

Drug Prior Authorization Form Rituxan (rituximab)

The purpose of this form is to obtain information required to assess your drug claim.

IMPORTANT: Please answer all questions. Your claim assessment will be delayed if this form is incomplete or contains errors.

Any costs incurred for the completion of this form are the responsibility of the plan member/patient.

Canada Life recognizes and respects the importance of privacy. Personal information collected is used for the purposes of assessing eligibility for this drug and for administering the group benefits plan. For a copy of our Privacy Guidelines, or if you have questions about Canada Life's personal information policies and practices (including with respect to service providers), refer to www.canadalife.com or write to Canada Life's Chief Compliance Officer.

I authorize Canada Life, any healthcare provider, my plan administrator, any insurance or reinsurance company, administrators of government benefits or patient assistance programs or other benefits programs, other organizations, or service providers working with Canada Life or any of the above, located inside or outside Canada, to exchange personal information when relevant and necessary for these purposes. I understand that personal information may be subject to disclosure to those authorized under applicable law within or outside Canada.

I acknowledge that the personal information is needed to assess eligibility for this drug and to administer the group benefits plan. I acknowledge that providing consent will help Canada Life to assess my claim and that refusing to consent may result in delay or denial of my claim. Canada Life reserves the right to audit the information provided on this form at any time and this consent extends to any audit of my claim. This consent may be revoked by me at any time by sending written instruction to that effect.

I also consent to the use of my personal information for Canada Life and its affiliates' internal data management and analytics purposes.

If the patient is a person other than myself, I confirm that the patient has given their consent to provide their personal information and for Canada Life to use and disclose it as set out above.

I certify that the information given below is true, correct, and complete to the best of my knowledge. Failure to provide true, correct and complete information on this form could result in revocation of any approval decision, a requirement to repay paid claims or other appropriate action.

Plan Member's signature: _	Date:

Form Completion Instructions:

- 1. Complete "Patient Information" sections.
- 2. Have the prescribing physician complete the "Physician Information" sections.
- 3. Send all pages of the completed form to us by mail, fax or email as noted below.

Note: As email is not a secure medium, any person with concerns about their prior authorization form/medical information being intercepted by an unauthorized party is encouraged to submit their form by other means.

Mail to: The Canada Life Assurance Company

Drug Claims Management

PO Box 6000

Winnipeg MB R3C 3A5

Email to: <u>cldrug.services@canadalife.com</u>

Attention: Drug Claims Management

Fax to: The Canada Life Assurance Company

Fax 1-204-946-7664

Attention: Drug Claims Management

For additional information regarding Prior Authorization and Health Case Management, please visit our Canada Life website at www.canadalife.com or contact Group Customer Contact Services at 1-800-957-9777. Deaf or hard of hearing and require access to a telecommunications relay service? Please contact us at 711 for TTY to Voice or 1-800-855-0511 for Voice to TTY.

(Continued on next page)



Patient Information Rituxan (rituximab)

Plan Member Information – Complete all sections of this page (please print)					
Plan Member:		Patient Name:			
Plan Name:	Plan Number:		Plan Member ID Number:		
Patient Date of Birth (DD/MM/YYYY):	Address (number, street,	, city, province, postal cod	de):		
,			,		
Please indicate preferred contact number and if the	here are any times when t	elephone contact with yo	u about your claim would be most convenient.		
May we contact you by email? (Note that some co	orrespondence may still n	eed to be sent by regular	mail).		
Yes No If yes, please provide email ad	ldress:				
Tell us if you have been on this drug b	pefore				
Is the patient currently on, or previously been on	this drug? ☐ Yes ☐ N	0			
If Yes, a) indicate start date (DD/MM/YYYY):					
b) coverage provided by:					
(if coverage is not provided by Canada Lif	e please provide pharmad	cy print-out showing purc	hase of this drug)		
Tell us if you have coverage with any	other benefits plan				
Does the patient have drug coverage under any If Yes, name of other Insurance Company: If other plan is with Canada Life, tell us the plan and the plan member: Relationship to patient:	and ID number:				
Provide details and attach documentation of acceptance or decline:					
Tell us about any Provincial or other of	coverage you may h	nave			
Does the patient have coverage under a provincial program or from any other source?					
Is the patient currently receiving disability benefits for the condition for which this drug has been prescribed? Yes No					
Tell us about any Patient Assistance Program you might be enrolled in					
Has the patient enrolled in the patient assistance	e program for this drug?	☐ Yes ☐ No			
If Yes, please provide the following information:					
Patient assistance program patient ID Number:					
2. Patient assistance program contact person name and phone number:					
Contact Name:		Phone Number:			

(Continued on next page)



Physician Information Rituxan (rituximab)

Note to Physician: In order to assess a patient's claim for this drug, we require detailed information on the patient's prescription drug history as requested below.

