♥aetna	Truxima Medicati Page 1 of 3	•				quest	Phone: FAX: For Media	ecertification Notification 1-866-752-7021 1-888-267-3277 care Advantage Part B:
	(All fields must be c			-				se Medicare Request Form
Please indicate: Start of tre		/ /	[Continua	tion of thera	py, date of l	ast treatment:	
Precertification Requested B	y:				Phone:		Fax	K:
A. PATIENT INFORMATION								
First Name:		Last Nam	ne:	1			1	
Address:	- 1			City:			State:	ZIP:
Home Phone:	Work Phone:		Cell Phone:		C	DOB:	E-mail:	
Current Weight: lbs or _	kgs Height:	inches or	cms	Allergies:				
B. INSURANCE INFORMATIO	N							
Member ID #:		Does patient have other coverage?						
Group #:			If yes, provide ID#: Carrier Name: _					
Insured:		Insured:						
Medicare: Yes No If ye	es, provide ID #:		М	edicaid:	Yes 🗌 N	o If yes, pro	ovide ID #:	
C. PRESCRIBER INFORMATI	ON						_	
First Name:		Last Nam	ne:	1		(Check one	e): 🗌 M.D. [□ D.O. □ N.P. □ P.A.
Address:				City:			State:	ZIP:
Phone:	Fax:	St Lic #:		NPI #:		DEA #:		UPIN:
Provider E-mail:		Office Co	ontact Name:				Phone:	
Specialty (Check one): Rhe	umatologist 🔲 Oncolog	jist 🗌 Oth	ner:					
Place of Administration: Self-administered Pł Outpatient Infusion Center Center Name: Home Infusion Center Agency Name: Administration code(s) (CPT Address:	Phone:			☐ Pf _ ☐ Sp _ Name - Addre - Phon	nysician's Offi pecialty Pharr e: ess: e:	ice [nacy [Fax:	
Request is for: Truxima (r	-				ns for Use:			
F. DIAGNOSIS INFORMATION	I - Please indicate primary	ICD code a	nd specify an	y other an	y other where	e applicable	(*).	
Primary ICD Code:				her ICD C				
G. CLINICAL INFORMATION	- Required clinical informat	ion must be	completed for	or ALL pred	ertification re	equests.		
For All Requests (clinical documentation required for all requests): Non-Oncology As part of a non-myeloablative conditioning regimen for allogeneic transplant Autoimmune hemolytic anemia Autoimmune blistering diseases Please select which applies to the patient: pemphigus foliaceus								
 □ epidermolysis bullosa acquisita □ paraneoplastic pemphigus □ none of the above □ Chronic graft versus host disease □ Cryoglobulinemia □ Yes □ No Have corticosteroids and other immunosuppressive agents been ineffective? □ Churg-Strauss syndrome □ Granulomatosis with polyangiitis (GPA) (Wegener's granulomatosis) □ Immune Checkpoint Inhibitor-related toxicities □ Immune or idiopathic thrombocytopenic purpura (ITP), refractory □ Multiple sclerosis (MS) □ Yes □ No Has the patient been diagnosed with relapsing-remitting multiple sclerosis (RRMS)? □ Yes □ No Is the patient taking the requested medication with any other medication used for the treatment of multiple sclerosis other than Ampyra? 								
 Myasthenia gravis, refractory Neuromyelitis optica (i.e., neuromyelitis optica spectrum disorder (NMOSD), Devic disease) Yes No Will the requested drug be used concomitantly with another biologic for the treatment of neuromyelitis optica spectrum disorder (NMOSD)? 								

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Truxima[™] (rituximab-abbs) Medication Precertification Request

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

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For Medicare Advantage Part B: Please Use Medicare Request Form

