

State of Oklahoma SoonerCare

Imbruvica[®] (Ibrutinib) Prior Authorization Form

Member Name:	Date of Birth:	Member ID#:		
Drug Information				
<u> </u>	(HCPCS code:)			
Dose:Regimen:	Star	rt Date (or date of next dose):		
Billing Provider Information				
Provider NPI:	rovider NPI: Provider Name:			
Provider Phone: Provider Fax:				
Prescriber Information				
Prescriber NPI: Prescriber Name:				
Prescriber Phone:	_ Prescriber Fax:	Specialty:		
Criteria				
 1. Will ibrutinib be used as a single-agent? Yes No				
A. Does member's diagnosis related B-cell lymphomas, Yes No		nphomas, human immunodeficiency virus (HIV)- rative disorders, or high grade B-cell lymphoma?		
B. Will ibrutinib be used as aC. Will ibrutinib be used as mcell transplant (HSCT)? Ye	component of aggressive indunaintenance therapy following a es No	aggressive induction therapy or hematopoietic stem		
Lymphoma	· non-germinal center diffuse la	iency Syndrome (AIDS)-Related B-Cell arge B-cell lymphoma? Yes No No		
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

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Pharm – 101 4/3/2024



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Member Name:	Date of Birth:	Member ID#:
Page 2 of 2—Please complete and re	eturn <u>all</u> pages. Failure to complete all pa	ges will result in processing delays.
 4. Please indicate the diagnosis and ir Post-Transplant Lymphoprol A. Is the member's diagnosis B. Please indicate member Partial therapy responsistent disease Progressive disease 	Ilferative Disorders sis non-germinal center B-cell type? Yes r's disease status: conse	_ No
☐ Chronic Lymphocytic Leuke	mia (CLL)/Small Lymphocytic Lymphoma combination with bendamustine, rituximab, c	
Hairy Cell Leukemia	third-line or subsequent therapy for refracto	
 □ Waldenström's Macroglobuli A. Will ibrutinib be used in a □ Primary Central Nervous System A. Is member not a candidate 	inemia (WM)/Lymphoplasmacytic Lymphocombination with rituximab (Rituxan®)? Yes_stem (CNS) Lymphoma ate for, or intolerant to high-dose methotrexater refractory or progressive disease? Yes	No ate? Yes No
If diagnosis is not listed above, pleas	se indicate diagnosis:	
Additional Information:		
3. Has the member experienced any a	of progressive disease while on ibrutinib? Yes adverse drug reactions related to ibrutinib the eactions:	erapy? Yes No
Additional Information:		
Please complete and return <u>all</u> pages Please do not send in chart notes. Spec	Page 2 of 2 s. Failure to complete all pages will result cific information will be requested if necessal	in processing delays.
Prescriber Signature:	Date:	
I certify that the indicated treatment is m Please do not send in chart notes. Specific processing delays.	edically necessary and all information is true information will be requested if necessary. Failure	and correct to the best of my knowledge. e to complete this form in full will result in

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