only in selected patients (49%), recommend docetaxel as first-line treatment.

radium-223. The panel did not recommend (33%) or recommended only in a minority of selected men with CRPC (55%) radium-223 as first-line treatment.

This may reflect the fact that the chemotherapy-naïve patient population included in the ALSYMPCA trial was mixed and not well defined: it is not possible to distinguish the groups of patients who were unfit for chemotherapy, unwilling to have chemotherapy, or who had no access to chemotherapy as these data were not collected. Furthermore, activity of radium-223 is limited to the bone environment and a significant proportion of men with CRPC have soft-tissue (nodal and/or visceral) disease (Figure 3B) [60].

Combination trials of this agent with other agents with a proven survival benefit are ongoing. Routine use of combination therapy should not be undertaken until these data are available.

Sixty-five percent of the panel felt that it was appropriate to extrapolate the results of the ALSYMPCA study to certain symptomatic men with CRPC with bone metastases who qualify as fit for chemotherapy. Furthermore, 56% of the panel felt that it was also appropriate to extrapolate the results of ALSYMPCA to certain asymptomatic patients with bone metastases, whereas 44% felt that this extrapolation was not appropriate.

sipuleucel-T. The panel was divided with regards to recommending sipuleucel-T as first-line therapy for otherwise healthy, asymptomatic men with CRPC without visceral metastases; 23% recommended it in the majority of patients, 33% only in a minority of selected patients, and 43% did not recommend it. This may be due to the fact that experience with the agent is limited to very few countries (and in fact, relatively few physicians), the fact that the agent has no detectable direct antitumor activity, and logistics are challenging. There was consensus (90% of the panel) that it is inappropriate to extrapolate the results of the IMPACT study to patients who are symptomatic and/or have visceral disease (Figure 3B).

Sipuleucel-T is currently only available in the United States and marketing authorisation in Europe was withdrawn recently.

cabazitaxel. In the second-line setting after first-line docetaxel (before abiraterone/enzalutamide/radium-223), 9% of the panel recommended cabazitaxel in a majority of patients, 55% in a minority of selected patients and 31% did not recommend it.

In third-line after second-line docetaxel (post-first-line abiraterone or enzalutamide), however, 73% of the panel recommended cabazitaxel in a majority of patients, while 24% recommended it in a minority of selected patients (Figure 3C).

staging and monitoring of treatment

baseline staging. Baseline staging and assessment of effects of anticancer-treatment of patients treated with agents with a proven survival benefit outside of clinical trials in daily clinical practice remains a challenge [61], and most current guidelines (e.g. ESMO, NCCN, EAU) do not provide clear recommendations. The Prostate Cancer Working Group 2 (PCWG2) recommendations provide clear and detailed instructions on baseline staging and treatment monitoring, but PCWG2 focused on clinical trials and was not intended as a guide for routine clinical care [62].

Laboratory and imaging parameters are subject to considerable uncertainty. Disease monitoring in the bone is especially difficult with well-described bone lesion flare phenomena both on CT and bone scans [63–65]. In addition, there are only well-defined criteria for progression on bone scans, with no specific criteria for the positive identification of benefit/response. PSA alone is not reliable enough for monitoring disease activity in advanced CRPC, since visceral metastases may develop in men without rising PSA [60].

PCWG2 recommends a combination of bone scintigraphy and CT scans, PSA measurements and clinical benefit in men with CRPC [62], while NCCN includes also MRI and PET, which are reported as ‘useful’ techniques.

Advanced MRI techniques include endorectal MRI, high magnetic field scanning (3-Tesla), high resolution $T_2$-weighted imaging, contrast enhanced MRI and diffusion-weighted imaging. Multi-parametric MRI ($T_2$-weighted imaging together with one or more of the before mentioned functional techniques) shows great promise for detecting local recurrence [66, 67]. Advanced spinal/whole-body MRI techniques are also better able to identify and gauge the extent of bone disease than planar bone scans [41, 43, 68].

