SENTARA HEALTH PLANS

PHARMACY/MEDICAL PRIOR AUTHORIZATION/STEP-EDIT REQUEST

Directions: 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; fax to 1-800-750-9692. No additional phone calls will be necessary if all information (including phone and fax #s) on this form is correct. If information provided is not complete, correct, or legible, authorization can be delayed.

Drug Requested: Benlysta® (belimumab) Subcutaneous Injection (Pharmacy)

ME	MBER & PRESCRIBER INFO	DRMATION: Authorization may be delayed if incomplete.	
Memb	ber Sentara #:	Date of Birth:	
Presci	eriber Name:		
Presci	eriber Signature:	Date:	
Office	e Contact Name:		
Phone Number:		Fax Number:	
DEA (OR NPI #:		
DRU	UG INFORMATION: Authorizat	tion may be delayed if incomplete.	
Drug	Form/Strength/Quantity:		
Dosing Schedule:		Length of Therapy:	
Diagn	nosis:	ICD Code:	
Weigh	ht:	Date:	
	ANTITY LIMITS: For adults with ritis: 400 mg once weekly for 4 doses, to	SLE: Maximum of 200 mg once weekly. For adults with lupus then 200 mg once weekly thereafter	
appro	oval. To support each line checked, all	GIS: Check below all that apply. All criteria must be met for documentation, including lab results, diagnostics, and/or chart denied. Check box below for the Diagnosis that applies.	
	Diagnosis - active systemic lupus tandard therapy	s erythematosus (SLE) in adults who are receiving	
<u>In</u>	nitial Authorization: 12 months	S	
	Must be prescribed by or in consulta	tion with a rheumatologist	
<u> </u>	1	vith a diagnosis of active, autoantibody-positive SLE confirmed by	
	☐ anti-nuclear antibody (ANA) tite		
	□ anti-double stranded DNA (anti-	$dsDNA) \ge 30 \text{ IU/mL}$	

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	Member's SLE activity has been confirmed by one of the following (submit results):				
	□ Safety of Estrogen in Lupus Index (SELENA-SLEDAI)	National Assessment – Systemic Lupuscore of 6-12	us Erythematosus Disease Activity		
	□ ≥2 British Isles Lupus Asses	ssment Group (BILAG) B organ doma	in scores		
	Member has tried three of the following and is established on two of the following therapies taken for the last 90 days (please submit chart notes documenting therapy trials with insufficient disease control):				
	□ mycophenolate	□ hydroxychloroquine	□ azathioprine		
	□ cyclophosphamide	□ methotrexate	□ cyclosporine		
	□ corticosteroids	□ Other			
	Member does not have any of the following limitations to therapy: severe active central nervous system lupus; current or previous diagnosis of progressive multifocal leukoencephalopathy (PML); or concurrent use with other biologics				
□ D	iagnosis - active lupus nep	hritis in adults who are receivin	ng standard therapy		
<u>I</u> 1	nitial Authorization: 12 m	onths			
	Must be prescribed by or in con	sultation with a nephrologist or rheum	atologist		
	Member is 18 years of age or older with a diagnosis of active lupus nephritis Class III, IV, or V as confirmed by renal biopsy				
	Member's diagnosis of active, autoantibody-positive SLE was confirmed by one of the following (submit lab results):				
	☐ anti-nuclear antibody (ANA) titer $\geq 1:80$			
	☐ anti-double stranded DNA (anti-dsDNA) ≥ 30 IU/mL			
	Member has active renal disease and has received standard therapy for the last 90 days with corticosteroids along with one of the following (chart notes documenting established therap submitted):				
	mycophenolate				
	cyclophosphamide				
	Provider must obtain a baseline measurement of one of the following collected within the last 30 days (labs must be submitted):				
	urine protein:creatinine rati	o (uPCR)			
	urine protein				
	•	ne following limitations to therapy: sevosis of progressive multifocal leukoen	_		

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Re		thorization Approval: 12 months. Check below all that apply. All criteria must be met for approval.			
		support each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be ovided or request may be denied.			
	Di	Diagnosis - systemic lupus erythematosus (SLE) in adults			
	_				
		All of the initial authorization criteria continues to be met			
		Member's response to therapy has been confirmed by one of the following (submit results):			
		□ Safety of Estrogen in Lupus National Assessment – Systemic Lupus Erythematosus Disease Activity Index (SELENA-SLEDAI) score has improved by and/or maintained at a level that is ≥4 points below baseline score			
		□ No new BILAG-A organ domain score OR 2 new BILAG-B organ domain scores			
		Member has absence of intolerable side effects such as serious infections, signs or symptoms of progressive multifocal leukoencephalopathy (PML), malignancy, severe hypersensitivity reactions/anaphylaxis, or serious infusion reactions			
app	orov	thorization Approval: 12 months. Check below all that apply. All criteria must be met for val. To support each line checked, all documentation, including lab results, diagnostics, and/or chart must be provided or request may be denied.			
	Di	agnosis - active lupus nephritis in adults			
		All of the initial authorization criteria continues to be met			
		Member has had improvement from baseline and/or stabilization since last approval of one of the following (submit current labs completed within the last 30 days):			
		☐ Urine protein:creatinine ratio (uPCR)			
		☐ Urine protein			
		Member has absence of intolerable side effects such as serious infections, signs or symptoms of			

Medication being provided by a Specialty Pharmacy - PropriumRx

progressive multifocal leukoencephalopathy (PML), malignancy, severe hypersensitivity

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.

reactions/anaphylaxis, or serious infusion reactions