Attach extra information if necessary. GENETIC TEST RESULTS ARE NOT REQUIRED

Physician's Information (please print)				
Name of prescribing physician:				
Specialty:				
Specialty.				
Address (number, street, city, province, postal code):				
Telephone Number (including area code):	Fax Number (including area code):			
Prescribed dosage and regimen:				
\square Two 1000mg infusions scheduled two weeks apart				
☐ 375mg/m² once weekly for 4 doses				
☐ Other (please specify):				
Provide rationale:				
2. Health Canada indication (include date of initial diagnosis) (MM/YY	YY):			
\square Relapsed or refractory low-grade or follicular non-Hodgkin's lym	phoma			
☐ Diffuse large B-cell lymphoma (DLBCL)				
☐ Stage III/IV follicular non-Hodgkin's lymphoma				
☐ B-cell chronic lymphocytic leukemia (B-CLL)				
☐ Rheumatoid arthritis				
☐ Granulomatosis with Polyangiitis (GPA, also known as Wegener's Granulomatosis) and Microscopic Polyangiitis (MPA)				
Other (Approved by Health Canada):				
Complete questions 1 - 6 and Other Condition (Health Canada approved)				
Genetic test results are not required				
Is this drug being presribed in accordance with approved Health Co	anada indications ¹ ?			
\square Yes, complete questions 1 - 6 and Physician's Information				
\square No, prescribed use is not approved by Health Canada:				
Complete questions 1 - 6 and Off Label use				



Physician Information Rituxan (rituximab)

Physician's Information (continued) (please print)

¹Approved Health Canada Indications and Clinical Use for Rituxan:

- The treatment of patients with relapsed or refractory low-grade or follicular, CD20 positive, B-cell non-Hodgkin's lymphoma.
- The treatment of patients with CD20 positive, diffuse large B-cell non-Hodgkin's lymphoma (DLBCL) in combination with CHOP (cyclophosphamide, doxorubicin, vincristine, and prednisone) chemotherapy.
- The treatment of patients with previously untreated Stage III/IV follicular, CD20 positive, Bcell non-Hodgkin's lymphoma in combination with CVP (cyclophosphamide, vincristine and prednisolone) chemotherapy.
- Maintenance treatment of patients with follicular non-Hodgkin's lymphoma who have responded to induction therapy with either CHOP or CHOP plus RITUXAN.
- Single-agent maintenance treatment of previously untreated patients with advanced follicular non-Hodgkin's lymphoma with high tumour burden and who have responded to induction therapy with either CHOP plus RITUXAN or CVP plus RITUXAN.
- The treatment of patients with previously untreated or previously treated B-cell chronic lymphocytic leukemia (B-CLL), Binet Stage B or C, in combination with fludarabine and cyclophosphamide.
- In combination with methotrexate is indicated in adult patients: to reduce signs and symptoms in adult patients with moderately to severely active rheumatoid arthritis who have had an inadequate response or intolerance to one or more tumour necrosis factor (TNF) inhibitor therapies.
- In combination with glucocorticoids is indicated for the induction of remission in adult patients with severely active Granulomatosis with Polyangiitis (GPA, also known as Wegener's Granulomatosis) and Microscopic Polyangiitis (MPA).

