

MEDICARE FORM

Prolia®, Xgeva® (denosumab) Injectable Medication Precertification Request

Page 1 of 3

(All fields must be completed and legible for precertification review.)

For Ohio MMP:

FAX: 1-855-734-9389 **PHONE:** 1-855-364-0974

For other lines of business:

Please use other form.

Note: Xgeva is non-preferred. The preferred products are pamidronate or zoledronic acid. Pamidronate and zoledronic acid do not require precertification.

Please indicate: Start of treatment: Start date: //			☐ Continuation of therapy: Date of last treatment//					
Precertification Requested By:			Phone:		Fax:			
A. PATIENT INFORMATION								
First Name:	Last Name:				DOB:			
Address:		City	:		State:	ZIP:		
Home Phone: Work Phone:		Cell	Phone:		Email:			
Current Weight: lbs or kgs H	leight: inches or	cm	ns Allergies:		1			
B. INSURANCE INFORMATION	ÿ <u>——</u> —		3					
			other coverage?					
Group #:		If yes, provide ID#: Carrier Name:						
Insured: Insured:								
C. PRESCRIBER INFORMATION	'							
First Name:	Last Name:	(Check or			e): ☐ M.D. ☐ D.O. ☐ N.P. ☐ P.A.			
Address:		City	:		State:	ZIP:		
Phone: Fax:	St Lic #:	NPI	#:	DEA #:	UPIN	N :		
Provider E-mail:	Office Contact Name:	ı		Phone:				
D. DISPENSING PROVIDER/ADMINISTRATION	INFORMATION							
Place of Administration:			Dispensing Provider	Pharmacy:				
☐ Self-administered ☐ Physician's Office	e		☐ Physician's Office		Retail Pharmac			
<u> </u>			☐ Specialty Pharmac	у 🗆	Other:			
Center Name:			Name:					
☐ Home Infusion Center Phone: Agency Name:			Address:					
Administration code(s) (CPT):			Phone:		Fax:			
Address:			TIN:		NPI:			
E. PRODUCT INFORMATION								
E. PRODUCT INFORMATION								
Request is for: Prolia Xgeva Dose:	Frequen	су:		H(CPCS Code:			
					CPCS Code:			
Request is for: Prolia Xgeva Dose: _	e primary ICD code and s	ecify	any other where appli	cable.				
Request is for: Prolia Xgeva Dose: F. DIAGNOSIS INFORMATION - Please indicate Primary ICD Code: S	e primary ICD code and s Secondary ICD Code:	pecify	any other where applic	cable. ner ICD Code	:			
Request is for: Prolia Xgeva Dose: F. DIAGNOSIS INFORMATION - Please indicate	e primary ICD code and s Secondary ICD Code: _ I information must be com	pleted	any other where applic	cable. ner ICD Code	:			
Request is for: Prolia Xgeva Dose: F. DIAGNOSIS INFORMATION – Please indicate Primary ICD Code: SG. CLINICAL INFORMATION – Required clinical For All Requests: (Clinical documentation r. Note: Xgeva is non-preferred. The preferred produ	e primary ICD code and specondary ICD Code: Information must be come required for all request are pamidronate or zo	pleted s)	any other where application of the contract of	cable. ner ICD Code recertification	: requests.			
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Request is for: Prolia Xgeva Dose: F. DIAGNOSIS INFORMATION – Please indicate Primary ICD Code: S. G. CLINICAL INFORMATION – Required clinical For All Requests: (Clinical documentation r. Note: Xgeva is non-preferred. The preferred produ Yes No Has the patient had prior therapy wit Yes No Has the patient had a trial and failure Please explain if there are any other medical reasons. Please provide the patient's Bone Mineral Density (B. Please indicate the location the BMD was measured. Yes No Is the patient receiving 1000mg of Ca. Yes No Will the patient have clinical evider. Yes No Will the patient be using denosumable Yes No Will the patient be using Prolia in color Yes No Is the patient at high risk for fracture. Yes No Has the patient had an osteoporotic Yes No Does the patient he Please explain (see Theumatoid article glucocorticoid of Yes No Does the patient have a high FRAX.	Exprimary ICD code and specific condary ICD Code: Information must be come required for all request acts are pamidronate or zo the Xgeva (denosumab) with e, intolerance, or contrainding (s) that the patient cannot use a compart of the Xgeva (denosumab) with e, intolerance, or contrainding (s) that the patient cannot use a compart of the Agent and 400 international conditions and 400 international conditions with Xgeva? The Agent Age	pleted s) ledron in the leation se par ed: T-se par spiral units ing hypenous leation fracture hol inta moking	other where application of the inits entirety for all pricacid. Pamidronate a last 365 days? to pamidronate or zoled midronate or zoled midronate.	er day □ pare	:requests. acid do not recommend and provide the commendation of the creased fall risk to fracture risk ≥	uire prece	ertification.	
Request is for: Prolia Xgeva Dose: F. DIAGNOSIS INFORMATION – Please indicate Primary ICD Code: S. G. CLINICAL INFORMATION – Required clinical For All Requests: (Clinical documentation r. Note: Xgeva is non-preferred. The preferred production Yes No Has the patient had prior therapy with Yes No Has the patient had a trial and failure Please explain if there are any other medical reasons. Please provide the patient's Bone Mineral Density (B. Please indicate the location the BMD was measured. Yes No Is the patient receiving 1000mg of Compared Yes No Will the patient be using denosumable Yes No Will the patient be using Prolia in compared Yes No Is the patient at high risk for fracture Yes No Has the patient had an osteoporotic Yes No Does the patient had an osteoporotic Yes No Does the patient had Indicated Information Info	BMD) score and date obtain (s) that the patient cannot use in combination with Xgeva? BMD) score and date obtain (s) that the patient cannot use in combination with Xgeva? BMD) score and date obtain (s) that the patient cannot use in combination with intraverse in combination with Xgeva? BMD) score and date obtain (s) that the patient cannot use in combination with Xgeva? BMD) score and date obtain (s) that the patient cannot use in combination with Xgeva? BMD) score and date obtain (s) that the patient cannot use in combination with Xgeva? BMD) score and date obtain (s) that the patient cannot use in combination with Xgeva? BMD) score and date obtain (s) that the patient cannot use in combination with Xgeva? BMD) score and date obtain (s) that the patient cannot use in combination with Xgeva? BMD) score and date obtain (s) that the patient cannot use in combination with Xgeva? BMD) score and date obtain (s) that the patient cannot use in combination with Xgeva? BMD) score and date obtain (s) that the patient cannot use in combination with Xgeva? BMD) score and date obtain (s) that the patient cannot use in combination with Xgeva? BMD) score and date obtain (s) that the patient cannot use in combination with Xgeva? BMD) score and date obtain (s) that the patient cannot use in combination with Xgeva? BMD) score and date obtain (s) that the patient cannot use in combination with Xgeva? BMD) score and date obtain (s) that the patient cannot use in combination with Xgeva? BMD) score and date obtain (s) that the patient cannot use in combination with Xgeva? BMD) score and date obtain (s) that the patient cannot use in combination with Xgeva?	pleted s) ledron in the leation se par ed: T-se par spiral units ing hypenous leation fracture hol inta moking r major month	other where application of the inits entirety for all pricacid. Pamidronate a last 365 days? to pamidronate or zoled midronate	er day □ pare of frailty □ incoment with denoted the pare of the	:requests. acid do not recommendate Date: tify: ental history of hereased fall risk of fracture risk ≥ psumab?	uire prece	ertification.	



