MEDICARE FORM Evenity [®] (romosozumab-aqqg) Injectable Medication Precertification Request Page 1 of 2 (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: Evenity is non-preferred. The preferred products for MA plans		
Please indicate:		atment: Start date ///					are Prolia and IV zoledronic acid. The preferred product for MAPD plans is Forteo.	
Precertification R	equested By:			Phone		Fax:		
A. PATIENT INFOR	RMATION							
First Name:				Last Name:		State:		
Address:					City:		ZIP:	
Home Phone:		Work Phone:			Cell Phone			
DOB:	Allergies:				E-mail:			
_	lbs or	kgs	Height:	inches c	or cm	IS		
B. INSURANCE IN	FORMATION #:	Do	es natient have o	ther coverage?	☐ Yes ☐ No			
	#		•	aller coverage:				
Insured:			sured:					
C. PRESCRIBER I	NFORMATION							
First Name:		Las	st Name:		(Check Or	ne): 🗌 M.D. 🗌 D.	0. 🗌 N.P. 🗌 P.A.	
Address:				City:		State:	ZIP:	
Phone:	Fax:	St	Lic #:	NPI #:	DEA #:	UP	IN:	
Provider Email:	ROVIDER/ADMINISTRATIO		ontact Name:		Phone:			
Home Infusion C Agency Na Administration c Address: City: Phone:	on Center Phone: me: Center Phone: ame: ode(s) (CPT): State Fax: PIN:	ZIP:		□ Specialty Name: Address: City: Phone:		Retail Pharma Other State: Fax: PIN:	ZIP:	
E. PRODUCT INFO	RMATION							
-	enity [®] (romosozumab-a					HCPCS	Code:	
	ORMATION – Please indica							
G. CLINICAL INFORMATION – Required clinical information must be completed in its <u>entirety</u> for all precertification requests. For Initiation Requests (clinical documentation required for all requests): Note: Evenity is non-preferred. The preferred products for MA plans are Prolia and IV zoledronic acid. The preferred product for MAPD plans is Forteo. Yes No Has the patient had prior therapy with Evenity (romosozumab-aqqg) within the last 365 days? Yes No Has the patient had a trial and failure, intolerance, or contraindication to any of the following? (select all that apply) Prolia (denosumab) IV zoledronic acid Yes No Has the patient had a trial and failure, intolerance, or contraindication to any of the following? (select all that apply) Forteo (teriparatide) IV zoledronic acid Yes No Has the patient completed two years of treatment with a parathyroid hormone medication? Please explain if there are any medical reason(s) that the patient cannot use any of the following preferred products when indicated for the patient's diagnosis (select all that apply). IV zoledronic acid Prolia (denosumab) IV zoledronic acid Prolia (denosumab) IV zoledronic acid								
(select all that apply		s) that the patier	nt cannot use any o	of the following prefer	red products whe	n indicated for the p	atient's diagnosis	



MEDICARE FORM

Evenity[®] (romosozumab-aqqg) Injectable Medication Precertification Request

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For other lines of business: Please use other form.

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB				
C CLINICAL INFORMATION (continued)	Poquirad alinical information must be con	polotod in its optiraty for all prov	portification requests				
G. CLINICAL INFORMATION (continued) - Required clinical information must be completed in its entirety for all precertification requests. Please explain if there are any medical reason(s) that the patient cannot use Forteo (teriparatide):							
		e disodium (Didronel) 🔲 Ibandro nate (Skelid) 🔲 Zoledronic acid	(Zometa, Reclast)				
For Continuation Requests: (Clinical document Yes No Does the patient have a hyperse Please indicate what type of response the patient	ensitivity to romosozumab-aqqg?		mal response ❑ Significant improvement				
H. ACKNOWLEDGEMENT							
Request Completed By (Signature Require	d):		Date: / /				
Any person who knowingly files a request for a insurance company by providing materially t insurance act, which is a crime and subjects s	alse information or conceals material i	nformation for the purpose of					

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