PET/CT can be carried out with different tracers, enabling exploration of different features of the cancer and its interactions with bone. At present, choline PET/CT (either labelled with $^{11}$Carbon or $^{18}$Fluoride) is the approach for which there is most information, with data suggesting good accuracy for detection of recurrence [69], but validation with randomised prospective trials is still lacking. Also, solid data on new promising tracers (such as PSMA) are still lacking.

There was clear consensus with the panel recommending unanimously (100%) that imaging should be undertaken in men with metastatic CRPC before starting a new line of treatment.
Baseline examinations should include history and clinical examinations as well as baseline blood tests such as blood count (haemoglobin, thrombocytes, total white blood cell count, neutrophil and lymphocyte counts), alkaline phosphatase (ALP) and lactate dehydrogenase (LDH). Some of these variables (Hb, LDH, ALP) are considered established prognostic factors; however, their value in direct treatment decisions is not established. In addition, other assessments such as renal and liver function as well as electrolytes should be carried out.

There was clear consensus that a CT scan of the chest thorax and abdomen-pelvis (91% of the panel) and bone scintigraphy (83% of the panel) should be recommended in the majority of patients before starting a new treatment.

It is recognised that planar bone scintigraphy has short-comings and is less sensitive than other newer imaging technologies (e.g. MRI of the axial skeleton for bone staging, MRI of the whole body and/or PET/CT for concurrent bone and node metastasis screening). But availability of these newer imaging technologies is limited and their added value over bone scintigraphy in the management of APC, i.e. impact of earlier and more reliable detection of metastasis on complications and survival, has not been proven.

Malignant spinal cord compression (MSCC) occurs at a frequency of 4%–8% in men with CRPC and is one of the most devastating complications [5]. The panel stressed the need for a risk-adapted approach to undertaking MRI of the entire spine. Retrospective, small studies have shown occult spinal cord compression or impingement in up to 30% of men with CRPC [70, 71]. Extensive bone metastatic disease on bone scan was shown in both studies to be an independent predictive factor for MSCC. A large randomised phase III clinical trial comparing screening MRI of the spine to standard of care is ongoing (CRUK/11/053).

Seventeen percent of the panel recommended a baseline MRI of the entire spine in the majority of patients and 54% in a minority of selected patients based on extent of spinal metastases on bone scintigraphy or CT scan.

In case of neurologic symptoms possibly indicating MSCC, a MRI of the whole spine should be done immediately.

In routine practice, the panel did not (47%) or only in a minority of selected patients (36%) recommend the use of newer imaging methods beyond bone scintigraphy and CT scan, namely whole-body MRI and/or PET/CT, as a components of baseline prostate cancer staging.

The variability in cost and availability of newer imaging modalities throughout the world, and lack of definitive data concerning their prognostic and/or predictive value makes this area an important unmet medical need for research.

treatment monitoring. There was consensus (83% of the panel) that regular monitoring of treatment (apart from clinical and laboratory assessment) is recommended for men with metastatic CRPC. This reflects the fact that the agents with a proven overall survival benefit all have potential toxicity and considerable costs, and patients with no objective benefit (including disease stabilisation) should not be further exposed to them. It was discussed that, for imaging, generally a risk-adapted approach should be considered depending on response to therapy, extent of disease and clinical situation (e.g. line of therapy, symptoms).

The panel recommended regular measurements of ALP (88%) and LDH (61%). Both markers may be helpful as serial values when it comes to interpretation of discordant results (e.g. PSA rise and clinical improvement).

The panel stressed the need for cautious interpretation of PSA values especially in the first 2–3 months after starting a new treatment. PSA flare has been described following initiation of chemotherapy or newer hormonal therapies with significant PSA falls after initial rise [72–74].

For sipuleucel-T and radium-223 no significant PSA declines have been reported despite an OS benefit. Education of patients and physicians about uncertainties in interpretation of PSA values is critical.

Eighty-three percent of the panelists did not recommend regular monitoring or monitoring at progression with MRI of the entire spine in patients with multiple spine lesions on bone scintigraphy but rather stressed the need for a risk-adapted approach largely dependent on the development of clinical symptoms.

