

MEDICARE FORM

Riabni[®] (rituximab-arrx), Rituxan[®] (rituximab), Ruxience (rituximab-pvvr), Truxima (rituximab-abbs) Medication Precertification Request Page 1 of 3

(All fields must be completed and return both pages for precertification review.)

For Illinois MMP: FAX: 1-855-320-8445 PHONE: 1-866-600-2139

For other lines of business: Please use other form.

Note: Rituxan, Rituxan Hycela, and Truxima are preferred. Riabni and Ruxience are non-preferred. For rheumatoid arthritis, all Rituxan and biosimilar products are non-preferred.

Please indicate: Start of treatment, star	t date:	/ /		ontinuation of therap	oy, date of la	st treatment:	<u> </u>	
Precertification Requested By:				Phone:		Fax:	:	
A. PATIENT INFORMATION								
First Name:		Last Name:					DOB:	
Address:			City:			State:	ZIP:	
	k Phone:			Phone:		E-mail:		
		inches or				L-man.		
Current Weight:lbs_orkgs	Height:	inches or	cms	Allergies:				
B. INSURANCE INFORMATION								
Member ID #:		Does patient have o		-	∕es □ No			
Group #: Insured:		If yes, provide ID#: Insured:		Ca	rier Name:			
C. PRESCRIBER INFORMATION								
First Name:		Last Name:		(Check one)] D.O. 🗌 N.P. 🗌 P.A.	
Address:		Lust Humo.	City:	(State:	ZIP:	
Phone: Fax:	St Lic	<i>щ</i> .	NPI #	4.	DEA #:	State.	UPIN:	
			INPI #	<i>t</i> .			UPIN.	
Provider Email:		Contact Name:			Phone:			
D. DISPENSING PROVIDER/ADMINISTRA Place of Administration:		IATION						
Self-administered Physician's Office			Dispensing Provider/Pharmacy: Outpatient Dialysis Center Physician's Retail Pharmacy Mail Order Name:			Pharmacy		
Address:				Address:		Stata	ZIP:	
City: Sta								
Phone: Fa				Phone: TIN:				
TIN: PI NPI:	N:			NPI:		FIN		
E. PRODUCT INFORMATION								
		(rituximah) 🔲 Bu	viana))	
Request is for: Riabni (rituximab-arrx)		Directions for Use		(intuximab-pvvi)		HCPCS (
F. DIAGNOSIS INFORMATION - Please ind	icate primary I	CD code and specify	any o	ther any other where	applicable (*).		
Primary ICD Code:			Other	ICD Code:				
G. CLINICAL INFORMATION - Required clin		•	d for A	LL precertification red	quests.			
For All Requests (clinical documentation Note: Rituxan, Rituxan Hycela, and Truxin For rheumatoid arthritis, all Rituxan and the second sec	na are preferr biosimilar pro janz/Xeljanz X erapy with Riak nd failure, intol Rituxan Hycela reason(s) tha	red for most indicati ducts are non-prefe (R are preferred for oni (rituximab-arrx) or lerance, or contraindi a (rituximab/hyaluroni t the patient cannot u	rred. I MAPE Ruxie cation dase h ise any	nflectra, Remicade, plans. nce (rituximab-pvvr) to any of the followin numan)	and Simpor within the las g? (select all (rituximab-ab erred produc	ni Aria are pret t 365 days? that apply) ibs) ts when indicat	ed for the patient's	
Yes No Has the patient had a trial an □ Remicade (infliximab) □ □ Yes No Has the patient had a trial an □ Ptess No Has the patient had a trial an □ Enbrel (etanercept) □ Please explain if there are any other medical diagnosis? (select all that apply) □ Please explain if there are any other medical □	☐ Inflectra (inf nd failure, intol Humira (adalir I reason(s) tha de (infliximab)	liximab-dyyb) □ Sir lerance, or contraindi numab) □ Kevzara t the patient cannot u □ Inflectra (inflixim	mponi cation (sarilu ise any iab-dy	Aria (golimumab) to any of the followin mab) ☐ Rinvoq (up v of the following pref yb) ☐ Simponi Aria	g? (select all adacitinib) [erred produc (golimumab)	that apply) Xeljanz/Xelja ts when indicat	ed for the patient's	
diagnosis? (select all that apply)	mab) 🗌 Kev	zara (sarilumab) 🔲	Rinvo	q (upadacitinib) 🔲 🛛	Xeljanz/Xelja	nz XR (tofacitin	ib)	
🗌 Yes 🔲 No 🛛 Will Rituxan (rituximab) be u	sed concomita	antly with apremilast,	tofacit	inib, or other biologic	DMARDs (e.	.g., adalimumal	b, infliximab)?	

