

MEDICARE FORM

Darzalex Faspro[™] (daratumumab and hyaluronidase-fihj) Medication Precertification Request Page 1 of 2

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

For Michigan MMP: FAX: 1-844-241-2495 PHONE: 1-855-676-5772

For other lines of business: Please use other form.

Note: Darzalex Faspro is nonpreferred. The preferred products are Bortezomib and Velcade.

Please indicate: Start of treatment: Start date / Continuation of therapy, Date of last treatment /							
Precertification Requested By:				Phone	:	Fax:	
A. PATIENT INFO	RMATION						
First Name:			Last Name:			DOB:	
Address:				City:		State:	ZIP:
Home Phone:		Work Phone:		Cell Phone:		Email:	
Patient Current Wei	ght: lbs or	kgs Patien	t Height: inches	or cms	Allergies:		
B. INSURANCE IN	FORMATION						
Aetna Member ID #: Group #: Insured:			Does patient have other coverage? □ Yes □ No If yes, provide ID#: Carrier Name: Insured:				
Medicare: 🗌 Yes	No If yes, provid	de ID #:	Ме	edicaid: 🗌 Yes	□ No If yes, pro	vide ID #:	
C. PRESCRIBER I	NFORMATION						
First Name:			Last Name:	-	(Check Or	ne): 🗌 M.D. 🗌	D.O. 🗌 N.P. 🗌 P.A.
Address:				City:		State:	ZIP:
Phone:	Fax:		St Lic #:	NPI #:	DEA #:		UPIN:
Provider Email:			Office Contact Name:			Phone:	
Specialty (Check one): Oncologist Hematologist Other:							
D. DISPENSING P	ROVIDER/ADMINIS	TRATION INFOR	RMATION				
Place of Administration: Self-administered Physician's Office Outpatient Infusion Center Center Name: Home Infusion Center Agency Name:				Dispensing Provider/Pharmacy: Patient Selected choice Physician's Office Retail Pharmacy Specialty Pharmacy Other Name:			
Administration code(s) (CPT):				-			
Address:			ID.			Fax: PIN:	
	City: State: ZIP: Phone: Fax:					1 IIN	
TIN:		PIN:					
NPI:							
E. PRODUCT INFO			d huelurenideee fiki)	Deee	Erro er		
			d hyaluronidase-fihj)			iency:	
	_	e indicate primary	y ICD code and specify				
Primary ICD Code:			Secondary ICD Coc			ICD Code:	
			tion must be complete	d in its <u>entirety</u> fo	or all precertification	n requests.	
For ALL Requests (clinical documentation required for all requests): Note: Darzalex Faspro is non-preferred. The preferred products are Bortezomib and Velcade. Yes No Has the patient had prior therapy with Darzalex Faspro 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) Velcade Bortezomib Please explain if there are any other 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) Velcade Bortezomib							



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

Patient First Name	Patient Last Name	Patient Phone	Patient DOB						
G. CLINICAL INFORMATION - Re	equired clinical information mus	t be completed in its entirety for a	Il precertification requests						
Light chain amyloidosis		lor di <u>ornaroty</u> for di							
☐ Yes ☐ No Is the patient newly diagnosed with light chain amyloidosis?									
\longrightarrow Yes \square No Is the patient's disease relapsed or refractory?									
Yes 🗌 No Will the requested drug be used in combination with bortezomib, cyclophosphamide and dexamethasone?									
Multiple myeloma									
What is the prescribed regimen?									
The requested medication in combination with bortezomib, thalidomide, and dexamethasone									
└────> ☐ Yes ☐ No Is the patient eligible for transplant?									
☐ Yes ☐ No Will the requested medication be used as primary therapy?									
Yes No Will the requested medication be used for a maximum of 16 doses?									
□ The requested medication in combination with lenalidomide and dexamethasone □ Yes □ No Is the patient eligible for transplant?									
Yes ☐ No Will the requested medication be used as primary therapy?									
☐ Yes ☐ No Has the patient received one or more prior therapies?									
The requested medication in combination with bortezomib, melphalan, and prednisone									
\square The requested medication in combination with borezonic, methatian, and preditionic									
☐ Yes ☐ No Will the requested medication be used as primary therapy?									
The requested medication in combination with bortezomib and dexamethasone									
└────────────────────────────────────									
The requested medication in combination with carfilzomib and dexamethasone									
\square Yes \square No Is the patient's disease relapsed or progressive?									
The requested medication in combination with pomalidomide and dexamethasone									
ag	jent?	two prior therapies, including a pro	oteasome inhibitor (PI) and an immunomodulatory						
The requested medication as a single agent									
	as the patient received at least lent?	three prior therapies, including a p	proteasome inhibitor (PI) and an immunomodulatory						
			hibitor (PI) and an immunomodulatory agent?						
The requested medication in combination with cyclophosphamide, bortezomib, and dexamethasone									
☐ The requested medication will be used in combination with bortezomib, lenalidomide and dexamethasone									
	ill the requested medication be								
	in the requested medication be	used as primary merapy?							
For Continuation Requests (clinical documentation required for all requests)									
☐ Yes ☐ No Has the patient experienced disease progression or unacceptable toxicity while on the current regimen?									
Please select: Disease progression Unacceptable toxicity									
For light chain amyloidosis only:									
☐ Yes ☐ No Will the treatment duration exceed 24 months of treatment?									
H. ACKNOWLEDGEMENT									
Request Completed By (Signatu			Date: / /						
Any person who knowingly files a request for authorization of coverage of a medical procedure or service with the intent to injure, defraud or deceive any insurance company by providing materially false information or conceals material information for the purpose of misleading, commits a fraudulent insurance act, which is a crime and subjects such person to criminal and civil penalties.									

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