Further studies are necessary to confirm the potential role of systematic disease monitoring using MRI or PET/CT to characterise early response to treatment (i.e. recognition of disease response, stability or progression and not only late confirmation of progression) and its potential impact on the sequential use of agents with a proven overall survival benefits [68, 75–78].
have worsening bone pain related to non-malignant processes such as degenerative disorders or osteoporotic fractures. Similarly, fatigue/asthenia can be a side-effect of treatment and may therefore not be a sign of disease progression.

use of osteoclast-targeted agents for reducing risk of SREs and SSEs in men with CRPC

In men with CRPC and bone metastases, two agents (zoledronic acid and denosumab) have been approved for reducing risk of SREs defined as: radiation to the bone, surgery to the bone, fractures, spinal cord compression and (for zoledronic acid only) change in antineoplastic therapy. In the pivotal trial by Saad et al., zoledronic acid was given at a dose of 4 mg every 3 weeks for a total of 20 cycles (15 months) with an option to continue for an additional 9 months extension (24 months) [9, 10]. It is of note that the trial included a third arm using 8 mg every 3 weeks, which was reduced subsequently to 4 mg because of renal toxicity, and this arm did not show significant reduction of SREs compared with controls.

The dose of denosumab in the pivotal trial by Fizazi et al. was 120 mg s.c. every 4 weeks until discontinuation or until the primary cut-off date, which occurred about 41 months after start of enrolment [79]. Both zoledronic acid and denosumab were developed and investigated before the era of agents with significant anti-tumour effect and survival impact for men with CRPC. For abiraterone post-docetaxel, or enzalutamide both pre- and post-docetaxel, a significant reduction in the time to first SRE in the active treatment arms was reported [7, 80, 81].

For radium-223, also a bone targeting agent, there was also a significant prolongation in time to first SSE in the overall trial population and also in the subgroup of patients receiving bisphosphonates [82].

The optimal timing for starting treatment, optimal treatment intensity (dose and frequency) and optimal duration of osteoclast-targeted agents for men with CRPC is unclear. Also treatment-related toxicity (e.g. osteonecrosis, hypocalcaemia and hypophosphataemia) has to be taken into consideration.

Overall, 62% of the panel recommended that the majority of men with CRPC with bone metastases should receive an osteoclast-targeted agent for prevention of SRE. Thirty-two percent of the panel recommended osteoclast-targeted treatment in a minority of selected patients. If these treatments are planned there was consensus (76% of the panel) that a professional dental check (dentist) should be undertaken at baseline before treatment start in the majority of patients (Figure 4B).

The panel was almost equally divided as to preference between zoledronic acid (30%), denosumab (42%) and either of the two options (27%). This voting result may be influenced by the fact that zoledronic acid has become generic influencing the cost of the treatment.

Regarding the frequency of treatment administration for zoledronic acid or denosumab, 31% of the panel recommended dosing every 3–4 weeks, 34% recommended less frequent administration from the beginning and 28% recommended a 3–4 weekly dosing for ~2 years and less frequently after that.

This is distinct from the regulatory approvals and likely represents the lack of data supporting a dose response for these agents.

Half of the panel was of the opinion that osteoclast-targeted therapy should be continued indefinitely, whereas 47% of the panel recommended a total duration of ~2 years for reducing risk of SREs/SSEs.

For men with metastatic CRPC and bone metastases responding to a treatment, 58% of the panel were of the opinion that osteoclast-targeted therapy should be continued at the same schedule, whereas 26% voted for decreased frequency and 16% recommended interruption or discontinuation of the osteoclast-targeted therapy.

Denosumab has been compared with placebo in men with CRPC without bone metastases. Its use led to increased bone metastases-free survival but no overall survival benefit in a large phase III clinical trial [8]. No country has approved denosumab for men without bone metastases.

predictive markers and clinically important factors for decision making in daily clinical practice

The panel stressed the need for predictive markers indicating sensitivity or resistance to a specific therapy.

