

MEDICARE FORM

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

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

For Illinois MMP:

FAX: 1-855-320-8445 **PHONE:** 1-866-600-2139

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	<i>r</i> :		State:	ZIP:	
Home Phone: Work Phone:		Cel	I Phone:		Email:	I	
Current Weight: lbs or kgs He	eight: inches or	cr	ns Allergies:		l		
B. INSURANCE INFORMATION	<u> </u>		J				
			coverage?	No			
Group #:		If yes, provide ID#: Carrier Name:					
Insured:	Insured:						
C. PRESCRIBER INFORMATION							
First Name:	Last Name:			(Check on	e): 🗌 M.D. 🔲 I	D.O. 🗌 N	l.P. 🗌 P.A.
Address:		City	r:		State:	ZIP:	
Phone: Fax:	St Lic #:	NP	#:	DEA #:	UP	IN:	
Provider E-mail:	Office Contact Name:	I		Phone:	•		
D. DISPENSING PROVIDER/ADMINISTRATION	INFORMATION						
Place of Administration:			Dispensing Provider/Pharmacy:				
☐ Self-administered ☐ Physician's Office			☐ Physician's Office] Retail Pharma	-	
			☐ Specialty Pharmac	-	Other:		
Center Name:			Name:				
☐ Home Infusion Center Phone: Agency Name:			Address:				
Administration code(s) (CPT):			Phone:		Fax:		
Address:			TIN:		NPI:		
E. PRODUCT INFORMATION							
Request is for: Prolia Xgeva Dose:	Frequer	ıcy: _			ICPCS Code:		
F. DIAGNOSIS INFORMATION - Please indicate	primary ICD code and s	pecify	any other where applic	able.			
Primary ICD Code: Se	econdary ICD Code:		Oth	er ICD Cod	e:		
G. CLINICAL INFORMATION - Required clinical i	nformation must be com	pletec	l in its entirety for all p	recertificatio	n requests.		
For All Requests: (Clinical documentation re					·		
Note: Xgeva is non-preferred. The preferred produc	ts are pamidronate or zo	ledror		nd zoledroni	c acid do not re	quire pre	certification.
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 reason(s							
Flease explain in there are any other medical reason(s	s) that the patient cannot t	ise pa	midionate of Zoledionic	aciu.			
Please provide the patient's Bone Mineral Density (BN	MD) score and date obtain	ed: T-	score.		Date:	1	1
Please indicate the location the BMD was measured:							
☐ Yes ☐ No Is the patient receiving 1000mg of ca				p			
Yes No Does the patient have clinical evidence							
Yes No Will the patient be using denosumab		enous	bisphosphonates?				
☐ Yes ☐ No Will the patient be using Prolia in com☐ Yes ☐ No Is the patient at high risk for fractures							
Yes No Has the patient had an osteoporotic f	•						
	racture?						
Yes No Does the patient ha	ive multiple risk factors for						
Yes No Does the patient hat Please explain (sel	ive multiple risk factors for ect all that apply): alco	hol int	ake of 4 or more units p				re
Yes No Does the patient hat Please explain (sel	ove multiple risk factors for ect all that apply): ☐ alco ritis ☐ current tobacco s	hol int mokin	ake of 4 or more units p				re
Yes No Does the patient hat Please explain (sel	ave multiple risk factors for ect all that apply): ☐ alco ritis ☐ current tobacco s se ☐ none of the above	hol int mokin	ake of 4 or more units p g □ advanced age □] frailty 🔲 i	ncreased fall risl	k	re
Yes No Does the patient hat Please explain (sel	ave multiple risk factors for ect all that apply): alcoritis current tobaccos se none of the above racture probability: 10-yea	hol int mokin r majo	ake of 4 or more units p g □ advanced age □ r osteoporotic fracture ri] frailty iı sk ≥ 20% or l	ncreased fall ris	k	re
Yes No Does the patient hat Please explain (sel	ave multiple risk factors for ect all that apply): alcoritis current tobaccosse none of the above racture probability: 10-yea become pregnant within 5	hol intomokin mokin r majo 5 mont	ake of 4 or more units pr g ☐ advanced age ☐ r osteoporotic fracture ri hs of discontinuing treat] frailty □ i sk ≥ 20% or l ment with de	ncreased fall risk nip fracture risk inosumab?	k	re



