

### **MEDICARE FORM**

# Tysabri® (natalizumab) **Medication Precertification Request**

Page 1 of 3

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

Start of treatment: Start date / / Please indicate:

For Michigan MMP: **FAX:** 1-844-241-2495 **PHONE:** 1-855-676-5772

For other lines of business:

Please use other form.

Note: For the treatment of Crohn's disease, Tysabri is non-preferred. Entyvio, Inflectra, and Remicade are preferred for MA plans and Humira and Skyrizi are preferred for MAPD

Continuation of therapy: Date of last treatment/						sclerosis, Tysabri is preferred.	
Precertification Requested	d By:		Р	hone:	Fax: _		
A. PATIENT INFORMATION							
First Name:			Last Name:				
Address:			City:		State:	ZIP:	
Home Phone:	W	/ork Phone:		Cell Pho	one:		
DOB:	Allergies:			E-mail:			
Current Weight:		Height: _	incl	nes or	cms		
B. INSURANCE INFORMATION	ON						
Member ID #:		Does patient have other coverage?			No		
Group #:			If yes, provide ID#: Carrier Name:				
Insured:		Insured:	Insured:				
C. PRESCRIBER INFORMAT	TION						
First Name:		Last Name:	1	(Chec	k One): M.D.	☐ D.O. ☐ N.P. ☐ P.A.	
Address:			City:		State:	ZIP:	
Phone:	Fax:	St Lic #:	NPI #:	DEA #	#:	UPIN:	
Provider Email:		Office Contact Name:		Phone	e:		
D. DISPENSING PROVIDER/ Place of Administration:	ADMINISTRATION INFOR	RMATION		sing Provider/Phar			
Self-administered ☐ Physician's Office   Outpatient Infusion Center Phone:   Center Name: ☐   Home Infusion Center Phone:   Agency Name: ☐   Administration code(s) (CPT):			Specialty Phar Name: Address:		ce Retail Pharmacy nacy Other:  State: ZIP:		
Address:		710.					
City: Phone:							
TIN:			''''				
NPI:			_   NPI:				
E. PRODUCT INFORMATION	N						
Request is for Tysabri: Do			-		HCPCS Cod	de:	
F. DIAGNOSIS INFORMATIO							
Primary ICD Code:							
G. CLINICAL INFORMATION			n its <u>entirety</u> for	all precertification re	equests.		
For All Requests (clinical do Note: For the treatment of			tuvio Infloctr	a and Pomicado	are professed for	MA plane and Humira	
and Skyrizi are preferred f					are preferred for	WA plans and numina	
	itient had prior therapy with			-			
☐ Yes ☐ No Has the patient had a trial and failure, intolerance, or contraindication to any of the following? (select all that apply) ☐ Entyvio (vedolizumab) ☐ Inflectra (infliximab-dyyb) ☐ Remicade (infliximab)							
☐ Yes ☐ No Has the pa	itient had a trial and failure, (adalimumab)	intolerance, or contraind			ct all that apply)		
Please explain if there are any diagnosis (select all that apply	y other medical reason(s) th	nat the patient cannot use	•		ucts when indicated	d for the patient's	
□ Entyvio	(vedolizumab)	а (піпіхіпіар-фур) 🔲 Ке	emicade (inilixin	iau)			

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## Tysabri® (natalizumab) Medication Precertification Request

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

FAX: 1-844-241-2495 PHONE: 1-855-676-5772

For other lines of business:

Please use other form.

Note: For the treatment of Crohn's disease, Tysabri is non-preferred. Entyvio, Inflectra, and Remicade are preferred for MA plans and Humira and Skyrizi are preferred for MAPD plans. For the treatment of multiple sclerosis, Tysabri is preferred.