There was clear consensus (94% of the panel) that, at present, there is no validated and established marker that can be used as a predictive factor in daily clinical practice to inform on treatment choices for men with CRPC.

Factors favouring chemotherapy instead of AR pathway inhibitors with a proven overall survival benefit were discussed. The panel was of the opinion that a Gleason score of ≥8 (88% no) and a circulating tumour cell count of ≥5/7.5 ml (97% no) as single factors should not influence this decision. Also, in patients with extensive disease on imaging, 68% of the panel was of the opinion that this factor alone should not influence treatment choice for men with CRPC.

For several other factors, the panel was divided as to whether these factors would favour use of chemotherapy instead of abiraterone or enzalutamide: expression of AR-V7 splice variants (47% yes, 44% no), presence of visceral metastases (50% yes, 50% no), short response (<12 months) to primary ADT (53% yes, 47% no) and low PSA (<20 ng/ml) in the setting of high tumour volume (65% yes, 35% no).

In daily clinical practice, decision making is based on many patient-, tumour- and environment-related factors as well as patient and physician preference and the cost and availability of drugs.

tumour biopsy

Fresh tumour biopsies in men with CRPC are increasingly incorporated into clinical trials and advances in imaging and in laboratory technology have made the processing of biopsies from bone metastases feasible [83–87].

It is unclear when a biopsy from a man with CRPC should be considered. Tumour biopsies should only be carried out, if the procedure can be done without unreasonable risks, if adequate handling and analysis of the tissue is ensured, and if the result has potential implications for patient management. In a subset of men with CRPC, an AR signalled-independent phenotype may develop which can be associated with small-cell carcinoma seen on biopsy and clinically with rapidly progressing visceral metastases, predominately lytic bone metastatic disease and stable or low PSA levels. Recommendations from a multidisciplinary committee on aggressive variants of CRPC have been published recently [88]. The optimal treatment in these patients is unclear but platinum-based therapy is often recommended for patients with small-cell carcinoma and sometimes for patients with aggressive clinical features. There are limited data regarding the management of men with CRPC with neuroendocrine differentiation or other atypical morphology.

The panel discussed a number of clinical factors that may indicate that a tumour biopsy should be considered, low PSA (<20 ng/ml) in the setting of high tumour volume (44% yes in the majority, 47% yes in a minority, 9% no); new development of visceral metastases (32% yes in the majority, 56% yes in a minority, 12% no); progressive lesions in case of discordant tumour response to treatment (49% yes in the majority, 46% yes in a minority, 6% no); predominately lytic bone metastatic disease (37% yes in the majority, 37% yes in a minority, 26% no); patients progressing on primary ADT within <6 months (23% yes in the majority, 31% yes in a minority, 46% no).

multidisciplinary care of men with prostate cancer

Multidisciplinary care of men with APC has become much more complex in recent years. The involvement of physicians with expertise in distinct
domains of cancer care is crucial in order to achieve the best possible care for men with APC. 

Half of the panel members recommended that patients should be discussed in a multidisciplinary team (MDT) before a new line of therapy is started and 44% recommended discussion by the MDT in a majority of selected patients. Although the importance of MDTs is acknowledged, time constraints are the main reason for not discussing all but only complex patient cases in a MDT. There are also clear differences between different health systems, since in some countries discussion in a MDT is a prerequisite for insurance companies to cover the costs of the treatments.

There was strong consensus (94% of the panel) that patients should be informed about the possibility of participating in a clinical trial to improve knowledge of the disease.

Also 64% of the panelists recommended early access of men with CRPC to an expert in symptom palliation or a dedicated palliative care service.

discussion

In the absence of level one evidence or in areas where there are conflicting data or conflicting interpretation of existing data, weighted expert recommendations are helpful for making treatment decisions in daily clinical practice. That was the motivation and goal to initiate the APCCC where prostate cancer experts discussed and voted for different management options for men with APC.

