♥aetr	Page 1 of 3	ce <sup>®</sup> (rituxima ition Precert	ification	•	Phone: 1 FAX: 1 For Medica	ertification Notification -866-752-7021 -888-267-3277 are Advantage Part B: Medicare Request Form	
		be completed and return bot			4 4		
	t of treatment, start date:	1 1		of therapy, date of las			
	sted By:		Pho	one:	Fax:		
A. PATIENT INFORMA	TION						
First Name:		Last Name:					
Address:	1	I	City:		State:	ZIP:	
Home Phone:	Work Phone:	Cell Phone:	[	OOB:	E-mail:		
Current Weight: Ib	os or kgs Height:	inches orcms	Allergies:				
<b>B. INSURANCE INFOR</b>	MATION						
Member ID #:		Does patient have o	ther coverage?	🗌 Yes 🔲 No			
Group #:		If yes, provide ID#:		Carrier Name:			
Insured:		Insured:					
Medicare: 🗌 Yes 🗌 N	lo If yes, provide ID #:		Medicaid: 🗌 Ye	s 🗌 No 🛛 If yes, provi	de ID #:		
C. PRESCRIBER INFO	RMATION						
First Name:		Last Name:		(Check one):	□ M.D. □	D.O. 🗌 N.P. 🗌 P.A.	
Address:			City:		State:	ZIP:	
Phone:	Fax:	St Lic #:	NPI #:	DEA #:		UPIN:	
Provider E-mail:		Office Contact Name	e:		Phone:		
Specialty (Check one)	🗌 Rheumatologist 🔲 Onco	ologist					
	IDER/ADMINISTRATION INF						
Center Name: Home Infusion Center Agency Name: Administration code(s Address: E. PRODUCT INFORM	Physician's Office Center Phone: er Phone: s) (CPT): ATION		□ Physicia           □ Specialt           □ Specialt           □ Address: _           Phone:	y Pharmacy	Retail Pharma Other Fax: _	cy	
	nce (rituximab-pvvr) <b>Dose:</b>			for Use:			
	MATION - Please indicate prim			er where applicable (*).			
Primary ICD Code:			er ICD Code:				
G. CLINICAL INFORMATION - Required clinical information must be completed for ALL precertification requests.         For All Requests (clinical documentation required for all requests) :         Yes       No         Has the patient failed treatment with Truxima due to a documented intolerable adverse event (e.g., rash, nausea, vomiting)?         Yes       No         Was the adverse event unexpected and not attributed to the active ingredient as described in the prescribing information (i.e., known adverse reaction for both the brand and biosimilar medication)?         Non-Oncology       As part of a non-myeloablative conditioning regimen for allogeneic transplant         Autoimmune hemolytic anemia       Autoimmune bilistering diseases         Please select which applies to the patient:       pemphigus vulgaris       pemphigus foliaceus       bullous pemphigoid							
<ul> <li>☐ Chronic graft versu</li> <li>☐ Cryoglobulinemia</li> <li>☐ Yes ☐ No Hav</li> <li>☐ Churg-Strauss synd</li> <li>☐ Granulomatosis witi</li> <li>☐ Immune Checkpoin</li> <li>☐ Immune or idiopath</li> <li>☐ Microscopic polyar</li> <li>☐ Multiple sclerosis (I</li> <li>☐ Yes ☐ No Has</li> </ul>	epic epic epices host disease ve corticosteroids and other im drome th polyangiitis (GPA) (Wegen th Inhibitor-related toxicities nic thrombocytopenic purput ngiitis (MPA) MS) s the patient been diagnosed of	dermolysis bullosa acquisit munosuppressive agents ner's granulomatosis) ra (ITP), refractory vith relapsing-remitting mu	ta ☐ paraneoplas been ineffective? Itiple sclerosis (RF	stic pemphigus 🗍 no	ne of the abov	re	
☐ Yes ☐ No Is th ☐ Myasthenia gravis,	he patient taking the requester <b>refractory</b>	d medication with any othe	r medication used	for the treatment of m	ultiple sclerosi	s other than Ampyra?	



