SENTARA HEALTH PLANS

PHARMACY PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

<u>Directions</u>: <u>The prescribing physician must sign and clearly print name (preprinted stamps not valid)</u> on this request. 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: Omvoh[™] SQ & IV (mirikizumab-mrkz)

MEMBER & PRESCRIBER INFORM	MATION: Authorization may be delayed if incomplete.	
Member Name:	, , ,	
Member Sentara #:		
Prescriber Signature:		
Phone Number:		
DRUG INFORMATION: Authorization	may be delayed if incomplete.	
Drug Name/Form/Strength:		
	Length of Therapy:	
Diagnosis:		
Weight (if applicable):		
<u>ATTENTION</u> : Omvoh IV induction (loading can only be billed under the <u>MEDICAL BENE</u>) Adult Dosing Ulcerative Colitis:	g dose) for treatment of Crohn's Disease & Ulcerative Colitis FIT . NDC: 00002-7575-01; J2267	
	voh (J2267) IV 300 mg/15 mL vial; 1 vial = 300 billable units	
	usion over at least 30 minutes at Week 0, Week 4, and Week 8	
☐ Maintenance SubQ:		
 200 mg administered by subcutaneous in Week 12, and every 4 weeks thereafter 	jection (given as two consecutive injections of 100 mg each) at	
o NDC: 00002-8011-27 – Omvoh 100	mg/mL prefilled pen	

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NDC: 00002-8870-27 – Omvoh 100 mg/mL prefilled syringe

- ☐ Induction IV: NDC: 00002-7575-01 Omvoh (J2267) IV 300 mg/15 mL vial; 3 vials = 900 billable units
 - 900 mg administered by intravenous infusion over at least 90 minutes at Week 0, Week 4, and Week 8
- **□** Maintenance SubQ:
 - 300 mg administered by subcutaneous injection (given as two consecutive injections of 100 mg and 200 mg in any order) at Week 12, and every 4 weeks thereafter
 - o NDC: 0002-7717-11 Omvoh 200 mg/2 mL + 100 mg/mL prefilled pen
 - NDC: 0002-7722-11 Omvoh 200 mg/2 mL + 100 mg/mL prefilled syringe

NOTE: The Health Plan considers the use of concomitant therapy with more than one biologic immunomodulator (e.g., Dupixent, Entyvio, Humira, Rinvoq, Stelara) prescribed for the same or different indications to be experimental and investigational. Safety and efficacy of these combinations has **NOT** been established and will **NOT** be permitted.

Sta	ablished and will NOT be permitted.					
•	Will the member be discontinuing a previously prescribed biologic if approved for requested medication?					
		□ Yes OR □ No				
•	If yes, please list the medication that will be discontinued and the medication that will be initiated upon approval along with the corresponding effective date.					
	Medication to be discontinued:	Effective date:				
	Medication to be initiated:	Effective date:				
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: Ulcerative Colitis					
		ered by subcutaneous injection (given as two at Week 12, and every 4 weeks thereafter				
A	uthorization Criteria: To be reviewed for	or approval under the pharmacy benefit				
	☐ Member has a diagnosis of Ulcerative Coliti	S				
	☐ Medication has been prescribed by a Gastro	enterologist				

5-aminosalicylates (balsalazide, olsalazine, sulfasalazine)

☐ Member meets <u>ONE</u> of the following:

months

□ oral mesalamine (Apriso, Asacol/HD, Delzicol, Lialda, Pentasa)

☐ Member has tried and failed budesonide or high dose steroids (40-60 mg prednisone)

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☐ Member has tried and failed at least **ONE** of the following **DMARD** therapies for at least **three** (3)

	Me	mber meets ONE of the following:					
	■ Member tried and failed, has a contraindication, or intolerance to <u>TWO</u> of the following <u>PREFERRED</u> biologics:						
		☐ Preferred adalimumab product	□ Rinvoq [®]	□ Simponi [®]			
		☐ Skyrizi® SC (on-body injector)	□ Stelara®	□ Tremfya [®]			
		□ Xeljanz [®] /XR [®]	□ Zymfentra [™]				
		Member has been established on Omvindicates at least a 90-day supply of chart notes or pharmacy paid claim	Omvoh was dispensed	s <u>AND</u> prescription claims history dwithin the past 130 days (verified by			
		enare notes of pharmacy para claim					
		nosis: Crohn's Disease					
C	□ Maintenance Dose – 300 mg administered by subcutaneous injection (given as two consecutive injections of 100 mg and 200 mg in any order) at Week 12, and every 4 weeks thereafter						
Aut	<u>hori</u>	zation Criteria: To be reviewe	d for approval und	er the pharmacy benefit			
	Mei	mber has a diagnosis of Crohn's Dise	ase				
	Me	dication has been prescribed by a Gas	troenterologist				
	☐ Member meets <u>ONE</u> of the following:						
☐ Member has tried and failed budesonide or high dose steroids (40-60 mg prednisone)							
		Member has tried and failed at least <u>C</u> months	<u>ONE</u> of the following D	MARD therapies for at least three (3)			
		☐ 5-aminosalicylates (balsalazide, o	lsalazine, sulfasalazine)			
		□ oral mesalamine (Apriso, Asacol/	HD, Delzicol, Lialda, P	'entasa)			
☐ Member meets <u>ONE</u> of the following:							
■ Member tried and failed, has a contraindication, or intolerance to <u>TWO</u> of the following <u>PREFERRED</u> biologics:							
		☐ Preferred adalimumab product	□ Cimzia [®]	☐ Skyrizi [®] SC (on-body injector)			
		□ Stelara [®]	□ Rinvoq [®]	□ Zymfentra [™]			

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	☐ Member has been established on Omvoh [™] for at least 90 days <u>AND</u> prescription claims history indicates at least a 90-day supply of Omvoh was dispensed within the past 130 days (verified by).			
	chart notes or pharmacy paid claims)			
☐ Induction Dose (If required) — One time approval for duration of 2 months, member to receive up to three (3) IV infusion doses				
Auth bene	norization Criteria: To be reviewed for one-time approval under the medical fit			
	Medication will be used as induction therapy			
	Medication being provided by:			
	□ Location/site of drug administration:			
	□ NPI or DEA # of administering location:			
	Member to receive FDA approved loading dose of $900~\text{mg}$ administered by intravenous infusion over at least $30~\text{minutes}$ at Week $4,~\text{and}$ Week 8			
Medication 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. *