In some areas, there was clear consensus (supplementary Appendix S2–S4, available at Annals of Oncology online) among the prostate cancer experts whereas, in other areas of approaches to patient care, there were divergent opinions. Some of the variation may be due to different patient management philosophies in different geographic regions; other differences may result from lack of data.

Areas of consensus included the definition of castration resistance where there was consensus that rising PSA with a documented testosterone level <50 ng/dl (<1.7 nmol/l) on ADT alone is sufficient and no secondary hormonal manipulations are required. There was also consensus that, in men with metastatic CRPC, staging examinations, including imaging, have to be carried out before each new line of treatment. Regular treatment monitoring with imaging was also recommended for men with CRPC. There was strong consensus that currently there is no validated predictive factor for treatment selection available. In men with castration-naïve prostate cancer, the use of osteoclast-targeted therapy with denosumab or zoledronic acid for reducing risk of SREs or SSES was not recommended. The same is true for men with CRPC without bone metastases. There was consensus that treatment of CRPC should generally only be stopped if two of three criteria (PSA-, radiographic or clinical progression) are fulfilled. There was strong consensus that patients should be informed about the possibility of joining a clinical trial.

The topics which generated disagreement among the panel should serve as potential areas for research and should be addressed in future clinical trials. Such areas include chemo-hormonal therapy in castration-naïve disease, but lack of consensus may be resolved by the results of the STAMPEDE trial. Another area of discrepant opinions is the treatment of M0 CRPC disease and three large phase III trials in this setting are ongoing.

In the first-line setting for men with CRPC, there seems to be considerable uncertainty. There was no consensus as to whether abiraterone or enzalutamide should be chosen, if the option of a newer AR pathway inhibitor is considered. Also the role of radium-223 in this setting seems controversial. There was not a single factor that the panel recommended to use for choosing chemotherapy first line. Biomarkers capable of informing specific applications and sequencing of treatments are of special interest for future research. Another open question is sequencing of therapies and specifically the use of abiraterone or enzalutamide after progression on the other one of them after initial response.

An important unmet medical need is the assessment of value of the newer imaging modalities in early diagnosis and treatment monitoring and their impact on patient outcome. The variability of cost and availability of these imaging modalities throughout the world are also an issue. Prospective multicentric translational studies embedding these newer modalities are needed to confirm the ability of these techniques to do better than bone scintigraphy and CT in terms of disease detection, demonstration of recurrence and to provide an earlier, more refined approach of response evaluation and defining the optimal timing to switch treatment or type of treatment. Importantly, they need to be assessed in terms of their impact on QoL measures and survival, although the latter is likely to be more challenging given the availability of multiple survival-prolonging treatments.

Furthermore, new ‘registration’ end points are needed in the light of several drugs that have been shown to prolong median OS.

Another area of uncertainty is the ‘best use’ of osteoclast-targeted agents in men with CRPC concerning initiation, dose, frequency and duration. It is also unclear if, in the time of potent new therapies, the effect of these agents is redundant or additive, and that may depend on the new therapy used. The management of men with oligometastatic disease is another area where trial data are lacking and no consensus could be achieved, which makes it an important research question. More data are necessary to address the above topics where consensus is lacking, and this conference pointed to clinical issues where physicians make decisions without certainty of benefit. If the gaps in knowledge are not filled with informative rigorous data, future regulators and payers may decide how a patient will be treated.

We did not capture differences in the opinion of panel members by geographic region, access to drugs or imaging or by different patient ethnic groups or cultures. This will be considered at the next meeting.

Additional important questions remain that have not been addressed in this meeting such as costs and cost-effectiveness of drugs, drug accessibility, ideal clinical settings, health economic issues, patient-reported QoL outcomes, implementation of precision medicine strategies in clinical practice as well as combination strategies.

