

Please indicate:

MEDICARE FORM

1

Entyvio[®] (vedolizumab) Injectable Medication Precertification Request

Page 1 of 3

Start of treatment: Start date

(All fields must be completed and legible for precertification review.)

1

For Ohio MMP:FAX:1-855-734-9389PHONE:1-855-364-0974

For other lines of business: Please use other form.

Note: Entyvio is preferred on MA and MAPD plans.

	ontinuation of therapy: D	Date of last treatment	/ /					
Precertification Reques	ted By:		PI	hone:		Fax:		
A. PATIENT INFORMATIO								
First Name:			Last Name:					
Address:			City:			State:	ZIP:	
Home Phone:		Work Phone:			Cell Phone:		1	
DOB:	Allergies:				Email:			
Current Weight:	-	kgs He	ight:	in	ches or		cms	
B. INSURANCE INFORMA		3	5					
		Does patient hav	e other coverad	e? □Y	′es □No			
			If yes, provide ID#: Carrier Name:					
Insured:		Insured:			_			
C. PRESCRIBER INFORM	ATION							
First Name:		Last Name:			(Check Or	ne): 🗌 M.D.	. 🗌 D.O. 🗌 N.I	P. 🗌 P.A
Address:			City:			State:	ZIP:	
Phone:	Fax:	St Lic #:	NPI #:		DEA #:	•	UPIN:	
Office Contact Name:					Phone:			
D. DISPENSING PROVIDE	ER/ADMINISTRATION IN	FORMATION						
	•	Image: Constraint of the second se		-		er		
City:	State:	ZIP:				Fax:		
Phone:			—			PIN:		
NPI:	FIN							
E. PRODUCT INFORMATI	ION							
Request is for Entyvio (v	/edolizumab): Dose:	Fre	quency:			HCPCS C	ode:	
F. DIAGNOSIS INFORMAT	TION – Please indicate pri	imary ICD Code and specify	any other where	applicable.				
Primary ICD Code: Secondary ICD Code:				Other ICD Code:				
		ormation must be completed						
For Initiation Requests (cl								
	atient had prior therapy wi	s. th Entyvio (vedolizumab) wit concomitantly with aprelima			: DMARDs (e	.g., adalimun	nab, infliximab)?	

Continued on next page



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB
C CUNICAL INFORMATION (confinue	d) Deguired aligical information much	a completed in its entirety for a	I proportification requests
G. CLINICAL INFORMATION (continued	<i>d)</i> – Required clinical information must a	be completed in its <u>entirety</u> for a	i precertification requests.
\longrightarrow Please indicate the seve	diagnosis of fistulizing Crohn's disease erity of the patient's Crohn's disease: □ clinical evidence that the disease is act	Mild Moderate Severe	
└────────────────────────────────────	□ No Is the Crohn's disease manifes	ted by at least one of the follow	ing?
	\longrightarrow Check all that apply: \Box abdom	ninal pain 🔲 arthritis 🔲 bleed	ing 🔲 diarrhea 🔲 internal fistulae
	☐ intestinal obstruction ☐ m	egacolon 🔲 perianal disease	🗌 spondylitis 🔲 weight loss
$ $ \rightarrow \square Yes	atment with corticosteroids ineffective?	cated	
			P hydrocortisone methylprednisolone
Which o	of the following corticosteroids was tried		prednisolone
☐ Yes ☐ No Was tre	atment with 6-mercaptopurine (6-MP) ine		
\rightarrow \Box Yes	No Was treatment with 6-mercapto	purine (6-MP) not tolerated or co	ntraindicated?
	→ ☐ not tolerated ☐ contraind atment with azathioprine ineffective?	cated	
$ \longrightarrow \Box$ Yes	No Was treatment with azathioprin		?
	\longrightarrow not tolerated \Box contraind	cated	
Ulcerative Colitis	d fulminant ulcerative colitis?		
	erity of the patient's ulcerative colitis:	Mild Moderate Severe	
	evidence that the disease is active?		
	atient refractory to immunosuppression No Does the patient require contin methylprednisolone, prednisor	uous immunosuppression with	cortisone, methylprednisolone, prednisone)? corticosteroids (e.g., hydrocortisone,
	\longrightarrow Name and dose: Name:	Dose:	
		Dral 🔲 IV	
	ind dose: Name:Oral IV	Dose:	
	atment with immunosuppressant agent	(e.g. azathioprine m6-mercapt	opurine) ineffective?
	No Was treatment with immunosu or contraindicated?	ppressant agent (e.g., azathiopr	
	→ ☐ not tolerated ☐ contraindi	cated	
Provid	e the name of the drug(s):	of the drug(3).	
🖓 Yes 📮 No 🛛 Was tre	atment with 5-aminosalicylic acid agent No Was treatment with 5-aminosa not tolerated or contraindicated	licylic acid agents (e.g., balsalaz 1?	
	\longrightarrow not tolerated \Box contraind		
	the name of the drug(s):	of the drug(s):	
Please select the sympt	oms the patient exhibit: \Box more than 10) stools per day 🔲 continuous	bleeding 🔲 abdominal pain 🗌 distension
For Continuation requests (clinical docu		e toxic symptoms, including feve	er and anorexia
		st. tofacitinib. or other biologic D	MARDs (e.g., adalimumab, infliximab)?
	est a result of the patient receiving sam		
	ntation supporting disease stability?		
	ntation supporting disease improvement	?	
	Entyvio (vedolizumab) within the past (
	he patient have a documented severe a ng the previous infusion?	nd/or potentially life-threatening	adverse event that occurred during or
	es \Box No Could the adverse reaction	be managed through pre-medic	ation in the home or office setting?
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H. ACKNOWLEDGEMENT					
Request Completed By (Signature Required):					/

Any person who knowingly files a request for authorization of coverage of a medical procedure or service with the intent to injure, defraud or deceive any insurance company by providing materially false information or conceals material information for the purpose of misleading, commits a fraudulent insurance act, which is a crime and subjects such person to criminal and civil penalties.

The plan may request additional information or clarification, if needed, to evaluate requests.