

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.)

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For Medicare Advantage Part B: FAX: 1-844-268-7263 PHONE: 1-866-503-0857

For other lines of business: Please use other form.

Note: Entyvio is preferred on MA and MAPD plans.

Continuation of therapy: Date	e of last treatment	/ /				
Precertification Requested By:		Phone:		Fax:		
A. PATIENT INFORMATION						
First Name:		Last Name:				
Address:		City:		State:	ZIP:	
Home Phone:	Work Phone:		Cell Phone:	•		
DOB: Allergies:			Email:			
Current Weight: Ibs or	kgs He	eight:	inches or		cms	
B. INSURANCE INFORMATION						
Aetna Member ID #:	Does patient hav	ve other coverage? []Yes ∏No			
Group #:		-	 Carrier Name: _			
Insured:	Insured:					
C. PRESCRIBER INFORMATION						
First Name:	Last Name:		(Check Oi	ne): 🗌 M.D. [] D.O. 🗌 N.P. 🗌 P.A	
Address:		City:		State:	ZIP:	
Phone: Fax:	St Lic #:	NPI #:	DEA #:	U	JPIN:	
Office Contact Name:	·		Phone:			
D. DISPENSING PROVIDER/ADMINISTRATION INFO	RMATION					
Place of Administration: Self-administered Physician's Office Outpatient Infusion Center Phone: Center Name:		Name: Address:	Office harmacy	_ Retail Pha ☐ Mail Order		
City: State:	ZIP:	Phone:		Fax:		
Phone: Fax:						
TIN: PIN: NPI:		NPI:				
E. PRODUCT INFORMATION						
Request is for Entyvio (vedolizumab): Dose:	Fre	quency:		HCPCS Cod	le:	
F. DIAGNOSIS INFORMATION – Please indicate prima						
Primary ICD Code: Secondary ICD Code:						
G. CLINICAL INFORMATION – Required clinical inform						
For Initiation Requests (clinical documentation requ						
Note: Entyvio is preferred on MA and MAPD plans. Yes No Has the patient had prior therapy with Yes No Will Entyvio (vedolizumab) be used con	Entyvio (vedolizumab) wit	,	ogic DMARDs (e	.g., adalimumal	b, infliximab)?	

Continued on next page



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB
C CLINICAL INFORMATION (continued)	aquirad alinical information must be comp	lated in its antiraty for all proces	tification requests
G. CLINICAL INFORMATION (continued) – R	equired clinical mormation must be comp	leted in its <u>entirety</u> for all prece	tilication requests.
Please indicate the severity o	nosis of fistulizing Crohn's disease? <i>If yes,</i> f the patient's Crohn's disease:	Moderate 🔲 Severe	diagnosis: / /
	→ Check all that apply: □ abdominal pair □ intestinal obstruction □ megacolo		
			drocortisone 🔲 methylprednisolone
Which of the	following corticosteroids was tried? hyc		olone lain:
└──> ☐ Yes ☐ N ☐ Yes ☐ No Was treatmer └──> ☐ Yes ☐ N	th with 6-mercaptopurine (6-MP) ineffective? o Was treatment with 6-mercaptopurine (6 → □ not tolerated □ contraindicated at with azathioprine ineffective? o Was treatment with azathioprine not tole	S-MP) not tolerated or contraindi	cated?
Ulcerative Colitis	> _ not tolerated _ contraindicated		
Yes No Is there evide	internative control ? the patient's ulcerative colitis: Mild nce that the disease is active? refractory to immunosuppression with cort		e methylprednisolone prednisone)?
$ \qquad \qquad$	o Does the patient require continuous im	munosuppression with corticos	teroids (e.g., hydrocortisone,
Name and do	methylprednisolone, prednisone)? → Name and dose: Name: Please indicate the route: □ Oral □ se: Name:	IV Dose:	
Yes □ No Was treatmer Yes □ N Yes □ N	se: Name: IV te the route: □ Oral □ IV nt with immunosuppressant agent (e.g., az o Was treatment with immunosuppressan or contraindicated? > □ not tolerated □ contraindicated > Provide the name of the d	athioprine, m6-mercaptopurine nt agent (e.g., azathioprine, m6) ineffective? -mercaptopurine) not tolerated
☐ Yes ☐ No Was treatmer	name of the drug(s):	id agents (e.g., balsalazide, me	
	ame of the drug(s): ne patient exhibit: □ more than 10 stools □ acute, severe toxic s	per day 🔲 continuous bleedir ymptoms, including fever and a	
For Continuation requests (clinical document			
	used concomitantly with aprelimast, tofac result of the patient receiving samples of E n supporting disease stability?	-	s (e.g., adalimumab, infliximab)?
□ Yes □ No Is there clinical documentation □ Yes □ No Has the patient received Enty	n supporting disease improvement? vio (vedolizumab) within the past 6 months		
following the	ient have a documented severe and/or po previous infusion? No Could the adverse reaction be man		-



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H. ACKNOWLEDGEMENT					
Request Completed By (Signature Required):			Date:	/	/

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.