The panel predicts that by the next APCCC in 2017 some of the unanswered questions will be better understood or hopefully clarified by emerging data. Additional questions will then be addressed to provide further recommendations useful for daily clinical practice: in particular selecting the right treatment, for the right patient and at the right time.

acknowledgements

We gratefully acknowledge all the participants in the Advanced Prostate Cancer Consensus Conference (APCCC) for their lively,
stimulating discussions. In addition to the panel members, we thank Beat Thürlimann for guidance in the development of the questions and for reviewing the manuscript. We also thank Thomas Cerny and the local organising committee, including Hans-Peter Schmid, Arnoud Templeton, Christian Rothermundt, Joachim Müller, Simon Wildermuth, Wolfram Jochem and Ludwig Plasswilm for their support and suggestion of speakers; Carmel Pezaro for reviewing the consensus questions; Lewis Rowett for his editorial assistance in the preparation of this report.

funding

We gratefully acknowledge the financial support of the following non-profit organisations for the Advanced Prostate Cancer Consensus Conference (APCCC): Cantonal Hospital St Gallen, City and Canton of St Gallen, Swiss Cancer Research Organisation, European School of Oncology (ESO), Swiss Cancer League, the Swiss Oncology Research Network SAKK, Swiss Cancer Foundation, Prostate Cancer Foundation (PCF) and the Günter and Regine Kelm Foundation. We thank the Movember Foundation for the sponsorship of the travel grants (no grant number).

disclosure

The full and detailed conflict of interest statements of all authors are included in supplementary Appendix S1, available at Annals of Oncology online.

references


Pulmonary neuroendocrine (carcinoid) tumors: European Neuroendocrine Tumor Society expert consensus and recommendations for best practice for typical and atypical pulmonary carcinoids

M. E. Caplin*, E. Baudin2, P. Ferolla3, P. Filosso4, M. Garcia-Yuste5, E. Lim6, K. Oberg7, G. Pelosi8, A. Perren9, R. E. Rossi1,10 & W. D. Travis11 the ENETS consensus conference participants1

1Neuroendocrine Tumour Unit, Royal Free Hospital, London, UK; 2Department of Nuclear Medicine, Endocrine Cancer and Interventional Radiology, Institut Gustave Roussy, Université Paris Sud, Villejuif Cedex, France; 3NET Center, Umbria Regional Cancer Network, Università degli Studi di Perugia, Perugia; 4Department of Thoracic Surgery, University of Torino, Torino, Italy; 5Department of Thoracic Surgery, University Hospital Valladolid, Valladolid, Spain; 6Imperial College and The Academic Division of Thoracic Surgery, The Royal Brompton Hospital, London, UK; 7Endocrine Oncology Unit, Department of Medicine, University Hospital, Uppsala, Sweden; 8Fondazione IRCCS Istituto Nazionale dei Tumori and Dipartimento di Scienze Biologiche e Cliniche Luigi Sacco, Università degli studi di Milano, Milan, Italy; 9Institute of Pathology, Surgery, University of Torino, Torino, Italy; 10Department of Thoracic Surgery, University Clinic Hospital, Valladolid, Spain; 11Imperial College and The Academic Division of Roussy, Université Paris Sud, Villejuif Cedex, France; 3NET Center, Umbria Regional Cancer Network, Università degli Studi di Perugia, Perugia; 78. Padhani AR, Makris A, Gall P et al. Therapy monitoring of skeletal metastases with whole-body diffusion MRI. J Magn Reson Imaging 2014; 39: 1049–1078.

Received 6 April 2014; revised 8 January 2015; accepted 22 January 2015

Background: Pulmonary carcinoids (PCs) are rare tumors. As there is a paucity of randomized studies, this expert consensus document represents an initiative by the European Neuroendocrine Tumor Society to provide guidance on their management.

*Correspondence to: Prof. Martyn Caplin, Neuroendocrine Tumour Unit, Royal Free Hospital, Pond Street, London NW3 2QG, UK. Tel: +44-20-7830-2727; Fax: +44-20-7372-6728; E-mail: m.caplin@ucl.ac.uk

†See the ENETS consensus group members in the Appendix section.

© The Author 2015. Published by Oxford University Press on behalf of the European Society for Medical Oncology. All rights reserved. For permissions, please email: journals.permissions@oup.com.