9	6	tI	a
đ	C	L	d

MEDICARE FORM

Riabni[®] (rituximab-arrx), Rituxan[®] (rituximab), Ruxience (rituximab-pvvr), Truxima (rituximab-abbs) Medication Precertification Request Page 2 of 3

For Illinois MMP: FAX: 1-855-320-8445 PHONE: 1-866-600-2139

For other lines of business: Please use other form.

Note: Rituxan, Rituxan Hycela, and Truxima are preferred. Riabni and Ruxience are non-preferred. For rheumatoid arthritis, all Rituxan and biosimilar products are non-preferred.

(All fields must be completed and return both pages for precertification review.)

Patient First Name	Patient Last Name	Patient Phone	Patient DOB		
G CLINICAL INFORMATION (Continued)	Required clinical information must be complete	d for ALL procortification request			
Acute lymphoid leukemia		ed for ALL precertification request	5.		
	umented diagnosis of Philadelphia chromosom	e-negative acute lymphoid leukem	nia (ALL)?		
☐ Yes ☐ No Is Rituxan (rituximab) being u					
Autoimmune hemolytic anemia					
☐ Yes ☐ No Does the patient have a docu	umented diagnosis of refractory autoimmune he	emolytic anemia?			
Anti-neutrophil cytoplasmic antibody-asso					
Please indicate which of the following applies	to the patient: U Wegener granulomatosis	Churg-Strauss syndrome pauci-immune glomerulonepl	britio		
│ │			IIIus		
Autoimmune blistering diseases, corticost					
	umented diagnosis of corticosteroid-refractory	autoimmune blistering disease?			
	to the patient: 🔲 pemphigus vulgaris 🗌 pem		nphigoid 🔲 cicatricial pemphigoid		
	🗌 epidermolysis bullosa acquis	sita 🔲 paraneoplastic pemphigus	None of the above		
B-cell lymphomas					
] AIDS-related B-cell lymphoma 🔲 Burkitt lyr				
] Gastric MALT lymphoma 🛛 High-grade B-C] Nodal marginal zone lymphoma 🔲 Nongasi				
	Splenic marginal zone lymphoma Other:				
Castleman's disease					
Yes No Does the patient have a docu	umented diagnosis of multicentric Castleman's	disease (angiofollicular lymph noc	de hyperplasia)?		
Central nervous system lymphomas	, and the second s				
	leptomeningeal metastases from lymphoma	🗌 primary CNS lymphoma 🛛 n	one of the above		
Chronic or small lymphocytic leukemia		_			
	chronic lymphocytic leukemia (CLL)	l lymphocytic leukemia 🛛 none d	of the above		
Cryoglobulinemia	mented diagnosis of an aglabulinemia?				
 ☐ Yes ☐ No Does the patient have a documented diagnosis of cryoglobulinemia? ☐ Yes ☐ No Is there clinical documentation that the treatment with corticosteroids and other immunosuppressive agents was ineffective? 					
Graft versus host disease, chronic					
	t Rituxan (rituximab) being used as last-resort t	treatment for chronic graft versus	host disease (GVHD)?		
Hairy cell leukemia	. , _	-			
Please select which applies to the patient:	relapsed hairy cell leukemia 🔲 refractory ha	iry cell leukemia 🛛 none of the a	above		
Heart and solid organ transplant					
Yes No Is there a documentation that Rituxan (rituximab) is being used for treatment or prevention (desensitization) of highly sensitized patients with					
antibody mediated rejection in heart transplant recipients and other solid organ transplant recipients? Please select which applies to the patient: heart transplant recipient other solid organ transplant recipient					
Immune checkpoint-inhibitor related encep			spon		
Please identify which immune check-point inhibitor caused the encephalitis: Bavencio (avelumab) Imfinzi (durvalumab) Keytruda (pembrolizumab)					
	🗌 Opdivo (nivo	olumab) 🔲 Tecentriq (atezolizum	nab) 🔲 Yervoy (ipilimumab)		
	Other:				
Immune or idiopathic thrombocytopenic pu	•				
	umented diagnosis of refractory immune or idio		(ITP)?		
	mbocytopenic purpura 🔲 idiopathic thromboo	cytopenic purpura (ITP)			
	used as rejection prophylaxis in sensitized kidn	ey transplant recipients with dono	r specific antibodies?		
·	10ma umented diagnosis of lymphocyte-predominant	Hodgkin's lymphoma?			
Multiple Sclerosis					
Please indicate the type of multiple sclerosis	the patient has been diagnosed with: ondary-progressive MS (SPMS) 囗 Primary-p		essive-relansing MS (DRMS)		
	other medications used for treating MS (not in				



