

Drier Authorization Deguest Form				
Prior Authorization Request Form Sendero Fax: 512-901-9724		Entyvio Phone: 855-297-9191		
		<u> </u>		
URGENCY: STANDARD URGENT (In checking this box, I attest to the fact that applying the standard review time frame may seriously jeopardize the customer's life, health or ability to regain maximum function)				
Provider Information		Patient Information		
Referring/Prescribing Physician:  PCP Specialist Name: Please identify SPECIALTY: DEA, NPI or TIN: Contact: Phone: ( ) Fax: ( )		Patient's Name: Birth Date: ID Number: Phone Number: Patient Height: Patient Weight:		
Indicate where the drug is being DISPENSED		Indicate where the drug is being ADMINISTERED		
☐ Ambulatory Surgery Center		☐ Ambulatory Surgery Center		
☐ Hom	e Care Agency	☐ Inpatient Hospital		
☐ Inpa	tient Hospital	☐ Long Term Care		
☐ Long	g Term Care	☐ Outpatient Hospital		
☐ Outpatient Hospital		□ Patient's Home		
☐ Patient's Home		☐ Pharmacy		
☐ Pharmacy		☐ Physician's Office		
_	sician's Office	☐ Other (explain):		
☐ Othe	er (explain):	Anticipated Date of Service:		
Approvals may be subject to dosing limits in accordance with FDA-approved labeling, accepted compendia, and/or evidence-based practice guidelines.				
		CAL INFORMATION		
	FATILITI CLINI	CAL IN ORMATION		
CRITERIA QUESTIONS:  1. Has the patient been diagnosed with any of the following?  □ Crohn's disease (CD)  □ Ulcerative colitis (UC)  □ Other:				
2.	What is the HCPCS code?What is the NDC#:	What is the ICD-10 code?		
3. Will the requested drug be used in combination with any other biologic or targeted synthetic DMARD (e.g., Olumiant, Xeljanz)? □ Yes □ No				
4.	4. Has the patient had a TB screening test (e.g., a tuberculosis skin test [PPD] or an interferon-release assay [IGRA]) within 6 months of initiating therapy? □ Yes □ No			
5.	5. What were the results of the TB screening test? □ Positive □ Negative			
6.	6. Does the patient have latent or active tuberculosis (TB)? □ Latent □ Active □ No/Neither			
7.	<ul> <li>7. If the patient has latent or active tuberculosis, has treatment been initiated or completed?</li> <li>□ Yes - treatment initiated</li> <li>□ Yes - treatment completed</li> <li>□ No</li> </ul>			
8.	Is this request for continuation of therapy?	Yes □ No If No, skip to diagnosis section.		
9.	<ol> <li>For continuation of therapy requests, has the patient achieved or maintained positive clinical response as evidenced by low disease activity or improvement in signs and symptoms since starting treatment with the requested drug? ☐ Yes ☐ No</li> </ol>			

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	ls the patient currently receiving the requested drug through samples or a manufacturer's patient assistance program? □ Yes □ No
	OSIS SECTION Please only complete sections below that are relevant to the patient's diagnosis.  A. Crohn's Disease
11.	There is documentation of moderate to severe Crohn's disease per the Crohn's Disease Activity index (CDAI):  Mild = CDAI <220  Moderate = CDAI 220-450  Severe = CDAI >450
	There is documentation of 1 or more of the following high-risk features:  Diagnosis at age <30 years  Ileal disease  Penetrating or stricturing disease  Perianal or severe rectal disease  Extra-intestinal manifestations  History of bowel resections  Initial extensive bowel involvement on endoscopy  None
13.	The treatment is prescribed by or in consultation with a gastroenterologist ☐ Yes ☐ No
	Has the patient previously received a biologic indicated for Crohn's disease? ☐ Yes ☐ No If Yes, please indicate the drug, duration, response, and intolerance/contraindication if applicable:
inhibitor these a	note, the preferred biologic class is a TNF inhibitor (specifically Cimzia), followed by Stelara after failure of a TNF. Please consider prescribing 1 of these drugs before Entyvio if clinically appropriate. If Entyvio is preferred over gents, please provide additional clinical reasoning documentation here:  Does the patient have perianal or fistulizing Crohn's disease?  Possible No
	Has the patient had an inadequate response to a minimum 3 month trial at the maximally indicated dose of 1 or more of the following within the last 6 months?  If Yes, indicate below and no further questions.  Sulfasalazine  Mesalamine (if primarily colonic disease)  Azathioprine at minimum dose 1.5 mg/kg daily  6-mercaptopurine at minimum dose 50mg daily  Methotrexate at minimum dose 15mg IM or SQ weekly  Systemic corticosteroids (prednisone, methylprednisolone)  None of the above therapies have been trialed
	Does the patient have a contraindication or intolerance to at least 2 options listed above? ☐ Yes ☐ No If yes, please document medications and respective contraindications/intolerances:
18.	E: Ulcerative Colitis  There is a diagnosis of moderate to severe ulcerative as evidenced by one of the following:  >4 loose and/or bloody bowel movements per day  Evidence of systemic toxicity (e.g., fever, tachycardia, anemia with Hgb<10.0 g/dL, weight loss, and/or elevated CRP or ESR).  Endoscopic findings of marked erythema, absent vascular pattern, friability, erosions, spontaneous bleeding, and/or ulceration (e.g., findings consistent with a Mayo endoscopic sub score of at least 2, or Ulcerative Colitis Endoscopic Index of Severity of at least 5)  Patients with corticosteroid dependent or corticosteroid refractory disease
	Patients at high risk for colectomy with clinical documentation of risk by prescribing provider

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19. The treatment is prescribed by or in consultation with a gastroenterologist □ Yes □ No				
20. Has the patient previously received a biologic or targeted synthetic disease modifying drug (e.g., Xeljanz) indicated for moderately to severely active ulcerative colitis? ☐ Yes ☐ No If Yes, please indicate the drug, duration, response, and intolerance/contraindication if applicable:				
* Please note, the preferred biologic class is a TNF inhibitor (specifically Simponi). Please consider prescribing a TNF inhibitor before Entyvio if clinically appropriate. If Entyvio is preferred, please provide additional clinical reasoning documentation here:				
21. Has the patient been hospitalized for acute severe ulcerative colitis (e.g., continuous bleeding, severe toxic symptoms, including fever and anorexia)? ☐ Yes ☐ No				
<ul> <li>22. Has the patient had an inadequate response to a minimum 2 month trial at the maximally indicated dose of 1 or more of the following within the last 6 months?  If Yes, indicate below and no further questions.  Oral 5-aminosalycylic acid (e.g., sulfasalazine, mesalamine) at a minimum dose of 2g daily  Rectal 5-aminosalycylic acid enemas with minimum dose of 1g daily  Note: failure of rectal 5-ASA suppositories alone will not meet criteria for biologic  Budesonide  Thiopurines (e.g., azathioprine, 6-mercaptopurine)  Methotrexate with a minimum dose of 15mg IM or SQ weekly  Systemic corticosteroids (e.g., prednisone, methylprednisolone)  None of the above therapies have been trialed</li> <li>23. Does the patient have a contraindication or intolerance to at least 2 options listed above?   Yes  No If yes, please document medications and respective contraindications/intolerances:</li> </ul>				
I attest that this information is accurate and true, and that documentation supporting this information is available for review if requested by Sendero Health Plans.				
Prescriber or Authorized Signature	DATE			