

## State of Oklahoma SoonerCare

## Darzalex<sup>®</sup> (Daratumumab) and Darzalex Faspro<sup>®</sup> (Daratumumab/Hyaluronidase-fihj) Prior Authorization Form

	Drug Information	1		
Physician billing (HCPCS o	ode:) Start Dat	te (or date of next dose):		
Dose:	Regimen	n:		
	Billing Provider Inform			
Provider NPI:	Provider Name:	:		
Provider Phone:	Provider Fax:			
Prescriber Information				
Prescriber NPI:	Prescriber Name:			
Prescriber Phone:	Prescriber Fax:	Specialty:		
	Criteria			
For Initial Authorization:  1. Please indicate the diagn  Light Chain Amyloide  A. Will daratumumats newly diagnosed of the diagn of the diagnosed of the diagn	be used as a single-agent in relapsed of be used in combination with bortezomibilisease? YesNo	de and dexamethasone as primary therapy for a nt (ASCT)? Yes No de and dexamethasone after at least 1 prior o, melphalan, and prednisone as primary therapy o, thalidomide, and dexamethasone as primary No thalidomide, and dexamethasone as primary No thenalidomide, and dexamethasone as primary No to the number of the following therapy indicate which therapy combination will be used:		
primary therapy?	Page 1 of 2			

### PLEASE PROVIDE THE INFORMATION REQUESTED AND RETURN TO:

University of Oklahoma College of Pharmacy Pharmacy Management Consultants Product Based Prior Authorization Unit

> Fax: 1-800-224-4014 Phone: 1-800-522-0114 Option 4

### **CONFIDENTIALITY NOTICE**

This document, including any attachments, contains information which is confidential or privileged. If you are not the intended recipient, be aware that any disclosure, copying, distribution, or use of the contents of this information is prohibited. If you have received this document in error, please notify the sender immediately by telephone to arrange for the return of the transmitted documents or to verify their destruction.



Manahar Nama.

## State of Oklahoma **SoonerCare**

# Darzalex<sup>®</sup> (Daratumumab) and Darzalex Faspro<sup>®</sup> (Daratumumab/Hyaluronidase-fihj) Prior Authorization Form

wember name:	Date of Birth:	iviember ID#:	
	Criteria		
same regimen? Yes No J. Will daratumumab be used a an immunomodulatory agen Yes No If diagnosis is not listed abov	information, continued:  for disease relapse after 6 m o as a single-agent after ≥3 pr nt, or double refractory to a P re, please indicate diagnos	nplete all pages will result in procession on the following primary induction the rior therapies, including a protease inlead an immunomodulatory agent?	erapy with the hibitor (PI) and
For Continued Authorization:  1. Date of last dose:  2. Does member have any evidence o  3. Has the member experienced adverse reactions	rse drug reactions related to	daratumumab therapy? Yes No_	
Additional Information:			

Prescriber Signature: I certify that the indicated treatment is medically necessary and all information is true and correct to the best of my knowledge. Please do not send in chart notes. Specific information will be requested if necessary. Failure to complete this form in full will result in processing delays.

Date:

Page 2 of 2

## PLEASE PROVIDE THE INFORMATION REQUESTED AND RETURN TO:

University of Oklahoma College of Pharmacy Pharmacy Management Consultants **Product Based Prior Authorization Unit** 

> Fax: 1-800-224-4014 Phone: 1-800-522-0114 Option 4

### **CONFIDENTIALITY NOTICE**

Manahar 104.

This document, including any attachments, contains information which is confidential or privileged. If you are not the intended recipient, be aware that any disclosure, copying, distribution, or use of the contents of this information is prohibited. If you have received this document in error, please notify the sender immediately by telephone to arrange for the return of the transmitted documents or to verify their destruction.