MEDICARE FORM

Riabni[®] (rituximab-arrx), Rituxan[®] (rituximab), Ruxience (rituximab-pvvr), Truxima (rituximab-abbs) Medication Precertification Request Page 3 of 3

For Illinois MMP: FAX: 1-855-320-8445 PHONE: 1-866-600-2139

For other lines of business: Please use other form.

Note: Rituxan, Rituxan Hycela, and Truxima are preferred. Riabni and Ruxience are non-preferred. For rheumatoid arthritis, all Rituxan and biosimilar products are non-preferred.

(All fields must be completed and return both pages for precertification review.)

Patient First Name	Patient Last Name	Patier	Patient Phone		Patie	Patient DOB			
G. CLINICAL INFORMATION (Continued)	- Required clinical information must be	completed for Al	l precertifi	cation r	requests				
Myasthenia gravis (MuSk-MG)		completed for Al		Cation	equests				
Yes No Does the patient have a doo		-		gravis (N	MuSK-N	IG)?			
Yes No Has the p	atient had an unsatisfactory response to	o initial immunoth	erapy?						
Neuromyelitis optica (Devic's disease)									
 ☐ Yes ☐ No Does the patient have a do ☐ Yes ☐ No Was the treatment with at I 		otica (Devic's dise	ease)?						
Opsocionus-myocionus-ataxia (opsocionu									
\square Yes \square No Does the patient have a do		oclonus-ataxia (O	MA) associ	ated wi	th neuro	blastor	ma?		
☐ Yes ☐ No Is the patient refractory to s									
Please provide the name	es and date ranges of medications tried	:							
Medication:			Dates:	/	1		/	1	-
Medication:			Dates:				1	1	-
			Dates:	/	/		/	1	-
Post-transplant lymphoproliferative disor		mah an reliferention	diagrada 20						
Yes No Is Rituxan (rituximab) being				\ noot t	rononlou	at lumanal	honrolife	arativa dia	ardarO
Rheumatoid Arthritis	n (rituximab) being used as prophylaxis	ior Epstein-barr	VIIUS (EDV) post-u	ranspiar	плитр	noprome	erative dis	sorder?
Please indicate the severity of the patient's	rheumatoid arthritis:	rate							
Yes No Is there evidence that the c									
☐ Yes ☐ No Will Rituxan (rituximab) be									
	tment with methotrexate ineffective, not			?					
Please sel	lect: ineffective in not tolerated] contraindicated							
☐ Yes ☐ No Was trea	tment with another conventional DMAR	D ineffective?							
	ect: azathioprine cyclosporine [quine 🗌 le	eflunom	nide 🗌	sulfasa	alazine		
Sjögren syndrome									
Yes No Does the patient have a do									
Yes No Was treatment with cortico				anto u					
	es and dates of the corticosteroids and o					-	1	1	
Medication:			Dates:	1	1		1	1	•
Thrombotic thrombocytopenic purpura									
Yes No Does the patient have a do	cumented diagnosis of refractory throm	botic thrombocyte	openic purp	ura (TT	P)?				
Waldenstrom's macroglobulinemia									
☐ Yes ☐ No Does the patient have a do	cumented diagnosis of Waldenström ma	acroglobulinemia	?						
For Continuation Requests:									
☐ Yes ☐ No Is this continuation request Please indicate the length of time on Rituxa	1 0 1	s of Rituxan (ritu	kimab)?						
For rheumatoid arthritis only:									
Please indicate the severity of the disease a		ituximab): 🗌 Milo	d 🗌 Mode	rate] Sever	e			
Yes No Is there clinical documenta									
☐ Yes ☐ No Is there clinical documenta	tion supporting disease improvement?								
For all other indications:	tion supporting dispass stability?								
☐ Yes ☐ No Is there clinical documenta									
H. ACKNOWLEDGEMENT									
							Deter	,	
Request Completed By (Signature Red							Date: _		/
Any person who knowingly files a reques									
any insurance company by providing ma insurance act, which is a crime and subje			AUOTI IOF THE	e purpo	use of h	nisiead	iiriy, col	mmus a i	nauuuent

The plan may request additional information or clarification, if needed, to evaluate requests.