Darolutamide for non-metastatic, castration-resistant prostate cancer

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LAY SUMMARY

Prostate cancer is cancer of the prostate gland (a small organ in a man’s pelvis) and is the second most common cancer in the UK. There are three stages: localised, locally-advanced and advanced (or metastatic) prostate cancer. The symptoms may vary depending on the stage of cancer but can include pain, tiredness, problems emptying the bladder and the bowels. About half of men diagnosed with locally-advanced prostate cancer will see their cancer spread to other body organs (i.e. becoming metastatic). Prostate cancer growth and spread depends on the hormone, testosterone. Cancer that do not respond to hormonal treatment to reduce the level of testosterone are known as castration-resistant prostate cancer.

Darolutamide is a hormonal drug under development for castration-resistant prostate cancer that has not spread (non-metastatic). It acts by blocking testosterone receptors from getting activated on the cancer cells and consequently reducing the size of the cancer. Darolutamide is taken orally twice a day. If licenced, darolutamide will provide an additional treatment option for men with non-metastatic castration-resistant prostate cancer that are at high risk of their cancer spreading to other parts of the body.

This briefing reflects the evidence available at the time of writing. A version of the briefing was sent to the company for a factual accuracy check. The company was available to provide comment. It is not intended to be a definitive statement on the safety, efficacy or effectiveness of the health technology covered and should not be used for commercial purposes or commissioning without additional information.

This briefing presents independent research funded by the National Institute for Health Research (NIHR). The views expressed are those of the author and not necessarily those of the NHS, the NIHR or the Department of Health.
TARGET GROUP
Non-metastatic, castration-resistant prostate cancer (nmCRPC); high risk as defined by a prostate specific antigen (PSA) doubling time of ≤ 10 months – first line

TECHNOLOGY
DESCRIPTION
Darolutamide (ODM-201, BAY-1841788) is a novel nonsteroidal androgen receptor (AR) inhibitor with potential antineoplastic activity. AR antagonist darolutamide binds to ARs in target tissues; subsequently, inhibiting androgen-induced receptor activation and facilitating the formation of inactive complexes that cannot translocate to the nucleus. This prevents binding to, and transcription of, AR-responsive genes that regulate prostate cancer cell proliferation. This ultimately leads to an inhibition of growth in AR-expressing prostate cancer cells.1

Darolutamide is under development for the treatment of non-metastatic castration-resistant prostate cancer (nmCRPC).1 The medicinal product is administered orally and the proposed place in treatment is as an add-on to standard therapy.

In the currently ongoing phase III trial, darolutamide is administered twice a day in 300 mg tablets (x2) until unacceptable toxicity, confirmed metastasis or death.19

This product is not currently licenced in the EU for any indication.

Darolutamide is also at phase III in clinical trials for metastatic hormone sensitive prostate cancer.2

INNOVATION and/or ADVANTAGES
If licensed, darolutamide will offer an additional treatment option for high risk non-metastatic castration-resistant prostate cancer patients; darolutamide has the potential to increase the metastasis-free survival time.

DEVELOPER
Bayer plc.

PATIENT GROUP
BACKGROUND
Prostate cancer is the most common male cancer in the UK. It affects the prostate gland which produces some of the fluid in the semen and plays a role in urine control in men. The cancer starts in the glandular cells in the prostate and are known as acinar adenocarcinomas. It is more common in black Caribbean and black African men than in white men, and very rare in Asian men. More than half of those diagnosed are aged 70 years and over.3 Because the cancer develops slowly, people often have no signs in the early stages and symptoms only become apparent when the prostate is large enough to affect the urethra.
General symptoms of prostate cancer include: urinary tract infection, urinary frequency, sensation of incomplete emptying, haematuria (presence of blood in urine), dysuria (painful urination), haematospermia (blood in sperm), symptoms of acute kidney injury or chronic kidney disease and impotence. Various risk factors for prostate cancer have been identified, including ageing, black ethnic origin and a family history. Other factors include diet, pattern of sexual behaviour, alcohol consumption, exposure to ultraviolet radiation, chronic inflammation and occupational exposure.

