## 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 may be delayed.</u>

Drug Requested: Skyrizi® SQ & IV (risankizumab) For CD & UC (Pharmacy) (Preferred)

MI	EMBER & PRESCRIBER I	NFORMATION: Authorization may be delayed if incomplete.
Men	nber Name:	
		Date of Birth:
Pres	criber Name:	
		Date:
Offic	ce Contact Name:	
	ne Number:	
NPI	#:	
		norization may be delayed if incomplete.
Drug	g Name/Form/Strength:	
	ng Schedule:	
		ICD Code, if applicable:
Weig	ght (if applicable):	Date weight obtained:
bille		dose for treatment of Crohn's disease & Ulcerative colitis can only be $\underline{\Gamma}$ . NDC: 00074-5015-01; J2327; 600 mg = 600 billable units, 1200 mg =
Adu	ult Dosing:	
		01 – Skyrizi IV 600 mg/10 mL – J2327
	Crohn's disease $-600 \text{ mg}$ adn and 8; $600 \text{ mg} = 600 \text{ billable ur}$	ninistered by IV infusion over a period of at least one hour at week 0, 4 nits per dose
	Ulcerative colitis – 1200 mg ac 8; 1200 mg = 1200 billable unit	dministered by IV infusion a period of at least two hours at week 0, 4 and ts per dose
0	<del>-</del>	-1069-01/00074-1070-01 — Skyrizi SQ 360 mg/ 2.4 mL cartridge; NDC mg/ 1.2 mL cartridge. *Use the lowest effective dosage to maintain
	360 mg administered at week 1	•
	180 mg administered at week 1	2 and every 8 weeks thereafter

(Continued on next page)

(Continued on next page)

indications to be experimental and investigational. S established and will <b>NOT</b> be permitted.	afety and efficacy of these combinations has <b>NOT</b> been
• Will the member be discontinuing a previously p	prescribed biologic if approved for requested medication?
	☐ Yes <b>OR</b> ☐ No
• If yes, please list the medication that will be disc approval along with the corresponding effective	continued and the medication that will be initiated upon date.
Medication to be discontinued:	Effective date:
Medication to be initiated:	Effective date:
	at apply. All criteria must be met for approval. To ding lab results, diagnostics, and/or chart notes, must be
Maintananaa Daga 190 mg ay 260 mg	administered by subartaneous injection of
week 12, and every 8 weeks thereafter	g administered by subcutaneous injection at
<b>Authorization Criteria:</b> To be reviewed for	or approval under the pharmacy benefit
☐ Member has <u>ONE</u> of the following diagnose	S
☐ Moderate-to-severe active Crohn's disea	ise
☐ Moderate-to-severe active Ulcerative col	litis
☐ Prescribed by or in consultation with a <b>Gast</b> ı	roenterologist
☐ Member meets <u>ONE</u> of the following:	
☐ Member has tried and failed budesonide	or high dose steroids (40-60 mg prednisone)
<ul><li>Member has tried and failed at least <u>ONI</u> months</li></ul>	$\underline{C}$ of the following <b>DMARD</b> therapies for at least $\underline{C}$
<ul><li>5-aminosalicylates (balsalazide, olsal</li></ul>	azine, sulfasalazine)
☐ oral mesalamine (Apriso, Asacol/HD	, Delzicol, Lialda, Pentasa)
☐ Induction Dose (If required) – One tine to receive up to three (3) IV infusion d	ne approval for duration of 2 months, member oses
<b>Authorization Criteria:</b> To be reviewed for	or one-time approval under the medical
benefit	
Medication will be used as induction therapy	

(Continued on next page)

**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

## PA Skyrizi for CD & UC (Pharmacy)(CORE) (Continued on next page)

	☐ Medication being provided by:		
	□ Location/site of drug administration:		
	□ NPI or DEA # of administering location:		
	Member to receive FDA approved loading dose for <b>ONE</b> of the following indications:		
	☐ Crohn's disease – 600 mg administered by IV infusion over a period of at least one hour at week 0, 4 and 8		
	☐ Ulcerative colitis – 1200 mg administered by IV infusion a period of at least two hours at week 0, 4 and 8		
Medication being provided by a Specialty Pharmacy – Proprium Rx			

\*\*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. \*