## Ruxience<sup>™</sup> (rituximab-pvvr) Medication Precertification Request Page 2 of 3

 Aetna Precertification Notification

 Phone:
 1-866-752-7021

 FAX:
 1-888-267-3277

For Medicare Advantage Part B: Please Use Medicare Request Form

(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) - Reg	uired clinical information must be completed for A	LL precertification requests					
☐ Yes ☐ No Has the patient experienced	methotrexate dose greater than or equal to 1 an inadequate response with another convention		loroquine, leflunomide,				
☐ Yes       No       Has the patient experienced an inadequate response with another conventional DMARD (e.g., hydroxychloroquine, leflunomide, sulfasalazine)?         ☐ Yes       No       Is the requested drug being prescribed in combination with methotrexate or leflunomide?         ☐ Yes       No       Has the patient experienced intolerance to methotrexate or leflunomide?         ☐ Yes       No       Has the patient experienced intolerance to methotrexate or leflunomide?         ☐ Yes       No       Does the patient have a contraindication to methotrexate or leflunomide?         ☐ Please indicate the contraindication:       ☐ History of intolerance or adverse event ☐ Renal impairment ☐ Hypersensitivity         ☐ Blood dyscrasias (e.g., thrombocytopenia, leukopenia, significant anemia)       ☐ Breastfeeding ☐ Elevated liver transaminases ☐ Myelodysplasia         ☐ Interstitial pneumonitis or clinically significant pulmonary fibrosis       ☐ Pregnancy or currently planning pregnancy ☐ Significant drug interaction         ☐ Clinical diagnosis of alcohol use disorder, alcoholic liver disease or other chronic liver disease							
☐ Yes ☐ No Will the requested drug be used with another biologic for the treatment of rheumatoid arthritis? ☐ Yes ☐ No Is the planned date of administration at least 16 weeks after the date of the last dose received?							
<ul> <li>Sjögren's syndrome</li> <li>Yes No Have corticosteroids and other immunosuppressive agents been ineffective?</li> <li>Solid organ transplant and prevention of antibody mediated rejection in solid organ transplant</li> <li>Yes No Is the requested drug being used for the treatment and prevention of antibody mediated rejection in solid organ transplant?</li> <li>Systemic Lupus Erythematosus (SLE)</li> <li>Yes No Is the disease refractory to immunosuppressive therapy?</li> <li>Thrombotic thrombocytopenic purpura (TTP)</li> </ul>							
	-related B-cell lymphoma ptomeningeal metastases from lymphomas	-	of the B-cell.				



## Ruxience<sup>™</sup> (rituximab-pvvr) Medication Precertification Request Page 3 of 3

 Aetna Precertification Notification

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## For Medicare Advantage Part B: Please Use Medicare Request Form

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

Patient First Name	Patient Last Name	Patient Phone	Patient DOB				
	d) - Required clinical information must be cc	ompleted for ALL precertification request	S.				
<ul> <li>Chronic lymphocytic leukemia (CLL)</li> <li>Diffuse large B-cell lymphoma (DLBCL)</li> </ul>							
Follicular lymphoma	-)						
Hairy cell leukemia							
☐ High-grade B-cell lymphoma with translocations of MYC and BCL2 and/or BCL6 (double/triple hit lymphoma)							
High-grade B-cell lymphoma, not othe	rwise specified						
Histological transformation of indolent	lymphomas to diffuse large B-cell lymphoma	a					
Hodgkin's lymphoma, nodular lymphocyte-predominant							
Mantle cell lymphoma							
Marginal zone lymphomas (nodal marg splenic marginal zone lymphoma)	ginal zone lymphoma, gastric mucosa assoc	ciated lymphoid tissue (MALT) lymphom	a, nongastric MALT lymphoma,				
Pediatric Aggressive Mature B-Cell Ly	mphomas						
Primary cutaneous B-cell lymphoma							
Post-transplant lymphoproliferative dis	order (PTLD)						
Rosai-Dorfman disease							
Small lymphocytic lymphoma (SLL)	anh an leann an tig luman hanna (I DI )						
Waldenström's macroglobulinemia/lym							
For Continuation Requests (clinical doo							
	atment with Truxima due to a documented in						
	adverse event unexpected and not attribute ion (i.e., known adverse reaction for both the		n the prescribing				
Rheumatoid Arthritis (RA)							
	the patient has received since starting treat						
	ation be used with another biologic for the tr						
•	tivity improvement from baseline in tender jo	oint count, swollen joint count, pain, or d	sability: %				
Multiple sclerosis (MS)							
	ng disease stability or improvement while rec	eiving the requested medication?					
Continuation, oncologic indications							
Yes No Is there evidence of unac							
	bitor-related toxicities and all other indica	ations					
Yes No Is the patient experiencin	g benefit from therapy?						
H. ACKNOWLEDGEMENT							
Request Completed By (Signature R	equired):		Date: / /				
	lest for authorization of coverage of a me naterially false information or conceals ma						

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

insurance act, which is a crime and subjects such person to criminal and civil penalties.