There are three stages of prostate cancer; localised (confined to the prostate gland), locally-advanced (spread outside the capsule of the gland) and advanced or metastatic (spread to other body organs). At an advanced stage the cancer most commonly spreads to lymph nodes in other parts of the body or to the bones. It can also spread to other organs. About 50% of men diagnosed with localised prostate cancer will get metastatic cancer during their lifetime. Finding cancer early and treating it can lower that risk.

Prostate cancer depends on testosterone to proliferate. Castration-resistant prostate cancer (CRPC) is a stage of the disease when progression occurs despite castrate levels of testosterone, induced for example by androgen deprivation therapy (ADT). Usually, the earliest sign of resistance to ADT is a rising serum PSA level before metastases are detectable. A raised level of PSA in the blood can be a sign of a problem with the prostate.

Men at high-risk of developing metastases can be identified by a short PSA doubling time (PSADT). For patients with a PSADT ≤ 10 months, treatment is recommended, with the goal of delaying time to development of metastases.

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**Clinical Need and Burden of Disease**

The number of men diagnosed with prostate cancer has been increasing over the last 10 years. This might be because more men are having the Prostate-specific Antigen (PSA) tests and the population is getting older. In adults, prostate cancer is the second most common cancer in the UK. In men, it is the most common cancer in the UK. In 2016, 40,484 men were newly diagnosed with prostate cancer and 10,178 deaths from prostate cancer were recorded. About nine in 10 (85%) men diagnosed with prostate cancer in England and Wales survived their disease for five years or more (2010-11). Using ONS population projections, the company estimated that in 2016 in England and Wales there would have been 41,768 cases of prostate cancer. For the UK, it is estimated that 11.2% of patients progress to non-metastatic CRPC (nmCRPC) this gives an estimate of 4,590 patients in England and Wales and a total of 5,141 including the population of Scotland and Northern Ireland. The proportion of nmCRPC patients classified as high risk (PSA doubling time ≤ 10 months) is 16% (Bayer data on file). The eligible UK patient population to receive darolutamide would equate to 16% * 5,141 = 823 patients.

The Hospital Episodes Statistics for 2016/2017 recorded 75,276 finished consultant episodes (FCE), 70,295 admissions and 97,382 FCE bed days for malignant neoplasm of the prostate (ICD-10 code C61). 48,567 of these admissions were day cases.

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*a Information provided by company in UK PharmaScan record*
PATIENT PATHWAY

RELEVANT GUIDANCE

NICE GUIDANCE


NHS ENGLAND and POLICY GUIDANCE


OTHER GUIDANCE

- Gnanapragasam VJ et al. Primary radical therapy selection in high-risk non-metastatic prostate cancer. 2015
- Parker C et al. Cancer of the prostate: ESMO Clinical Practice Guidelines for diagnosis, treatment and follow-up. 2015

CURRENT TREATMENT OPTIONS

There are several treatment options available to those with non-metastatic castration-resistant prostate cancer. Many men with localised prostate cancer will not benefit from definitive treatment, and 45% of men with PSA-detected prostate cancer are candidates for deferred management (watchful waiting). In men with comorbidity and limited life expectancy, treatment of localised prostate cancer may be deferred to avoid loss of quality of life. Guidelines recommend the use of:

- Watchful waiting or observation
- Radical prostatectomy (surgical removal of the entire prostate gland between the urethra and bladder)
- External beam radiotherapy
- Brachytherapy (trans-perineal implantation of radioactive seeds into the prostate)
- Cryotherapy (local or general use of low temperatures in medical therapy)
- Hormone therapy (androgen deprivation or anti-androgens)

Treatment recommendations are dependent on the disease and patient characteristics, currently there is no standard treatment for castrated patients with rising PSA and no evidence of metastases.