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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 pre	ecertification requests.				
For Prolia Requests:	•						
Post-menopausal osteoporosis							
Please select which of the following medication	n(s) was ineffective, not tolerated or contrain	indicated:					
Select all that apply: Alendronate (Binosto,			onate (Boniva)				
1 ' ' '	Actonel with Calcium or Atelvia) Terip	,	,				
	☐ Tamoxifen (Nolvadex/Soltamox) ☐ Tol		,				
☐ Other: Please identify:							
Prevention or treatment of osteoporosis in p	atients receiving endocrine therapy for	breast cancer					
Yes No Is the patient receiving endocr	ine therapy for breast cancer?						
	following endocrine therapy (aromatase inl						
	🗌 exemestane (Aromasin) 🔲 letrozole (F						
Yes No Is there documentation that the							
	the medication trial: \square Continued bone los						
	ge: <i> </i>	<u></u>					
Bisphosphonate #2 Date ran	ge:/ /						
Yes No Is there documented evidence		•					
☐ Yes ☐ No Is there documented evidence	•	• •					
Please select which of the following bisphosph							
Select all that apply: Alendronate (Binosto,	• • •	` , _	nate (Boniva)				
	Actonel with Calcium or Atelvia) Zoled	dronic acid (Zometa, Reciast)					
Other: Please identify:							
Treatment to increase bone mass in men red ☐ Yes ☐ No Does the patient have prostate							
Yes No Is the patient receiving androg							
Treatment of bone loss in men with osteopo	• • • • • • • • • • • • • • • • • • • •						
☐ Yes ☐ No Is there documentation that the		phonate trial of at least 1-vear durati	on?				
Please identify the failure of t							
	ge:/						
	ge:/						
Yes No Is there documented evidence							
Yes No Is there documented evidence	that the patient has a contraindication to b	pisphosphonates?					
Please select which of the following bisphosph							
Select all that apply: Alendronate (Binosto,	Fosamax or Fosamax plus D)	ate disodium (Didronel) 🔲 Ibandro	onate (Boniva)				
	Actonel with Calcium or Atelvia) Zoled	dronic acid (Zometa, Reclast)					
Other: Please identify:							
Treatment of glucocorticoid-induced osteop							
Yes No Is the patient initiating or continuor more?	nuing systemic glucocorticolds at a daily do	osage equivalent to 2.5 mg or greate	er of prednisone for 3 months				
Please select: ☐ initiating sy	stemic alucocorticoids	temic alucocorticoids					
	at expected to remain on glucocorticoids fo						
☐ Yes ☐ No Is there documentation that the							
	he medication trial: Continued bone los						
Bisphosphonate #1 Date ran	ge:/						
Bisphosphonate #2 Date ran	ge:/						
☐ Yes ☐ No Is there documented evidence		hosphonates?					
Yes No Is there documented evidence		•					
Please select which of the following bisphosph							
Select all that apply: Alendronate (Binosto,	Select all that apply: Alendronate (Binosto, Fosamax or Fosamax plus D)						
	Actonel with Calcium or Atelvia) Zoled	dronic acid (Zometa, Reclast)					
☐ Other: Please identify:							

Continued on next page



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G. CLINICAL INFORMATION (c	ontinued) - Required clinical informatio	on must be completed for ALL prece	ertification requests
For Xgeva Requests:			,
Bone metastases from solid tumo	ors		
Please indicate which of the followi	ng pertains to the patient: Bladder canc	er □ Breast cancer □ Kidnev can	cer
		ell lung cancer □ Prostate cancer [
		e specify:	
☐ Giant cell tumor of the bone			
☐ Prevention of skeletal-related	events in patients with multiple myelon	na	
Treatment of hypercalcemia of m			
☐ Yes ☐ No Has the patient be	en treated with intravenous bisphosphonate	e therapy?	
> Please indicate the	ne date range of therapy:/_/	- / /	
	nia of malignancy refractory to intravenous		
Yes No Has the albumin-co	orrected serum calcium level been tested?		
Please provide th	ne albumin-corrected serum calcium level:	mg/dL Date: /	1
·			
	Clinical documentation required for	an requests)	
1 — —	ave a hypersensitivity to denosumab?		
Please indicate what type of respon	nse the patient has experienced while on d		
		☐ Significant improveme	nt
H. ACKNOWLEDGEMENT			
Request Completed By (Sign	ature Required):		Date: /
Any person who knowingly files	a request for authorization of coverage	of a medical procedure or service v	with the intent to injure, defraud or deceive
			urpose of misleading, commits a fraudulent
insurance act, which is a crime a	nd subjects such person to criminal and	civil penalties.	-

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