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB					
G. CLINICAL INFORMATION (continued	/) – Required clinical information must be	completed in its entirety for all pr	ecertification requests					
For Prolia Requests:	1							
Post-menopausal osteoporosis								
Please select which of the following medication	n(s) was ineffective, not tolerated or contra	aindicated:						
Select all that apply: Alendronate (Binosto								
	I, Actonel with Calcium or Atelvia)		ronic acid (Zometa, Reclast)					
	☐ Raloxifene (Evista) ☐ Tamoxifen (Nolvadex/Soltamox) ☐ Toremifene citrate (Fareston)							
	Other: Please identify: Prevention or treatment of osteoporosis in patients receiving endocrine therapy for breast cancer							
☐ Yes ☐ No Is the patient receiving endoc		. Diodot odiloo.						
	e following endocrine therapy (aromatase in	hibitors) is being used:						
	☐ exemestane (Aromasin) ☐ letrozole (I							
☐ Yes ☐ No Is there documentation that the								
	the medication trial: Continued bone los							
	nge://							
	nge: / / - / /							
Yes No Is there documented evidence		•						
Yes No Is there documented evidence that the patient has a contraindication to bisphosphonates?								
Please select which of the following bisphosphonates was ineffective, not tolerated or contraindicated: Select all that apply: Alendronate (Binosto, Fosamax or Fosamax plus D) Etidronate disodium (Didronel) Ibandronate (Boniva)								
	Risedronate (Actonel, Actonel with Calcium or Atelvia) Zoledronic acid (Zometa, Reclast)							
☐ Other: Please identify	y:							
Treatment to increase bone mass in men re								
Yes No Does the patient have prosta								
Yes No Is the patient receiving andro								
Treatment of bone loss in men with osteop ☐ Yes ☐ No Is there documentation that the		phonato trial of at least 1 year durat	ion?					
	the medication trial: Continued bone los							
	nge:/ / _// /		_					
	nge:/ / // /							
Yes No Is there documented evidence		bhosphonates?						
Yes No Is there documented evidence								
Please select which of the following bisphosphonates was ineffective, not tolerated or contraindicated:								
Select all that apply: Alendronate (Binosto, Fosamax or Fosamax plus D) Etidronate disodium (Didronel) Ibandronate (Boniva)								
Risedronate (Actonel, Actonel with Calcium or Atelvia) Zoledronic acid (Zometa, Reclast)								
Other: Please identify								
Treatment of glucocorticoid-induced osteo ☐ Yes ☐ No Is the patient initiating or con-		losago oguivalent to 2.5 mg or groot	or of produisons for 3 months					
or more?	inding systemic glucocorticolds at a daily d	losage equivalent to 2.5 mg of great	er of predifisorie for 5 months					
Please select: ☐ initiating s	systemic glucocorticoids							
	ent expected to remain on glucocorticoids fo							
☐ Yes ☐ No Is there documentation that the								
	the medication trial: Continued bone los							
Bisphosphonate #1 Date ra	nge:							
	nge: / / - / /							
Yes No Is there documented evidence	·	•						
Yes No Is there documented evidence Please select which of the following bisphosp	•	• •						
Select all that apply: Alendronate (Binosto			onate (Boniva)					
	I, Actonel with Calcium or Atelvia) ☐ Zole		,					
Other: Please identif	ý:	· · · · · · · · · · · · · · · · · · ·						

