

BRUKINSA (zanubrutinib)

Instructions

Plan Member Signature

Please complete Part A and have your physician complete Part B. Completion and submission is not a guarantee of approval. Any fees related to the completion of this form are the responsibility of the plan member. Drugs in the Prior Authorization Program may be eligible for reimbursement if the patient does not qualify for coverage under a primary plan or a government program. Drugs used for indications not approved by Health Canada may be denied. For Quebec plan members, RAMQ exception drug criteria may apply. The decision for approval versus denial is based on pre-defined clinical criteria, primarily based on Health Canada approved indication(s) and on supporting evidence-based clinical protocols. The plan member will be notified whether their request has been approved or denied. Please note that you have the right to appeal the decision made by Express Scripts Canada.

Part A - Patient Patient information First Name: Last Name: Insurance Carrier Name/Number: Group Number: Client ID: Date of Birth (YYYY/MM/DD): Relationship: | Employee | Spouse | Dependent Language: English French Gender: | | Male | | Female Address: City: Province: Postal Code: Email address: Telephone (home): Telephone (cell): Telephone (work): Coordination of benefits **Patient** Is the patient enrolled in any patient assistance program? Yes No **Assistance Program** Contact Name: _ Has the patient applied for reimbursement under a provincial plan? Yes No N/A **Provincial** Coverage What is the coverage decision of the drug? Approved Denied *Attach decision letter* Has the patient applied for reimbursement under a primary plan? Yes No N/A **Primary** Coverage What is the coverage decision of the drug? Approved Denied *Attach decision letter* Authorization On behalf of myself and my eligible dependents, I authorize my group benefit provider, and its agents, to exchange the personal information contained on this form. I give my consent on the understanding that the information will be used solely for purposes of administration and management of my group benefit plan. This consent shall continue so long as my dependents and I are covered by, or are claiming benefits under the present group contract, or any modification, renewal, or reinstatement thereof.

Date



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Part B - Prescriber

Please see instructions on page 1 and complete all sections below. <u>Incomplete forms may result in automatic denial</u>. Please do **not** provide genetic test information or results.

RUKINSA (zanubrutinib)		New request	Renewal request*
Dose	Administration (ex: oral, IV, etc)	Frequency	Duration
Site of drug administration:			
Home Physicia	n's office/Infusion clinic	Hospital (outpatient)	Hospital (inpatient)
* Please submit proof of prior	coverage if available		
ECTION 2 – ELIGIBILITY (CRITERIA		
1. Please indicate if the pation	ent satisfies the below criteria:		
Waldenström's Macroglobuline		0.44.D.:	
_	Valdenström's macroglobulinemia		
prior therapies in the	n inadequate response or has a d chart below), AND	ocumented intolerance to a	prior therapy for WM (<i>Please list</i>
BRUKINSA will be use	d as monotherapy		
Mantle Cell Lymphoma			
For the treatment of re	elapsed or refractory mantle cell ly	mphoma (MCL) in an adult,	AND
The patient has had a list prior therapies in t	n inadequate response or has a d he chart below)	ocumented intolerance to a	prior therapy for MCL (<i>Plea</i> se
Marginal Zone Lymphoma			
For the treatment of n	narginal zone lymphoma (MZL) in a	an adult, AND	
The patient has had a list prior therapies in t	n inadequate response or has a d he chart below)	ocumented intolerance to pr	rior anti-CD20 therapy (Please
Chronic Lymphocytic Leukemia	a – Previously Untreated		
For the treatment of c	hronic lymphocytic leukemia (CLL)) in an adult, AND	
The patient has not pr	reviously received treatment for CL	L, AND	
The patient is not a ca	andidate for chemoimmunotherapy	y with fludarabine, cyclophos	sphamide, and rituximab (FCR),
AND			



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Chronic Lymphocytic Leukemia - Previou	usly Treated						
For the treatment of relapsed or refractory chronic lymphocytic leukemia (CLL) in an adult, AND							
The patient relapsed after, or was refractory to a prior therapy for CLL (<i>Please list prior therapies in the chart below</i>), AND							
BRUKINSA will be used as mond	otherapy						
Follicular Lymphoma							
For the treatment of relapsed or	For the treatment of relapsed or refractory grade 1, 2, or 3a follicular lymphoma (FL) in an adult, AND						
The patient has had an inadeque for FL (Please list prior therapies			lerance to at le	east 2 prior system	nic therapies		
BRUKINSA will be used in comb	ination with GAZYVA (ob	inutuzumab)					
None of the above criteria applies. Relevant additional information:							
Please list previously tried therapies	;						
Please list previously tried therapies		Duration	of therapy		r cessation		
	Dosage and administration	Duration From	of therapy To	Reason for Inadequate response	r cessation Allergy/ Intolerance		
Please list previously tried therapies	Dosage and			Inadequate	Allergy/		
Please list previously tried therapies	Dosage and			Inadequate	Allergy/		
Please list previously tried therapies	Dosage and			Inadequate	Allergy/		
Please list previously tried therapies	Dosage and			Inadequate	Allergy/ Intolerance		
Please list previously tried therapies	Dosage and			Inadequate	Allergy/ Intolerance		



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SECTION 3 - PRESCRIBER INFORMATION

Physician's Name:	
Address:	
Tel:	Fax:
License No.:	Specialty:
Physician Signature:	Date:

Please fax or mail the completed form to Express Scripts Canada®

Fax: Express Scripts Canada Clinical Services 1 (855) 712-6329

Mail: Express Scripts Canada Clinical Services 5770 Hurontario Street, 10th Floor Mississauga, ON L5R 3G5