SENTARA COMMUNITY PLAN (MEDICAID)

PHARMACY PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

<u>Directions</u>: <u>The prescribing physician must sign and clearly print name (preprinted stamps not valid) on this request</u>. All other information may be filled in by office staff; <u>fax to 1-800-750-9692</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 the information provided is not complete, correct, or legible, the authorization process can be delayed.</u>

Drug Requested: Joenja® (leniolisib)

MEMBER & PRESCRIBER INFORMATION: Authorization may be delayed if incomplete.				
Meml	ber Name:			
Member Sentara #:				
Presc	riber Name:			
Prescriber Signature:				
Office	e Contact Name:			
Phone Number:		Fax Number:		
NPI #	:			
DRU	UG INFORMATION: Authorizat	tion may be delayed if incomplete.		
Drug	Name/Form/Strength:			
Dosing Schedule:				
Diagn	osis:	ICD Code, if applicable:		
Weight (if applicable):		Date weight obtained:		
Quai	ntity Limit: 2 tablets per day			
supp		w all that apply. All criteria must be met for approval. To on, including lab results, diagnostics, and/or chart notes, must be		
Init	ial Authorization: 6 months			
	Member is ≥ 12 years of age			
	Member is $\geq 45 \text{ kg}$			
	Provider is an immunologist, has bee activated phosphoinositide 3-kinase (n in consultation with one, or is a specialist in treating patients with PI3K) delta syndrome (APDS)		
	confirmed by submission of laborator	phosphoinositide 3-kinase (PI3K) delta syndrome (APDS) ry results confirming the presence of an activated phosphoinositide ociated genetic PI3Kδ mutation with a documented variant in either		

(Continued on next page)

	sin	ovider must submit documentation of member's medical history which may note recurrent opulmonary infections, sinusitis, pneumonia, bronchitis, chronic Epstein-Barr virus and comegalovirus (CMV) viremia, autoimmune cytopenia, and/or lymphadenopathy/hepatomegaly
		r members with splenomegaly, the provider has submitted measured volume of spleen size using mputed tomography (CT) or magnetic resonance imaging (MRI) scan
		r members with evidence of nodal lesions, the provider has submitted results from a computed nography (CT) or magnetic resonance imaging (MRI) scan
		ember is <u>NOT</u> on concurrent immunosuppressive therapy (e.g., mTOR inhibitors, B-cell depleters, acocorticoids [doses > 25 mg/day of prednisone equivalent], cyclophosphamide, mycophenolate)
		ovider must submit BOTH the following baseline objective clinical laboratory documentation prior to tiating treatment (obtained within the past 30 days):
		Results of a complete blood count with differential to include a T and B cell (lymphocytes) screening
		Measurement of serum immunoglobulin M (IgM) level
e de	nied	ked all documentation (lab results, diagnostics, and/or chart notes) must be provided or request may l. ember is NOT on concurrent immunosuppressive therapy (e.g., mTOR inhibitors, B-cell depleters,
_		acocorticoids [doses > 25 mg/day of prednisone equivalent], cyclophosphamide, mycophenolate)
	by	ember has been observed to have a positive clinical response since the beginning of therapy evidenced disease stability, or mild progression, in any of the following (check all that apply; submitted in cumentation and charted in clinical notes):
		Computed tomography (CT) or magnetic resonance imaging (MRI) scan observing the reduction in lymph node lesions if present at baseline
		Follow-up complete blood count observing normalization in T and B cells
		Computed tomography (CT) or magnetic resonance imaging (MRI) scan observing the reduction in spleen size if applicable
		Follow-up serum immunoglobulin screening observing normalization in IgM levels
		Submission of progress notes documenting frequency or severity of infections
		ember has experienced an absence of unacceptable toxicity from the drug (e.g., severe neutropenia NC $<$ 500 cells/ μ L), skin rash)
1ed	ica	tion being provided by Specialty Pharmacy – Proprium Rx

Not all drugs may be covered under every Plan.

If a drug is non-formulary on a Plan, documentation of medical necessity will be required.

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