

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.)

Virginia (HMO D-SNP)
FAX: 1-833-280-5224
PHONE: 1-855-463-0933
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.

Procest interest (Price of Agricultural Information New John Regulated By: Last Name: Last Name: Call Phone: E-mail:	Please indicate:	☐ Start of treatme	nt, start da	te:	1 1	□ C	ontinuation of th	nerapy, date of la	st treatment	:	
	Precertification	Requested By:					Phone:		Fa	X:	
	A. PATIENT IN	FORMATION									
Home Phone:					Last Name:					DOB:	
Current Weight:	Address:				1	City:			State:	ZIP:	
Does patient have other coverage? Yes No	Home Phone:		Work Pl	none:		,			E-mail:		
Member ID #:	Current Weight:	Ibs or	kgs I	leight:	inches or	cms	Allergies:		.1		
fives.provide D#:	B. INSURANCE	INFORMATION									
fives.provide D#:	Member ID #:				Does patient hav	e other o	overage?	☐ Yes ☐ No			
Insured:					•		-	Carrier Name: _			
Last Name: Check one): M.D. D.O. N.P. P.	Insured:				Insured:						
Address: City: DEA #: UPIN:	C. PRESCRIBE	R INFORMATION									
Phone: Fax: St Lic #: NPI #: DEA #: UPIN:	First Name:				Last Name:			(Check one)): M.D. [☐ D.O. ☐ N.P. ☐ P.A.	
Provider Email: Office Contact Name: Phone:	Address:					City:			State:	ZIP:	
Dispensing Provider/Pharmacy Gelf-administration Gelf-admini	Phone:	Fax:		St Lic	#:	NPI :	# :	DEA #:		UPIN:	
Rece of Administration:	Provider Email:	•		Office	Contact Name:			Phone:		•	
Rece of Administration:	D. DISPENSING	G PROVIDER/ADMIN	IISTRATIO	N INFORM	ATION						
Request is for: Riabni (rituximab-arrx) Rituxan (rituximab) Ruxience (rituximab-pvvr) Truxima (rituximab-abbs) Dose: Directions for Use: HCPCS Code: F. DIAGNOSIS INFORMATION - Please indicate primary ICD code and specify any other any other where applicable (*). Primary ICD Code: Other ICD Code: G. CLINICAL INFORMATION - Required clinical information must be completed for ALL precertification requests. For All Requests (clinical documentation required for all requests): Note: Rituxan, Rituxan, Rituxan All ycola, and Truxima are preferred for most indications. Riabni and Ruxience are non-preferred. For rheumatoid arthritis, all Rituxan and biosimilar products are non-preferred. Inflectra, Remicade, and Simponi Aria are preferred for MAPD plans. Enbrel, Humira, Kevzara, Rinvoq, and Xeljanz/Xeljanz XR are preferred for MAPD plans. Pyes No Has the patient had prior therapy with Riabni (rituximab-arax) or Ruxience (rituximab-pvvr) within the last 365 days? Yes No Has the patient had a trial and failure, intolerance, or contraindication to any of the following? (select all that apply) Rituxan (rituximab) Rituxan Hycela (rituximab/hyaluronidase human) Truxima (rituximab-abbs) Please explain if there are any other medical reason(s) that the patient cannot use any of the following preferred products when indicated for the patient's diagnosis? (select all that apply) Rituxan (rituximab) Rituxan Hycela (rituximab/hyaluronidase human) Truxima (rituximab-abbs) Yes No Has the patient had a trial and failure, intolerance, or contraindication to any of the following? (select all that apply) Remicade (infliximab) Inflectra (infliximab-dyyb) Simponi Aria (golimumab) Yes No Has the patient had a trial and failure, intolerance, or contraindication to any of the following? (select all that apply) Enbrel (etanercept) Humira (adalimumab) Kevzara (sarilumab) Rinvoq (upadactitnib) Rinvoq (upadactitnib) Rinvoq (upadactitnib) Rinvoq (upadactitnib) Rinvoq (upadactitnib	☐ Outpatient Infusion Center Name: ☐ Home Infusion Center Phone: Agency Name:				ZIP:	Retail Pharmacy Mail Order Name: Address: City: Phone: TIN:			Specialty Pharmacy Other: State: Fax:		
Directions for Use:											
F. DIAGNOSIS INFORMATION - Please indicate primary ICD code and specify any other any other where applicable (*). Primary ICD Code:	-	: 🗌 Riabni (rituxim	ab-arrx)								
Ctinical Information - Required clinical information must be completed for ALL precertification requests.										6 Code:	
G. CLINICAL INFORMATION - Required clinical information must be completed for ALL precertification requests. For All Requests (clinical documentation required for all requests): Note: Rituxan, Rituxan Hycela, and Truxima are preferred for most indications. Riabni and Ruxience are non-preferred. For rheumatoid arthritis, all Rituxan and biosimilar products are non-preferred. Inflectra, Remicade, and Simponi Aria are preferred for MA plans. Enbrel, Humira, Kevzara, Rinvoq, and Xeljanz/Xeljanz XR are preferred for MAPD plans. Yes No Has the patient had prior therapy with Riabni (rituximab-arrx) or Ruxience (rituximab-pvvr) within the last 365 days? Hease explain if there are any other medical reason(s) that the patient