

Brukinsa (zanubrutinib) Prior Authorization Request Form



Caterpillar Prescription Drug Benefit Phone: 877-228-7909 Fax: 800-424-7640

Instructions: Please fill out all applicable sections completely and legibly. Attach any additional documentation that is important for the review (e.g., chart notes or lab data, to support the authorization request). Information contained in this form is Protected Health Information under HIPAA.

MEMBER INFORMATION				
LAST NAME:	FIRST NAME:			
PHONE NUMBER:	DATE OF BIRTH:			
STREET ADDRESS:				
CITY:	STATE: ZIP CODE:			
PATIENT INSURANCE ID NUMBER:				
MALE FEMALE HEIGHT (IN/CM): WEIG	iHT (LB/KG): ALLERGIES:			

IF YOU ARE NOT THE PATIENT OR THE PRESCRIBER, YOU WILL NEED TO SUBMIT A PHI DISCLOSURE AUTHORIZATION FORM WITH THIS REQUEST WHICH CAN BE FOUND AT THE FOLLOWING LINK: <u>https://magellanrx.com/member/external/commercial/common/doc/en-us/phi_disclosure_authorization.pdf</u>

PATIENT'S AUTHORIZED REPRESENTATIVE (IF APPLICABLE): _______AUTHORIZED REPRESENTATIVE'S PHONE NUMBER: ______

PRESCRIBER INFORMATION			
LAST NAME:	FIRST NAME:		
PRESCRIBER SPECIALTY:	EMAIL ADDRESS:		
NPI NUMBER:	DEA NUMBER:		
PHONE NUMBER:	FAX NUMBER:		
STREET ADDRESS:			
CITY:	STATE: ZIP CODE:		
REQUESTOR (if different than prescriber):	OFFICE CONTACT PERSON:		

MEDICATION OR MEDICAL DISPENSING INFORMATION					
MEDICATION NAME:					
DOSE/STRENGTH:	FREQUENCY:	LENGTH OF THERAPY/REFILLS:	QUANTITY:		
NEW THERAPY	RENEWAL	IF RENEWAL: DATE THERAP	(INITIATED:		
DURATION OF THERAPY (SPECIFIC DATES):					

Continued on next page.

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1. HAS THE PATIENT TRIED ANY OTHE	R MEDICATIONS FOR THIS CONDITION?	YES (if yes, complete below) NO			
MEDICATION/THERAPY (SPECIFY	DURATION OF THERAPY (SPECIFY	RESPONSE/REASON FOR			
DRUG NAME AND DOSAGE):	DATES):	FAILURE/ALLERGY:			
2. LIST DIAGNOSES:		ICD-10:			
Mantle Cell Lymphoma (MCL)					
□ Waldenstrom's Macroglubinemia(WM					
□ Marginal Zone Lymphoma(MZL)	5 1 1				
□ Chronic Lymphacytic Leukemia(CLL / S	11)				
Relapsed or Refractory CLL/SLL	,				
□ Other diagnosis:	ICD-10 Code(s):				
	N: PLEASE PROVIDE ALL RELEVANT CLIN	ICAL INFORMATION TO SUPPORT A			
PRIOR AUTHORIZATION.					
Clinical Information:					
In this during haing properties of the this r	ations as well of a two stars at was income				
trial? • Yes • No	patient as part of a treatment regimen	specified within a sponsored clinical			
Founding and a state of Manuala Call Luman have					
	na(MCL), please answer the following:				
	ease as confirmed by a computed tomo	ography/magnetic resonance imaging			
laboratory report? □ Yes □ No Plea	se submit documentation				
Does the patient's disease have histo	logic evidence of MCL morphology? 🗆 Y	es \square No Please submit documentation			
	(11; 14) translocation AND/OR overex	pression of cyclin D1?			
Yes No Please submit documents	ition				
Use the notions foiled one provides the					
Has the patient falled one previous t	herapy for MCL? □ Yes □ No Please	submit documentation			
Use the restant over is rely herd to set					
Has the patient previously had treatment with another BTK inhibitor (such as Calquence [®] / acalabrutinib or					
Imbruvica ® / ibrutinib)? 🗆 Yes 🗆 No					
Is the patient Eastern Cooperative Oncology Group (ECOG) performance status of 0, 1 or 2 (is ambulatory and					
•	arry out any work activities; up and ab	out more than 50% of waking hours?			
□ Yes □ No Please submit documentation					
For diagnosis of Waldenstrom's Macroglubinemia, please answer the following:					
Has patient had at least one prior therapy? Yes No Please submit documentation.					
Has patient had prior treatment with a BTK inhibitor such as Imbruvica(ibrutinib) or Calquence(acalabrutinib)? 🗆					
Yes 🗆 No Please submit documentation.					
For Marginal Zone Lymphoma, please	e answer the following:				
, picco					

