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: Tremfya[™] (guselkumab) Injection

MEMBER & PRESCRIBER INF	FORMATION: Authorization may be delayed if incomplete.
Member Name:	
Member Sentara #:	
Prescriber Name:	
Prescriber Signature:	
Office Contact Name:	
Phone Number:	
NPI #:	
DRUG INFORMATION: Authoriz	
Drug Name/Form/Strength:	
	Length of Therapy:
Diagnosis:	ICD Code, if applicable:
Weight (if applicable):	Date weight obtained:
ATTENTION: Tremfya IV induction (lo	pading dose) for treatment of ulcerative colitis can only be billed under

ATTENTION: Tremfya IV induction (loading dose) for treatment of ulcerative colitis can only be billed under the MEDICAL BENEFIT. NDC: 57894-0650-01/02: J1628; 200 mg/20 mL= 200 billable units

Recommended Dosage:

Indication	Dosage	
Moderate-to-Severe Plaque Psoriasis (Adults) – who are candidates for systemic therapy or phototherapy	100mg administered by subcutaneous injection at Week 0, Week 4 and every 8 weeks thereafter	
Active Psoriatic Arthritis (Adults)	100mg administered by subcutaneous injection at Week 0, Week 4 then every 8 weeks	
Moderate-to-Severe active ulcerative colitis (Adults)	200 mg IV induction dose at weeks 0, 4, & 8 by an HCP, recommended maintenance dosage is 100 mg SC at week 16 and every 8 weeks thereafter, or 200 mg SC at week 12 and every 4 weeks thereafter	
Moderate-to-Severe active Crohn's disease (Adults)	200 mg IV or 400mg SC induction dose at weeks 0, 4, & 8, recommended maintenance dosage is 100 mg SC at week 16 and every 8 weeks thereafter, or 200 mg SC at week 12 and every 4 weeks thereafter	

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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.

	□ DIAGNOSIS: Moderate-to-Severe Chronic Plaque Psoriasis				
	Moderate-to-severe plaque psoriasis who are candidates for systemic therapy or phototherapy				
	Member is ≥ 18 years				
	 Diagnosis of moderate to severe plaque psoriasis for ≥ 6 months with ≥ 1 of the following: Affected body surface area (BSA) of ≥ 10% Psoriasis Area and Severity Index (PASI) score ≥ 10 Incapacitation due to plaque location (e.g., head and neck, palms, soles or genitalia) 				
	Member did not respond adequately (or is not a candidate) to a 3-month minimum trial of ≥ 1 systemic agent (e.g., immunosuppressives, and/or methotrexate)				
	Member did not respond adequately (or is not a candidate) to a 3-month minimum trial of phototherapy (e.g., psoralens with UVA light (PUVA) or UVB with coal tar or dithranol)				
	Member is not receiving guselkumab in combination with another biologic agent for psoriasis or non-biologic immunomodulator (e.g., apremilast, tofacitinib, baricitinib)				
	Trial and failure of <u>TWO (2)</u> of the <u>PREFERRED</u> drugs below:				
	☐ Humira [®]	□ Enbrel [®]	□ Infliximab		
D	□ DIAGNOSIS: Active Psoriatic Arthritis (Adults)				
	Member has a diagnosis of active p	osoriatic arthritis			
	Member is ≥ 18 years				
	Trial and failure of methotrexate OR contraindication to methotrexate (e.g., alcohol abuse, cirrhosis, chronic liver disease, or other contraindication)				
	Trial and failure of TWO (2) of the PREFERRED drugs below:				
	☐ Humira [®]	□ Enbrel®	□ Infliximab		

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□]	□ DIAGNOSIS: Moderate-to-Severe Ulcerative Colitis (UC)			
	☐ Member has a diagnosis of moderate to severe ulcerative colitis			
	Men	nber is ≥ 18 years		
	Trial and failure to ONE conventional agent (i.e., 6-mercaptopurine, azathioprine, balsalazide, corticosteroids, cyclosporine, mesalamine, sulfasalazine) used in the treatment of UC after at least a 3-month duration of therapy			
	Member is not receiving guselkumab in combination with another biologic agent for UC or non-biologic immunomodulator (e.g., upadacitinib)			
	Men	nber has tried and failed both:		
		Humira [®]		Infliximab
o I	DIAG	NOSIS: Moderate-to-Severe Crohn	's D	isease (CD)
	Men	nber has a diagnosis of moderate to severe C	rohn	's disease
	Men	nber is ≥ 18 years		
	Trial and failure to of a compliant regimen of oral corticosteroids (unless contraindicated) or intravenous corticosteroids			
	Member is not receiving guselkumab in combination with another biologic agent for CD or non-biologic immunomodulator (e.g., upadacitinib)			
	Member has tried and failed both:			
		Humira [®]		Infliximab
	_			
				oval for duration of 2 months, member
		eive up to three (3) IV infusion dose cation Criteria: To be reviewed for o		time annroyal under the medical
	efit	ation Criteria. To be reviewed for t	iic-	ame approvar under the medicar
	Med	ication will be used as induction therapy		
	Med	ication being provided by:		
		NPI or DEA # of administering location: _		
		aber to receive FDA approved loading dose of 1 hour at Week 0, Week 4, and Week 8	of 20	Omg administered by intravenous infusion over at

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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.