cannot use any of the following preferred products when indicated for the patient's diagnosis? (select all that apply) Remicade (infliximab) Rituxan Hycela (rituximab/hyaluronidase human) Truxima (rituximab-abbs) The Remicade (infliximab) Inflectra (infliximab-dyyb) Simponi Aria (golimumab) Remicade (infliximab) Inflectra (infliximab-dyyb) Rituxan (golimumab) Rituxan (golimumab) Remicade (infliximab) Remicade (infliximab) Remicade (infliximab) Rituxan Hycela (infliximab-dyyb) Simponi Aria (golimumab) Remicade (infliximab) Remicade (infliximab) Rituxan Hycela (infliximab-dyyb) Simponi Aria (golimumab) Remicade (infliximab) Remicade (infliximab) Rituxan Hycela (infliximab-dyyb) Simponi Aria (golimumab) Remicade (infliximab) Remicade (infliximab) Rituxan Hycela (infliximab-dyyb) Simponi Aria (golimumab) Remicade (infliximab) Remicade (infliximab) Remicade (infliximab) Rituxan Hycela (infliximab-dyyb) Simponi Aria (golimumab)						ify any o	ther any other w	here applicable (*	').		
Note: Rituxan, Rituxan Hycela, and Truxima are preferred for most indications. Riabni and Ruxience are non-preferred. For rheumatoid arthritis, all Rituxan and biosimilar products are non-preferred. Inflectra, Remicade, and Simponi Aria are preferred for MA plans. Enbrel, Humira, Kevzara, Rinvoq, and Xeljanz/Xeljanz XR are preferred for MAPD plans. Yes	Primary ICD C	ode:				Other	ICD Code:				
Note: Rituxan, Rituxan Hycela, and Truxima are preferred for most indications. Riabni and Ruxience are non-preferred. For rheumatoid arthritis, all Rituxan and biosimilar products are non-preferred. Inflectra, Remicade, and Simponi Aria are preferred for MA plans. Enbrel, Humira, Kevzara, Rinvoq, and Xeljanz/Xeljanz XR are preferred for MAPD plans. Yes No Has the patient had prior therapy with Riabni (rituximab-arrx) or Ruxience (rituximab-pvvr) within the last 365 days? Rituxan (rituximab) Rituxan Hycela (rituximab/hyaluronidase human) Truxima (rituximab-abbs) Please explain if there are any other medical reason(s) that the patient cannot use any of the following preferred products when indicated for the patient's diagnosis? (select all that apply) Rituxan (rituximab) Rituxan Hycela (rituximab/hyaluronidase human) Truxima (rituximab-abbs) Please explain if there are any other medical reason(s) that the patient cannot use any of the following? (select all that apply) Remicade (infliximab) Inflectra (infliximab-dyyb) Simponi Aria (golimumab) Remicade (infliximab) Revzara (sarilumab) Rituxon (upadacitinib) Related for the patient's diagnosis? (select all that apply) Remicade (infliximab) Remicade (infliximab) Revzara (sarilumab) Rituxon (upadacitinib) Related for the patient's diagnosis? (select all that apply) Remicade (infliximab) Inflectra (infliximab-dyyb) Simponi Aria (golimumab) Please explain if there are any other medical reason(s) that the patient cannot use any of the following preferred products when indicated for the patient's diagnosis? (select all that apply)	G. CLINICAL IN	NFORMATION - Req	uired clinica	l informatio	on must be comple	ted for A	LL precertificatio	on requests.			
Remicade (infliximab)	Note: Rituxan, For rheumatoic Enbrel, Humira Yes No Yes No	Rituxan Hycela, and arthritis, all Rituxa, Kevzara, Rinvoq, a Has the patient had Has the patient had Rituxan (rituxima if there are any other	d Truxima a and and Xeljana prior therap a trial and fab)	are preferrimilar procedure imilar procedure imilar procedure imilare, intologican Hycela ison(s) that	ed for most indic ducts are non-pre (R are preferred fo oni (rituximab-arrx) erance, or contrair i (rituximab/hyaluro t the patient canno	eferred. or MAPI or Ruxion indication onidase I	Inflectra, Remic Diplans. Price (rituximab-p to any of the foll- numan) Trux Trux Trux Trux Trux	eade, and Simpo ovvr) within the last lowing? (select all kima (rituximab-ak preferred produc	ni Aria are pr st 365 days? that apply) obs) cts when indic	ated for the patient's	
diagnosis? (select all that apply)	☐ Yes ☐ No	☐ Remicade (inflixi Has the patient had ☐ Enbrel (etanerce if there are any other	mab) ☐ Ir a trial and fa pt) ☐ Hun medical rea	iflectra (infl ailure, intol nira (adalim ison(s) that	iximab-dyyb) ☐ : erance, or contrair numab) ☐ Kevzai t the patient canno	Simponi ndication ra (sarilu t use an	Aria (golimumab to any of the follomab)	o) lowing? (select all q (upadacitinib) ı preferred produc	that apply) Xeljanz/Xets when indic		
Yes No Will Rituxan (rituximab) be used concomitantly with apremilast, tofacitinib, or other biologic DMARDs (e.g., adalimumab, infliximab)?	diagnosis? (sele	ect all that apply) ercept)	adalimumal	o) 🗌 Kevz	zara (sarilumab) [☐ Rinvo	q (upadacitinib)	Xeljanz/Xelja	ınz XR (tofacit	tinib)	



