Available for both men and postmenopausal women with osteoporosis under the New Brunswick Prescription Drug Program via Special Authorization¹

Prolia® 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 (AI) 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 and in men who meet the following criteria:¹

• Have a contraindication to oral bisphosphonates;

AND

- High risk for fracture*, or refractory or intolerant to other available osteoporosis therapies.
- * High fracture risk is defined as:
- A moderate 10-year fracture risk (10% to 20%) as defined by the Canadian Association of Radiologists and Osteoporosis Canada (CAROC) tool or the World Health Organization's Fracture Risk Assessment (FRAX) tool with a prior fragility fracture; or
- A high 10-year fracture risk (\geq 20%) as defined by either the CAROC or FRAX tool.



References: 1. New Brunswick Drug Plans Formulary Bulletin #965, January 22, 2018. Accessed February 7, 2018. http://www2.gnb.ca/content/dam/ gnb/Departments/h-s/pdf/en/NBDrugPlan/FormularyUpdates/NBDrugPlansBulletin965.pdf. **2.** Prolia Product Monograph. Amgen Canada Inc., June 25, 2019.

New Brunswick Form – "How To"



New Brunswick Prescription Drug Program (NBPDP) SPECIAL AUTHORIZATION REQUEST FORM

Please complete all required sections to allow your request to be processed without delay. This form must be completed by a Prescriber

Date: DD/MM/YYYY							
	F	PATIENT INFO	ORMATION				
Patient's Last Name:			First:		MI:		
Medicare or NBPDP ID Number:			Date of Birth: DD/MM/YYYY				
Street address:							
P.O. Box:	City:		Postal C		Postal Co	ode:	
		DRUG REQ	UESTED				
Drug Name/Strength/Form:		Dosage Schedule:		Expe	Expected Duration of Therapy:		
Relevant Previous Drug T	herapies:						
Other Relevant Informatio	n:						,
REQUESTOR INFORMATION				PLEAS	SE RETUR	N FORM TO:	
Requestor Address:		e Number: 'SNB, NANB, NBF	PhS, etc.)	P.O. Box 644 Main Moncton, Inquiry Li	690, Street, NB E1C 8 ine: 1-800-3	332-3691	
Requestor signature:					506-867-48 ax: 1-888-4		

The information collected, used and disclosed by this request is collected, used and disclosed pursuant to section 4(4) and 4.1of the New Brunswick Prescription Drug Payment Act. If you have any questions please contact 1-800-332-3691. For use in male patients with osteoporosis or female patients with postmenopausal osteoporosis in whom bisphosphonate use is contraindicated and where the patient is at high risk of fracture, or refractory or intolerant to other available osteoporosis therapies.

Form **MUST** indicate previous therapies that the patient was refractory to or could not tolerate.

Form **MUST** indicate that bisphosphonate use is contraindicated.

