## Available under PEI Pharmacare when patients meet Special Authority criteria ${ }^{1}$

## Prolia ${ }^{\circledR}$ is indicated: ${ }^{2}$

- For the treatment of postmenopausal women with osteoporosis at high risk for fracture, defined as a history of osteoporotic fracture, or multiple risk factors for fracture; or patients who have failed or are intolerant to other available osteoporosis therapy. In postmenopausal women with osteoporosis, Prolia reduces the incidence of vertebral, nonvertebral and hip fractures.
- As a treatment to increase bone mass in men with osteoporosis at high risk for fracture, defined as a history of osteoporotic fracture, or multiple risk factors for fracture; or patients who have failed or are intolerant to other available osteoporosis therapy.
- As a treatment to increase bone mass in men with nonmetastatic prostate cancer receiving androgen deprivation therapy (ADT), who are at high risk for fracture.
- As a treatment to increase bone mass in women with nonmetastatic breast cancer receiving adjuvant aromatase inhibitor (Al) therapy, who have low bone mass and are at high risk for fracture.
- As a treatment to increase bone mass in women and men at high risk for fracture due to sustained systemic glucocorticoid therapy.
- As a treatment to increase bone mass in women and men at high risk for fracture who are starting or have recently started long-term glucocorticoid therapy.

Consult the Product Monograph at www.amgen.ca/Prolia_PM.pdf for important information relating to contraindications, warnings, precautions, adverse reactions, interactions, dosing and conditions of clinical use. The Product Monograph is also available by calling us at 1-866-502-6436.


## CRITERIA: ${ }^{1}$

For the treatment of osteoporosis in postmenopausal women who were previously approved or would otherwise be eligible for coverage of oral bisphosphonates and who:

1. Have experienced a further significant decline in bone mineral density (BMD) after 1 year of continuous bisphosphonate therapy and meet at least TWO of the following:

- Age >75 years old
- Prior fragility fracture
- BMD T-score $\leq-2.5$

OR
2. Have a contraindication to bisphosphonates due to hypersensitivity or abnormalities of the esophagus (e.g. esophageal stricture or achalasia) and have at least TWO of the following:

- Age >75 years old
- Prior fragility fracture
- BMD T-score $\leq-2.5$

NOTE: Hypersensitivity or abnormalities are defined as esophageal ulceration, erosion or stricture, or lower gastrointestinal symptoms severe enough to cause discontinuation of oral bisphosphonates, or swallowing disorders that will increase the risk of esophageal ulceration from oral bisphosphonates.

In all cases, patients receiving Prolia (denosumab) must not be receiving concomitant bisphosphonate therapy.

The recommended dose of Prolia (denosumab) is a single subcutaneous injection of 60 mg , once every 6 months.

For full details regarding coverage, visit www.healthpei.ca/formulary.

PAAB

## PEI Form - "How To"

## Health PEI

## SPECIAL AUTHORIZATION REQUEST

 STANDARD SPECIAL AUTHORIZATIONFax requests to (902) 368 -4905 OR mail requests to PEI Phammacare, P.O. Box 2000, Charloteltown, PE, C1A TNB

## SECTION 1 - PATIENT INFORMATION



SECTION 2 - PRESCRIBER INFORMATION


SECTION 3 - MEDICATION DETAIL INFORMATION
REQUESTED DRUG (PLEASE PRINT)


REASON FOR REQUEST (PLEASE EXPLAIN)

- Contraindicioion
$\square$ Adverse Event
$\square$ Other
OTHER COMMENTS, INCLUDING COPIES OF CUITURE \& SENSITIVITY REPORTS FOR ANTIBIOTIC REQUESTS, COPIES OF RELEVANT TEST RESULTS, AND RELEVANT ADVICE RECEVED FROM CONSULTANTS/SPECILLLSTS (IF APPLICABLE) Pince Edward Island's Freedom of fltormation \& Protection of Privacy (FOIPP) Act as itrelates directiy to and is necessay for providing sevices under the PEI High-Cost Drugs Program.
fyou have any questions about this collection of personal information, you may contact the program office at $902-368-4947$ or at the address at the top of the fom
PRESCRIBER SIGNATURE (REQUIRED)
DATE

Age: If older than 75 years, the patient meets one of the 3 additional criteria.

If older than 75 years of age, the patient must indicate that EITHER the T-score is $\leq-2.5$ or they have had a prior fragility fracture.
Otherwise patients MUST meet BOTH of these additional criteria.

Form MUST indicate that the patient has experienced a significant decline in bone mineral density (BMD), in which case bisphosphonate use is contraindicated due to hypersensitivity or patient having abnormalities of the esophagus.