### EFFICACY and SAFETY

<table>
<thead>
<tr>
<th>Trial</th>
<th>ARAMIS, NCT02200614; darolutamide vs placebo; phase III</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sponsor</td>
<td>Bayer plc</td>
</tr>
<tr>
<td>Status</td>
<td>Ongoing, recruiting</td>
</tr>
<tr>
<td>Source of Information</td>
<td>Trial registry(^{19})</td>
</tr>
<tr>
<td>Location</td>
<td>EU countries (incl UK), USA, Canada and other countries.</td>
</tr>
<tr>
<td>Design</td>
<td>Randomised, placebo-controlled</td>
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<tr>
<td>Participants</td>
<td>n=1500 (planned); aged 18 years or older; male; prostate cancer; castration-resistant with castrate level of serum testosterone; prostate-specific antigen doubling time of ≤ 10 months and PSA &gt; 2ng/ml</td>
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<tr>
<td>Schedule</td>
<td>Randomised to:</td>
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<tr>
<td></td>
<td>- Arm 1: Darolutamide (BAY1841788; ODM-201) tablets 2 x 300mg twice a day, orally</td>
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<tr>
<td></td>
<td>- Arm 2: Matching placebo tablets x 2 twice a day, orally</td>
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<tr>
<td>Follow-up</td>
<td>Until unacceptable toxicity, confirmed metastasis or death</td>
</tr>
<tr>
<td>Primary Outcomes</td>
<td>Metastasis-Free Survival (up to 72 months)</td>
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<tr>
<td>Secondary Outcomes</td>
<td>• Overall Survival (up to 72 months)</td>
</tr>
<tr>
<td></td>
<td>• Time to first symptomatic skeletal event (up to 72 months)</td>
</tr>
<tr>
<td></td>
<td>• Time to initiation of first cytotoxic chemotherapy for prostate cancer (up to 72 months)</td>
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<tr>
<td></td>
<td>• Time to pain progression (up to 72 months)</td>
</tr>
<tr>
<td></td>
<td>• Safety and tolerability (up to 72 months)</td>
</tr>
<tr>
<td></td>
<td>• Changes in health-related QoL are assessed by Functional Assessment of Cancer Therapy-Prostate (FACT-P), European Organization for Research and Treatment of Cancer Quality of life Questionnaire – Prostate Cancer Module (EORTC-QLQ-PR25) and European QoL 5-domain scale (EQ-5D-3L) questionnaire.(^b)</td>
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<tr>
<td>Key Results</td>
<td>-</td>
</tr>
<tr>
<td>Adverse effects</td>
<td>-</td>
</tr>
<tr>
<td>Expected reporting date</td>
<td>Study primary completion date 16(^{th}) April 2018, study estimated completion date 30(^{th}) June 2020.</td>
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</tbody>
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\(^b\) Information provided by company
**ESTIMATED COST and IMPACT**

**COST**

The cost of darolutamide is not yet known.

The following anti-androgens are currently marketed and used in the prostate cancer broader population in the UK:\textsuperscript{20,21,22}

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dose</th>
<th>Unit cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Xtandi (enzalutamide) 40 mg capsules</td>
<td>160 mg once daily</td>
<td>£24.40</td>
</tr>
<tr>
<td>Casodex (bicalutamide) 150mg tablets</td>
<td>150 mg once daily</td>
<td>15p</td>
</tr>
</tbody>
</table>

**IMPACT – SPECULATIVE**

**IMPACT ON PATIENTS AND CARERS**

☐ Reduced mortality/increased length of survival

☐ Reduced symptoms or disability

☐ Other: *improved quality of life for patients*

☐ No impact identified

**IMPACT ON HEALTH and SOCIAL CARE SERVICES**

☐ Increased use of existing services

☐ Decreased use of existing services

☐ Re-organisation of existing services

☐ Need for new services

☐ Other

☒ None identified

**IMPACT ON COSTS and OTHER RESOURCE USE**

☐ Increased drug treatment costs

☐ Reduced drug treatment costs

☐ Other increase in costs

☒ Other reduction in costs
☐ Other: uncertain unit cost compared to existing treatments

☐ None identified

OTHER ISSUES

☒ Clinical uncertainty or other research question identified: trial results not yet published

☒ None identified

REFERENCES


