## **AvMed**

## 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 1-305-671-0200</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>

<u>Drug Requested</u>: Entyvio<sup>®</sup> Pen (vedolizumab) (Pharmacy)

MEMBER & PRESCRIBER INF	<b>ORMATION:</b> Authorization may be delayed if incomplete.
Member Name:	
	Date of Birth:
Prescriber Name:	
	Date:
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: Entyvio IV induction (loonly be billed under the MEDICAL BENI	oading dose) for treatment of Crohn's disease & ulcerative colitis can E <u>FIT</u> . NDC: 64764-0300-20; J3380
Quantity Limits: 2 pens per 28 days	
Adult Dosing:  Induction IV: NDC: 64764-0300-20	– Entyvio IV 300 mg vial – J3380

- 300 mg infused intravenously over approximately 30 minutes at Week 0 and Week 2
- ☐ Maintenance SubQ: NDC: 64764-0108-20/21 Entyvio 108 mg/ 0.68 mL prefilled pen
  - Following the first two Entyvio IV doses administered at Week 0 and Week 2 in UC, Entyvio may be switched to subcutaneous (SC) injection at Week 6
  - 108 mg administered by subcutaneous injection starting at Week 6, and then every 2 weeks thereafter
  - Entyvio may be switched from IV infusion to SC injection, for patients in clinical response or remission beyond Week 6. To switch patients to Entyvio SC injection, administer the first SC dose in place of the next scheduled IV infusion and every two weeks thereafter.
  - Discontinue Entyvio in patients who do not show evidence of therapeutic benefit by Week 14

	s to be experimental and investigational. S d and will <b>NOT</b> be permitted.	afety and efficacy of these combinations has <b>NOT</b> been			
• Will th	e member be discontinuing a previously p	rescribed 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.					
Medica	ation to be discontinued:	Effective date:			
Medica	ation to be initiated:	Effective date:			
support ea		at apply. All criteria must be met for approval. To ding lab results, diagnostics, and/or chart notes, must be			
Weel	k 6, and then every 2 weeks therea	ered by subcutaneous injection starting at after or approval under the pharmacy benefit			
□ Me	ember is 18 years of age or older				
□ Me	ember has <b>ONE</b> of the following diagnoses	s:			
	Moderate-to-severe Crohn's disease				
	Moderate-to-severe ulcerative colitis				
□ Pre	escribed by or in consultation with a Gastr	oenterologist			
□ Me	ember meets <b>ONE</b> of the following:				
	Member has tried and failed budesonide	or high dose steroids (40-60 mg prednisone)			
	Member has tried and failed at least <b>ONE</b> months	of the following <b>DMARD</b> therapies for at least <b>three (3)</b>			
	☐ 5-aminosalicylates (balsalazide, olsala	azine, sulfasalazine)			
	□ oral mesalamine (Apriso, Asacol/HD	, Delzicol, Lialda, Pentasa)			

**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

(Continued on next page)

		Member tried and failed, has a contraindication, or intolerance to <u>TWO</u> of the <u>PREFERRED</u> biologics below (verified by chart notes or pharmacy paid claims):					
		☐ Preferred adalimumab product	□ Cimzia <sup>®</sup>	☐ Skyrizi <sup>®</sup> SC (on-body injector)			
		□ Stelara <sup>®</sup>	□ Rinvoq®	□ Zymfentra <sup>™</sup>			
		*NOTE: COMM/FAMIS preferreds = not approved, NDC's starting with 00074 or adalimumab-adbm		a - Humira NDC's starting with 83457 are rred; SG/IP/HIX preferreds = Simlandi			
		Member has been established on Enty- a 90-day supply of Entyvio was disper pharmacy paid claims)	•				
		r ulcerative colitis diagnosis: Member					
	☐ Member tried and failed, has a contraindication, or intolerance to <u>TWO</u> of the <u>PREFERRED</u> biologics below (verified by chart notes or pharmacy paid claims):						
		☐ Preferred adalimumab product	□ Rinvoq®	Skyrizi® SC (on-body injector)			
		□ Simponi <sup>®</sup>	□ Stelara <sup>®</sup>	☐ Tremfya <sup>®</sup>			
		□ Xeljanz <sup>®</sup> /XR <sup>®</sup>	□ Zymfentra <sup>™</sup>				
		*NOTE: COMM/FAMIS preferreds = not approved, NDC's starting with 00074 or adalimumab-adbm		a - Humira NDC's starting with 83457 are rred; SG/IP/HIX preferreds = Simlandi			
		Member has been established on Entyvio <sup>®</sup> for at least 90 days <u>AND</u> claims history indicates <u>at least a 90-day supply of Entyvio was dispensed within the past 130 days</u> (verified by chart notes or pharmacy paid claims)					
		tion Dose (If required) – One ti re up to two (2) IV infusion dose		ration of 1 month, member to			
<u>uth</u>	<u>oriz</u>	zation Criteria: To be reviewed	for one-time appro	val under the medical benefit			
	Me	edication will be used as induction thera	npy				
	Me	Medication being provided by:					
		Location/site of drug administration	ı:				
		NPI or DEA # of administering loca	tion:				
		ember to receive FDA approved loading st 30 minutes at Week 0 and Week 2	g dose of 300 mg admin	istered by intravenous infusion over at			

□ For Crohn's disease diagnosis: Member meets ONE of the following:

(Continued on next page)

	PA Entyvio Pen (AvMed)
(	(Continued from previous page)



\*\*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.\*