INTRODUCING



30-DAY FREE SAMPLE PROGRAM

Start your appropriate patients on treatment right in your office

The NUBEQA Free Sample
Program may allow you
to start your patients on
NUBEQA quickly

Please read the following pages for important information for prescribers, patients, and pharmacies.



INDICATION

NUBEQA® (darolutamide) is an androgen receptor inhibitor indicated for the treatment of patients with non-metastatic castration-resistant prostate cancer.

IMPORTANT SAFETY INFORMATION

Embryo-Fetal Toxicity: Safety and efficacy of NUBEQA have not been established in females. NUBEQA can cause fetal harm and loss of pregnancy. Advise males with female partners of reproductive potential to use effective contraception during treatment with NUBEQA and for 1 week after the last dose.

Please see additional Important Safety Information throughout and full **Prescribing Information**.

START TREATMENT WITH THE NUBEQA® 30-DAY FREE SAMPLE PROGRAM

The NUBEQA Free Sample Program will give you the opportunity to

- Start your patients on NUBEQA right in your office
- Provide your patients with a 30-day supply of NUBEQA at no cost

Ordering a sample is simple

- Contact your sales representative, who will process your sample request
- Your order will be shipped within 24 to 48 hours



For illustrative purposes only.

IMPORTANT SAFETY INFORMATION (cont'd)

Adverse Reactions

Serious adverse reactions occurred in 25% of patients receiving NUBEQA and in 20% of patients receiving placebo. Serious adverse reactions in $\geq 1\%$ of patients who received NUBEQA were urinary retention, pneumonia, and hematuria. Overall, 3.9% of patients receiving NUBEQA and 3.2% of patients receiving placebo died from adverse reactions, which included death (0.4%), cardiac failure (0.3%), cardiac arrest (0.2%), general physical health deterioration (0.2%), and pulmonary embolism (0.2%) for NUBEQA.

How to dispense the sample

- Identify patients who are appropriate for NUBEQA
- Give the appropriate patients both the sample and the patient brochure
- Patient starter kits are also available from your sales representative



After dispensing the sample, initiate the prescription process for your patients

 Visit <u>NUBEQAhcp.com</u> to download the patient request form



- Use <u>covermymeds.com</u>
- Access Services by Bayer[™] can help you with reimbursement, access, benefit verification, prior authorization assistance, and appeal support. Please call 1-800-288-8374 Monday-Friday, 9:00 AM-6:00 PM (ET)

IMPORTANT SAFETY INFORMATION (cont'd)

Adverse Reactions (cont'd)

Adverse reactions occurring more frequently in the NUBEQA arm (\geq 2% over placebo) were fatigue (16% vs 11%), pain in extremity (6% vs 3%) and rash (3% vs 1%).

Clinically significant adverse reactions occurring in ≥2% of patients treated with NUBEQA included ischemic heart disease (4.0% vs 3.4% on placebo) and heart failure (2.1% vs 0.9% on placebo).

Please see additional Important Safety Information throughout and full Prescribing Information.



NUBEQA® Free Sample Program Requirements

- The NUBEQA Free Sample Program provides NUBEQA at no cost for up to 30 days to eligible patients
- The NUBEQA Free Sample Program is only valid in the United States and Puerto Rico and is void where restricted or prohibited by law
- Qualifying prescriptions for the NUBEQA Free Sample Program must be for an on-label use of NUBEQA
- Participation in the NUBEQA Free Sample Program is not contingent on any past, present, or future prescriptions for, or purchase of, NUBEQA or any other Bayer product. No purchase or refills are required
- If applicable, patient is responsible for any taxes or costs associated with shipment of product
- The program is available to all patients, irrespective of drug coverage
- Bayer reserves the right to rescind, revoke, or amend the NUBEQA Free Sample Program at any time for any reason without prior notice
- By enrolling in the NUBEQA Free Sample Program, the prescribing physician agrees to these requirements
- Bayer cannot ensure continuity of care after the initial sample is dispensed for Medicare Part D patients who may experience co-pay and/or reimbursement challenge with their federally funded insurance program

IMPORTANT SAFETY INFORMATION (cont'd)

Drug Interactions

<u>Effect of Other Drugs on NUBEQA</u> – Combined P-gp and strong or moderate CYP3A4 inducers decrease NUBEQA exposure, which may decrease NUBEQA activity. Avoid concomitant use.

Combined P-gp and strong CYP3A4 inhibitors increase NUBEQA exposure, which may increase the risk of NUBEQA adverse reactions. Monitor more frequently and modify NUBEQA dose as needed.

No-cost Supply of NUBEQA Requirements

- The no-cost supply of NUBEQA is not insurance
- The no-cost supply of NUBEQA may not be billed in whole or part to any patient or insurer
- The no-cost supply of NUBEQA may not be sold, purchased, or traded
- By accepting the no-cost product, the patient agrees to these requirements

IMPORTANT SAFETY INFORMATION (cont'd)

Drug Interactions (cont'd)

<u>Effects of NUBEQA on Other Drugs</u> – NUBEQA inhibits breast cancer resistance protein (BCRP) transporter. Concomitant use increases exposure (AUC) and maximal concentration of BCRP substrates, which may increase the risk of BCRP substrate-related toxicities. Avoid concomitant use where possible. If used together, monitor more frequently for adverse reactions, and consider dose reduction of the BCRP substrate.

NUBEQA inhibits OATP1B1 and OATP1B3 transporters. Concomitant use may increase plasma concentrations of OATP1B1 or OATP1B3 substrates. Monitor more frequently for adverse reactions and consider dose reduction of these substrates.

Review the prescribing information of drugs that are BCRP, OATP1B1, and OATP1B3 substrates when used concomitantly with NUBEQA.

Please see additional Important Safety Information throughout and full Prescribing Information.





Please see Important Safety Information throughout and full <u>Prescribing Information</u>.