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Has patient had at least one prior therapy with an anti-CD20-based regimen such as rituximab, ibritumomab(Zevalin), obinutuzumab(Gazyva), Tositumomab (Bexxar) or ofatumumab(Arzerra ?
Yes No Please submit documentation.

Has patient had prior treatment with a BTK inhibitor such as Imbruvica(ibrutinib) or Calquence(acalabrutinib)? Yes \Box No *Please submit documentation.*

<u>For diagnosis of Chronic Lymphacytic Leukemia(CLL / SLL), please answer the following</u>: Does patient have a diagnosis of CD20 positive CLL/SLL?
□ Yes □ No Please submit documentation.

Has patient been previously treated for their CLL/ SLL?
□ Yes
□ No Please submit documentation.

Has patient been previously treated with at least one systemic therapy?
Subscript Yes In No Please submit documentation.

If patient had been previously treated, did patient have at least 2 cycles of treatment?
□ Yes □ No Please submit documentation.

Is patient unable to be treated with fludarabine, cyclophosphamide, and rituximab(FCR)?

Yes O Please submit documentation.

Does patient have an Eastern Cooperative Oncology Group(ECOG) performance score of 0,1, or 2?
Yes No Please submit documentation.

Has patient been previously treated with a BTK inhibitor such as ibrutinib(Imbruvica), acalabrutinib(Calquence), zanubrutinib(Brukinsa), tirabrutinib(Velexbru) or orleabrutinib(Hibruka)?
Yes ON *Please submit documentation*.

For diagnosis of Follicular Lymphoma(FL), please answer the following: Has patient been treated with 2 or more prior systemic treatments for follicular lymphoma?
• Yes • No Please submit documentation.

Was patient previously receiving an anti-CD20 antibody and an appropriate alkylator-based combination therapy? • Yes • No *Please submit documentation.*

Does patient have disease progression after completion of the most recent therapy or refractory disease?

Yes
Please submit documentation.

Does patient have an Eastern Cooperative Oncology Group (ECOG) performance status of 0,1 or 2?
Yes ON

Has patient had prior treatment with a Bruton's tyrosine kinase (BTK) inhibitor such as tirabrutinib(Velexbru), orleabrutinib(Hibruka), Imbruvica(ibrutinib), Calquence(acalabrutinib) or Brukinsa(zanubrutinib)?
• Yes • No Please submit documentation.

Will patient use Brukinsa(zanubrutinib) in combination with obinutuzumab(Gazyva)?
Ves
No Please submit documentation.





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Are there any other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to this review?

Please note: Not all drugs/diagnosis are covered on all plans. This request may be denied unless all required information is received.

ATTESTATION: I attest the information provided is true and accurate to the best of my knowledge. I understand that the Health Plan, insurer, Medical Group or its designees may perform a routine audit and request the medical information necessary to verify the accuracy of the information reported on this form.

Prescriber Signature or Electronic I.D. Verification: _

Date:

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FAX THIS FORM TO: 800-424-7640

MAIL REQUESTS TO: Magellan Rx Management Prior Authorization Program

Attn:CP-4201 P.O.Box 64811 St. Paul, MN 55164-0811 Phone: 877-228-7909



