

State of Oklahoma Oklahoma Health Care Authority Folotyn[®] (Pralatrexate) Prior Authorization Form

Member Name:	Date of Birth:	Member ID#:
	Drug Information	
Physician billing (HCPCS code:) Start Date (or date of next dose):		te of next dose):
	Regimen:	
Billing Provider Information		
Provider NPI:	Provider Name:	
	Provider Fax:	
Prescriber Information		
Prescriber NPI:	Prescriber Name:	
Prescriber Phone:	Prescriber Fax:	Specialty:
Criteria		
B. Does member have related a primary Cutaneous Lymphoma, Ex. A. Will pralatrexate be a combination chemical and the combination be a combinat	ed as a single-agent? Yes No apsed or refractory disease? Yes s and information: a/Lymphoma Lymphoma (ALCL), Primary Cutane e multifocal lesions or regional nodes? e used as primary treatment? Yes	eous ? Yes No No al Type g additional therapy with an alternate ? Yes No IF)/Sézary Syndrome (SS) _ No
3. Has the member experienced	ence of progressive disease while on pedical description of any adverse drug reactions related to rerse reactions:	o pralatrexate therapy? Yes No
Prescriber Signature:	Da	ate:
knowledge. Please do not send in complete this form in full will result in	chart notes. Specific information will b	nation is true and correct to the best of my be requested if necessary. Failure to

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

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