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

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB					
G. CLINICAL INFORMATION (Continued) - Required clinical information must be completed for ALL precertification requests.								
Acute lymphoid leukemia Yes No Does the patient have a documented diagnosis of Philadelphia chromosome-negative acute lymphoid leukemia (ALL)? Yes No Is Rituxan (rituximab) being used as induction/consolidation therapy?								
Autoimmune hemolytic anemia Yes No Does the patient have a docu	mented diagnosis of refractory autoimmune he	emolytic anemia?						
Anti-neutrophil cytoplasmic antibody-association	ciated (ANCA-associated) vasculitides							
Please indicate which of the following applies	☐ microscopic polyangiitis	☐ Churg-Strauss syndrome☐ pauci-immune glomeruloneph	ritis					
☐ Yes ☐ No Will Rituxan (rituximab) be giv	ven in conjunction with glucocorticoids?							
Autoimmune blistering diseases, corticosteroid-refractory Yes No Does the patient have a documented diagnosis of corticosteroid-refractory autoimmune blistering disease? Please select which applies to the patient: pemphigus vulgaris pemphigus folliaceus bullous pemphigoid cicatricial pemphigoid epidermolysis bullosa acquisita paraneoplastic pemphigus None of the above								
B-cell lymphomas								
	Please select which applies to the patient: AIDS-related B-cell lymphoma Burkitt lymphoma Diffuse large B-cell lymphoma Follicular lymphoma Gastric MALT lymphoma High-grade B-Cell lymphoma Mantle cell lymphoma Nodal marginal zone lymphoma Nongastric MALT lymphoma Primary cutaneous B-cell lymphomas Splenic marginal zone lymphoma Other:							
Castleman's disease								
☐ Yes ☐ No Does the patient have a docu	mented diagnosis of multicentric Castleman's	disease (angiofollicular lymph nod	e hyperplasia)?					
Central nervous system lymphomas Please select which applies to the patient:	leptomeningeal metastases from lymphoma	☐ primary CNS lymphoma ☐ no	one of the above					
Chronic or small lymphocytic leukemia	chronic lymphopytic loukomic (CLL)	lumphositic loukemie. ¬ none e	of the above					
Please select which applies to the patient: chronic lymphocytic leukemia (CLL) small lymphocytic leukemia none of the above cryoglobulinemia Yes No Does the patient have a documented diagnosis of cryoglobulinemia?								
	n that the treatment with corticosteroids and ot	her immunosuppressive agents w	as ineffective?					
Graft versus host disease, chronic Yes No Is there a documentation that Rituxan (rituximab) being used as last-resort treatment for chronic graft versus host disease (GVHD)?								
Hairy cell leukemia Please select which applies to the patient: ☐ relapsed hairy cell leukemia ☐ refractory hairy cell leukemia ☐ none of the 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 encephalitis								
	ibitor caused the encephalitis: ☐ Bavencio (av ☐ Opdivo (nivo ☐ Other:							
Immune or idiopathic thrombocytopenic purpura Yes No Does the patient have a documented diagnosis of refractory immune or idiopathic thrombocytopenic purpura (ITP)? refractory immune thrombocytopenic purpura idiopathic thrombocytopenic purpura (ITP)								
Kidney transplant, rejection prophylaxis ☐ Yes ☐ No Is Rituxan (rituximab) being used as rejection prophylaxis in sensitized kidney transplant recipients with donor specific antibodies?								
Lymphocyte-predominant Hodgkin's lymphoma ☐ Yes ☐ No Does the patient have a documented diagnosis of lymphocyte-predominant Hodgkin's lymphoma?								
Multiple Sclerosis								
Please indicate the type of multiple sclerosis the patient has been diagnosed with: Relapsing-remitting MS (RRMS) Secondary-progressive MS (SPMS) Primary-progressive MS (PPMS) Progressive-relapsing MS (PRMS) No Has the patient discontinued other medications used for treating MS (not including Ampyra)?								
☐ 163 ☐ 140 Has the patient discontinued	other medications used for treating ivid (flot flit	nading Ampyra):						