3. What is the anticipated duration of treatment with this drug?						
6. Drug and Treatment History – must be completed for every request.						
Drug(s) and Treatment(s) past and present	Dosing Regimen	Start Date (DD/MM/YYYY)	End Date (DD/MM/YYYY)	Clinical Results/Outcome		
				☐ Failure ☐ Intolerance ☐ Other Clinical details:		
				☐ Failure ☐ Intolerance ☐ Other Clinical details:		
				☐ Failure ☐ Intolerance ☐ Other Clinical details:		

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Physician Information Rituxan (rituximab)

Non-Hodgkin's lymphoma – Genetic test results are not required						
Does the patient have:						
Relapsed or refractory low-grade or follicular non-Hodgkin's lymphoma?						
Diffuse large B-cell non-Hodgki	n's lymphoma (DLBCL) ai	nd Rituxan will be p	rescribed in combin	ation with CHOP (cyclophosphamide, doxorubicin,		
vincristine, and prednisone) che	emotherapy?	□ No				
Previously untreated Stage III/IV follicular non-Hodgkin's lymphoma and Rituxan will be prescribed in combination with CVP						
(cyclophosphamide, vincristine and prednisolone) chemotherapy? $\ \square$ Yes $\ \square$ No						
Follicular non-Hodgkin's lympho	oma who have responded	d to induction therap	y with either CHOP	or CHOP plus RITUXAN and Rituxan will		
be prescribed as Maintenance t	reatment?	□ No				
Advanced follicular non-Hodgkin	's lymphoma with high tun	nour burden and has	responded to induct	ion therapy with either CHOP plus		
RITUXAN or CVP plus RITUXAN	and is previously untreated	d and Rituxan will be	prescribed as single	-agent maintenance treatment? ☐ Yes ☐ No		
D. cell above the bounds and	- II	0				
B-cell chronic lymphocytic				•		
		mia Binet Stage B o	C and Rituxan will	be prescribed in combination with fludarabine and		
cyclophosphamide?] No					
Rheumatoid Arthritis - Ge	netic test results a	re not required				
Swollen joint count Results of the following and date (DD/MM/YYYY)						
Rheumatoid Factor Positive	Yes		ESR			
Current results and date of one of the following (DD/MM/YYYY) CDAI DAS28 HAQ		CRP				
Drug(s) and Treatment(s) past and present	Dosing Regimen	Start Date (DD/MM/YYYY)	End Date (DD/MM/YYYY)	Clinical Results/Outcome		
				☐ Failure ☐ Intolerance ☐ Other		
				Clinical details:		
				☐ Failure ☐ Intolerance ☐ Other		
				Clinical details:		
				☐ Failure ☐ Intolerance ☐ Other		
				Clinical details:		
				☐ Failure ☐ Intolerance ☐ Other Clinical details:		

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Email to: <u>cldrug.services@canadalife.com</u>

Attention: Drug Claims Management

Physician Information Rituxan (rituximab)

	omatosis with Polyangiitis (GPA, also known as W tic test results are not required	'egener's Grar	nulomatosis) and	Microscopic Polyangiitis (MPA)
Does the	e patient have severely active disease?		☐Yes	□No
Will Ritux	xan be prescribed in combination with glucocorticoids for the i	nduction of remis	sion? Yes	□No
Other c	condition (Health Canada approved) – Genetic	test results a	re not required	
Please p	provide any relevant information related to the disease and at	tach supporting o	documentation.	
Off-labe	el use - Genetic test results are not required			
Question	ns 1-6 must be completed.			
Is there o	clinical evidence supporting the off-label use of this drug? \Box	Yes No		
Provide o	clinical literature/studies to support the request for off-label u	se, such as:		
•	At least two Phase II or two Phase III clinical trials showing	consistent result	s of efficacy; and	
•	Published recommendations in evidence-based guidelines	supporting its us	e.	
Provide r	medical rationale why Rituxan has been prescribed off-label in	nstead of an alter	rnate drug with an ap	proved indication for this condition.
Provide a	any pertinent medical history or information to support this of	f-label request.		
If this is a	a renewal request, provide documentation showing treatment	t efficacy since pr	revious request.	
authoriza informati I certify t	Physician: To be eligible for reimbursement, Canadation from a pharmacy designated by Canada Life. ion. That the information provided is true, correct, and can be seen as a signature:	If applicable, a	health case man	
•			Date	
License N	Number:		_	
	rtant to provide the requested information in detail to hear to audit. The completed form can be returned to Car			ns for the above drug. This form may
	email is not a secure medium, any person with concer ercepted by an unauthorized party is encouraged to su			form/medical information being
Mail to:	The Canada Life Assurance Company Drug Claims Management PO Box 6000 Winnipeg MB R3C 3A5	Fax to:	Fax 1-204-946-7	e Assurance Company 1664 Claims Management

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