Patient First Name	Patient Last Name	Patient Phone	Patient DOB			
G. CLINICAL INFORMATION (Continued) - Requ	uired clinical information must be completed for AL	L precertification requests.				
C. CLINICAL INFORMATION (Continued) - Required clinical information must be completed for ALL precertification requests. C. Opsocionus-myocionus ataxia C. Yes No Is the requested drug being used for associated opsocionus-myocionus ataxia associated with neuroblastoma? C. Yes No Is the patient refractory to steroids and chemotherapy? Pauci-immune glomerulonephritis Prevention of Epstein-Barr virus (EBV) related post-transplant lymphoproliferative disorder (PTLD) Rheumatoid arthritis (RA) C. Yes No Has the patient been diagnosed with moderately to severely active rheumatoid arthritis (RA)? C. Yes No Has the patient received a biologic DMARD or targeted synthetic DMARD (e.g., Xeljanz) that is indicated for moderately to severely active rheumatoid arthritis? C. Yes No Has the patient received two full doses of the requested medication, with the most recent dose being 6 months before this request? C. Yes No Has the patient been tested for the rheumatoid factor (RF) biomarker? Please indicate the test result: Dositive Degative Degative Disorder (CRP) biomarker? Please indicate the test result: Desitive Degative Degative Disorder (CRP) biomarker? Please indicate the test result: Desitive Degative Disorder (CRP) biomarker? Please indicate the test result: Desitive Disorder (CRP) biomarker? Please indicate the test result: Desitive Disorder (CRP) biomarker? Please indicate the test result: Desitive Disorder (CRP) biomarker? Please indicate the test result: Desitive Disorder (CRP) biomarker? Please indicate the test result: Desitive Disorder (CRP) biomarker? Please indicate the test result: Desitive Disorder (CRP) biomarker? Please indicate the test result: Disorder Disorder Disorder (CRP) biomarker? Please indicate the test result: Disorder Disorder Disorder Disorder (CRP) biomarker? Please indicate the test result: Disorder Disorde						
☐ Yes ☐ No Has the patient experienced sulfasalazine)? ☐ Yes ☐ No Is the requested drug being p ☐ Yes ☐ No Has the patie	 No Has the patient experienced an inadequate remethotrexate dose greater than or equal to 15 an inadequate response with another conventional prescribed in combination with methotrexate or left experienced intolerance to methotrexate or left No Does the patient have a contraindication to methotrexate indicate the contraindication: ☐ History of intolerance or adverse event ☐ Blood dyscrasias (e.g., thrombocytopenia, ☐ Breastfeeding ☐ Elevated liver transamin ☐ Interstitial pneumonitis or clinically signification or clinical diagnosis of alcohol use disorder, and other: 	5 mg per week? al DMARD (e.g., hydroxychloroq lunomide? lunomide? ethotrexate or leflunomide?] Renal impairment ☐ Hypers leukopenia, significant anemia) nases ☐ Myelodysplasia ant pulmonary fibrosis y ☐ Significant drug interaction alcoholic liver disease or other o	uine, leflunomide, ensitivity n			
 ☐ Yes ☐ No Is the planned date of admining ☐ Sjögren's syndrome ☐ Yes ☐ No Have corticosteroids and oth ☐ Solid organ transplant and prevention of and ☐ Yes ☐ No Is the requested drug being to ☐ Systemic Lupus Erythematosus (SLE) ☐ Yes ☐ No Is the disease refractory to in ☐ Thrombotic thrombocytopenic purpura (TT Oncology: ☐ Yes ☐ No Does the patient have CD20 position 	used for the treatment and prevention of antibody in munosuppressive therapy? P) tive disease that was confirmed by testing or anal or results of testing or analysis confirming CD20 is: related B-cell lymphoma	t dose received? splant mediated rejection in solid organ lysis?				



Truxima[™] (rituximab-abbs) Medication Precertification Request

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For Medicare Advantage Part B: Please Use Medicare Request Form

Patient First Name	Patient Last Name	Patient Phone	Patient DOB			
G. CLINICAL INFORMATION (Continued) - Red Hairy cell leukemia High-grade B-cell lymphoma with translocatio	ns of MYC and BCL2 and/or BCL6		sts.			
 High-grade B-cell lymphoma, not otherwise sp Histological transformation of indolent lympho Hodgkin's lymphoma, nodular lymphocyte-pre Mantle cell lymphoma Marginal zone lymphomas (nodal marginal zone splenic marginal zone lymphoma) Pediatric Aggressive Mature B-Cell Lymphom Post-transplant lymphoproliferative disorder (f Primary cutaneous B-cell lymphoma Rosai-Dorfman disease Small lymphocytic lymphoma (SLL) 	mas to diffuse large B-cell lymphor dominant ne lymphoma, gastric mucosa asso as		na, nongastric MALT lymphoma,			
Waldenström's macroglobulinemia/lymphoplasmacytic lymphoma (LPL)						
For Continuation Requests (clinical document Rheumatoid Arthritis (RA) Please indicate the number of total doses the para Yes No Will the requested medication be Please indicate the percent of disease activity im Multiple sclerosis (MS) Yes No Is the patient experiencing disea Continuation, oncologic indications Yes No Has the patient experienced and Continuation, immune checkpoint inhibitor-re Yes No Is the patient experiencing benefitied	tient has received since starting trea o used with another biologic for the provement from baseline in tender se stability or improvement while re unacceptable toxicity from treatmen lated toxicities and all other indi-	treatment of rheumatoid arthritis? joint count, swollen joint count, pain, or o ceiving the requested medication? t with the requested drug?				
H. ACKNOWLEDGEMENT						
Request Completed By (Signature Require	d):		Date: / / /			
Any person who knowingly files a request for authorization of coverage of a medical procedure or service with the intent to injure, defraud or deceive						

Any person who knowingly files a request for authorization of coverage of a medical procedure or service with the intent to injure, defraud or deceive any insurance company by providing materially false information or conceals material information for the purpose of misleading, commits a fraudulent insurance act, which is a crime and subjects such person to criminal and civil penalties.

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