Continued on next page



MEDICARE FORM

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

Page 3 of 3

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Patient First Name	Patient Last Name Patient F		ent Phone P			Patient DOB			
G. CLINICAL INFORMATION (Continued) - Required clinical information must be completed for ALL precertification requests.									
Myasthenia gravis (MuSk-MG)									
Yes No Doe's the patient have a documented diagnosis of muscle-specific tyrosine kinase myasthenia gravis (MuSK-MG)?									
Yes No Has the patient had an unsatisfactory response to initial immunotherapy?									
Neuromyelitis optica (Devic's disease)									
☐ Yes ☐ No Does the patient have a docu	Yes No Does the patient have a documented diagnosis of neuromyelitis optica (Devic's disease)?								
☐ Yes ☐ No Was the treatment with at lea	ast one immunotherapy ineffective?								
Opsoclonus-myoclonus-ataxia (opsoclonus	myoclonus syndrome)								
Yes No Does the patient have a doc	umented diagnosis of opsoclonus-my	oclonus-ataxia (O	MA) assoc	iated wi	th neurob	lastoma?			
Yes No Is the patient refractory to ste			s?						
Please provide the names									
Medication:			Dates:	/	1	/			
Medication:			Dates:	/	/	/	/		
Medication:			Dates:	/		/_	/		
Post-transplant lymphoproliferative disord	er								
☐ Yes ☐ No Is Rituxan (rituximab) being t									
└────────────────────────────────────	(rituximab) being used as prophylaxi	s for Epstein-Barr	virus (EBV	/) post-tı	ransplant	lymphopro	liferative	disorder?	
Rheumatoid Arthritis									
Please indicate the severity of the patient's rh		erate Severe							
Yes No Is there evidence that the dis									
Yes No Will Rituxan (rituximab) be us				10					
	nent with methotrexate ineffective, no ct:			1?					
Please sele	ct. menective not tolerated								
☐ Yes ☐ No. Was treatr	ment with another conventional DMAF	RD ineffective?							
	ct: azathioprine cyclosporine		quine 🔲 l	eflunom	ide 🗌 s	ulfasalazin	е		
Sjögren syndrome	_ , _ , ,	_ , ,	. –						
Yes No Does the patient have a doct	umented diagnosis of Sjögren's synd	rome?							
☐ Yes ☐ No Was treatment with corticost			?						
Please provide the names									
Medication:			Dates:	/	1	/			
Medication:			Dates:	/		/_			
Thrombotic thrombocytopenic purpura									
Yes No Does the patient have a documented diagnosis of refractory thrombotic thrombocytopenic purpura (TTP)?									
Waldenstrom's macroglobulinemia									
☐ Yes ☐ No Does the patient have a documented diagnosis of Waldenström macroglobulinemia?									
For Continuation Requests:									
☐ Yes ☐ No Is this continuation request a		es of Rituxan (ritux	kimab)?						
Please indicate the length of time on Rituxan	(rituximab):						-		
For rheumatoid arthritis only:									
Please indicate the severity of the disease at baseline (pretreatment with Rituxan (rituximab): Mild Moderate Severe Yes No Is there clinical documentation supporting disease stability?									
Yes No Is there clinical documentation	•								
For all other indications:	on supporting disease improvement:								
Yes ☐ No Is there clinical documentation	on supporting disease stability?								
☐ Yes ☐ No Is there clinical documentation									
	an supporting disease improvement:								
H. ACKNOWLEDGEMENT									
Request Completed By (Signature Requ	ıired):					Date:		<u> </u>	
Any person who knowingly files a request any insurance company by providing mate insurance act, which is a crime and subject	erially false information or conceals	s material informa							

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