SENTARA COMMUNITY PLAN (MEDICAID)

MEDICAL PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

<u>Directions:</u> The prescribing physician must sign and clearly print name (preprinted stamps not valid) on this request. All other information may be filled in by office staff; <u>fax to 1-844-305-2331</u>. No additional phone calls will be necessary if all information (<u>including phone and fax #s</u>) on this form is correct. <u>If information provided is not complete, correct, or legible, authorization can be delayed</u>.

FOR AN ONCOLOGY DIAGNOSIS

The Sentara Community Plan Oncology Program is administered by Carelon Medical Benefits Management

For any Oncology indications including the use of Rituxan Hycela, the most efficient way to submit a prior authorization request is through the Carelon Provider Portal at www.providerportal.com

Rituxan[®] (rituximah) (19312)

<u>Drug Requested</u>: Select drug below (Medical) (Non-Preferred)

□ Riahni[™] (rituximah-arrx) (O5123)

- Trabin (maximas anx) (20120)				
□ Ruxience [™] (rituximab-pvvr) (Q5119)	□ Truxima® (rituximab-abbs) (Q5115)			
MEMBER & PRESCRIBER INFORMAT	ION: Authorization may be delayed if incomplete.			
Member Name:				
Member Sentara #:	ber Sentara #: Date of Birth:			
Prescriber Name:				
Prescriber Signature:	Date:			
Office Contact Name:				
Phone Number:	hone Number: Fax Number:			
DEA OR NPI #:				
DRUG INFORMATION: Authorization may b	pe delayed if incomplete.			
Drug Form/Strength:				
Dosing Schedule:	Length of Therapy:			
Diagnosis:	ICD Code, if applicable:			
Weight:	Date:			
	Frame does not jeopardize the life or health of the member ion and would not subject the member to severe pain.			

CLINICAL CRITERIA: Check below all that apply. All criteria must be met for approval. To
support each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be
provided or request may be denied.

DIAGNOSES: Check below one of the diagnoses that applies to qualify.					
	□ Rheumatoid Arthritis (RA)				
		Prescriber is a rheumatologist			
		<u>AND</u>			
		Member has a diagnosis of mode	rate- to-severe rheum	atoid arthritis	
		AND			
		Trial and failure of at least three	(3) months of methotr	rexate therapy	,
		<u>AND</u>			
		Trial and failure of TWO (2) of t		biologics:	
		☐ Humira [®]	□ Enbrel [®]		1 Infliximab
	Gr	anulomatosis with Polyang	iitis OR Microsco	pic Polyan	giitis
	IN	ITIAL THERAPY [Ruxien	ce, Truxima, Ritu	ıxan, Riabı	ni]
		Prescriber is a rheumatologist or	nephrologist		
		AND			
		Member has a diagnosis of mode	rate- to-severe granul	omatosis with	n polyangiitis
		<u>AND</u>			
		Member will receive concurrent	therapy with corticost	eroids	
		AND			
		Member failed cyclophosphamid	e therapy		
		OR Mambanhas a continuidisation to	o avalambaambamida tl	h amana v	
	ш	Member has a contraindication to	o cyclopnospnamide u	nerapy:	

Gr	anulomatosis with Polyangiitis OR Microscopic Polyangiitis
M	AINTENANCE THERAPY [Ruxience, Truxima, Rituxan, Riabni]
	Prescriber is a rheumatologist or nephrologist
	AND
	Induction occurred at least 4 months prior
	<u>AND</u>
	Total duration of treatment does not exceed 24 months
	AND
	Member failed methotrexate or azathioprine therapy
	<u>OR</u>
	Member has a contraindication to methotrexate or azathioprine therapy:
Ne	euromyelitis Optica Spectrum Disorder (NMOSD) [Ruxience, Truxima, Rituxan,
Ri	abni]
	Prescribing physician must be a neurologist
	AND
	Member must be 18 years of age or older
	<u>AND</u>
	Provider must submit medical records (e.g., chart notes) confirming the member has a diagnosis of Neuromyelitis Optica Spectrum Disorder (NMOSD) based on the following:
	☐ Member was found to be seropositive for aquaporin-4 (AQP4) IgG antibodies; AND
	☐ Member has at least one core clinical characteristic (see chart below)
	<u>OR</u>
	☐ Member was found to be seronegative for AQP-4 IgG antibodies OR has unknown AQP-4- IgG status
	AND

Member has at least two core clinical characteristics occurring as a result of one or more clinical attacks (see chart below)
AND
Member experienced ALL of the following:
☐ At least 1 core clinical characteristic must be optic neuritis, acute myelitis with LETM*, or area postrema syndrome
☐ Dissemination in space (≥2 different core clinical characteristics)
☐ Fulfillment of additional MRI requirements, as applicable (see chart below)
AND
Alternative diagnoses have been excluded (e.g., multiple sclerosis, sarcoidosis, cancer, chronic infection, etc.)
AND
Member must have failed at least ONE of the following treatment options:

Core Clinical Characteristics of NMOSD

□ Oral therapy with azathioprine, methotrexate, mitoxantrone or mycophenolate
 □ Member has required plasmapheresis or plasma exchange (PE) or intravenous

- Optic neuritis
- Acute myelitis
- Area postrema syndrome: episode of otherwise unexplained hiccups or nausea and vomiting
- Acute brainstem syndrome
- Symptomatic narcolepsy or acute diencephalic clinical syndrome with NMOSD-typical diencephalic MRI lesions
- Symptomatic cerebral syndrome with NMOSD-typical brain lesions

Immunoglobulin (IVIG).

Core Clinical Characteristics of NMOSD

- Acute optic neuritis: requires brain MRI showing (a) normal findings or only nonspecific white matter lesions, OR (b) optic nerve MRI with T2-hyperintense lesion or T1-weighted gadolinium- enhancing lesion extending over >1/2 optic nerve length or involving optic chiasm
- Acute myelitis: requires associated intramedullary MRI lesion extending over ≥3 contiguous segments (LETM) OR ≥3 contiguous segments of focal spinal cord atrophy in patients with history compatible with acute myelitis
- Area postrema syndrome: requires associated dorsal medulla/area postrema lesions
- Acute brainstem syndrome: requires associated peri-ependymal brainstem lesions

Medication being provided by: Please check applicable box below.	
	Location/site of drug administration:
	NPI or DEA # of administering location:
	<u>OR</u>
	Specialty Pharmacy – Proprium Rx

For urgent reviews: Practitioner should call Sentara Health Plans Pre-Authorization Department if they believe a standard review would subject the member to adverse health consequences. Sentara Health Plan's definition of urgent is a lack of treatment that could seriously jeopardize the life or health of the member or the member's ability to regain maximum function.

**Use of samples to initiate therapy does not meet step edit/preauthorization criteria. **

Previous therapies will be verified through pharmacy paid claims or submitted chart notes.