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB
G. CLINICAL INFORMATION	(continued) - Required clinical information	n must be completed for ALL prece	ertification requests.
For Xgeva Requests:			
Bone metastases from solid tu	mors		
Please indicate which of the follo	owing pertains to the patient: 🗌 Bladder cance	er 🔲 Breast cancer 🔲 Kidney cai	ncer 🔲 Ovarian cancer
	☐ Non-small cel	Il lung cancer ☐ Prostate cancer	☐ Thyroid cancer
	☐ Other: Please	e specify:	
☐ Giant cell tumor of the bon	е		
1 —	ed events in patients with multiple myelom	a	
Treatment of hypercalcemia of			
Yes No Has the patient	been treated with intravenous bisphosphonate	e therapy?	
	e the date range of therapy://		
	emia of malignancy refractory to intravenous t	oisphosphonate therapy?	
	n-corrected serum calcium level been tested?		
→ Please provide	e the albumin-corrected serum calcium level: _	mg/dL Date:/	
For Continuation Requests	: (Clinical documentation required for	all requests)	
	t have a hypersensitivity to denosumab?		
<u> </u>	ponse the patient has experienced while on de	enosumab: No response Min Significant improveme	
H. ACKNOWLEDGEMENT			
Request Completed By (Si	gnature Required):		Date://
any insurance company by pr		eals material information for the p	with the intent to injure, defraud or deceive urpose of misleading, commits a fraudulent

The plan may request additional information or clarification, if needed, to evaluate requests.