Patient First Name	Patient Last Name	Patient Phone	Patient DOB			
G. CLINICAL INFORMATION (continued	Required clinical information must	be completed in its entirety for all p	recertification requests.			
Please explain if there are any other medic diagnosis (select all that apply).			*			
Please indicate the result   ☐ Yes ☐ No Will the patient have doc ☐ Yes ☐ No Is this infusion request in	e of the anti-JCV antibody test:/ ults of the anti-JCV antibody test with E umented anti-JCV antibody testing wit an outpatient hospital setting? patient medically unstable for infusions history of any cardiopulmonary conditions	/ ELISA: ☐ positive ☐ negative h ELISA annually after initiating trea at alternate levels of care?				
	ocumentation of unstable vascular according that the patient has an inability to safe nability to tolerate intravenous volume a document the following:	ess? ely tolerate intravenous volume load load due to unstable renal function?	? ed: / /			
Please select: ☐ Less	diagnosis of fistulizing Crohn's disease ng the patient has been diagnosed with than 1 month ☐ 1 month ☐ 2 mon	e? n fistulizing Crohn's disease:				
Yes						
megacolon   perianal disease   spondylitis   weight loss   None of the above						
☐ Yes ☐ No Will Tysabri (natalizumat			ors) (e.g., adalimumab, infliximab)?			
Multiple Sclerosis  Which of the following types of MS has the  ☐ Relapsing-Remitting MS (RRMS) ☐ F ☐ Yes ☐ No Has the patient discontin How many of the following preferred altern Aubagio (teriflunomide), Avonex (interferor (alemtuzumab), Plegridy (peginterferon bei ☐ 0 ☐ 1 ☐ 2 ☐ 3 ☐ 4 or more	rimary-Progressive MS (PPMS)	g MS (not including Ampyra (dalfan e trial been ineffective, not tolerated 1b), Gilenya (fingolimod), Glatopa/C	npridine))? d or is contraindicated?			

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### Tysabri<sup>®</sup> (natalizumab) Medication Precertification Request

Page 3 of 3

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For Michigan MMP: FAX: 1-844-241-2495 PHONE: 1-855-676-5772

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Note: For the treatment of Crohn's disease, Tysabri is non-preferred. Entyvio, Inflectra, and Remicade are preferred for MA plans and Humira and Skyrizi are preferred for MAPD plans. For the treatment of multiple sclerosis, Tysabri is preferred.

Patient First Name	Patient Last Name	Patient Phone	Patient DOB					
G CLINICAL INFORMATION (continued) - Re	equired clinical information must be comple	ted in its entirety for all preceptific	ration requests					
G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its entirety for all precertification requests.  For Continuation Requests (clinical documentation required for all requests):								
Please indicate the length of time on Tysabri (natalizumab):								
Yes No Is this continuation request a result of the patient receiving samples of Tysabri (natalizumab)?								
Yes No Has the patient had a documented anti-JCV antibody test with ELISA within the last 12 months?  Please indicate the date of the last anti-JCV antibody test with ELISA://								
Please indicate the results of the anti-JCV antibody test with ELISA: positive negative								
Yes No Has the patient received Tysabri (natalizumab) within the past 6 months?								
Yes No Does the patient have a documented severe and/or potentially life-threatening adverse event that occurred during or following the previous infusion?								
Yes No Could the adverse reaction be managed through pre-medication in the office setting?  Yes No Is there clinical documentation supporting disease stability?								
	•							
Yes No Is there clinical documentation supporting disease improvement?								
For Crohn's Disease: Please indicate the severity of the disease at bas	seline (pretreatment with Tysahri (patalizum	ash)): $\square$ mild $\square$ moderate $\square$	savara					
·	, ,	iab)). [ Illing [ Illogerate [	Severe					
For Crohn's Disease or Fistulizing Crohn's Disease:  Yes No Will Tysabri (natalizumab) be used concomitantly with immunosuppressants or TNF inhibitors (e.g., adalimumab, infliximab)?								
For Multiple Sclerosis:								
Which of the following types of MS has the patier		D						
☐ Relapsing-Remitting MS (RRMS) ☐ Primary-Progressive MS (PPMS) ☐ Progressive-Relapsing MS (PRMS) ☐ Secondary-Progressive MS (SPMS) ☐ Yes ☐ No Has the patient discontinued other medications used for treating MS (not including Ampyra (dalfampridine))?								
Yes No Has the patient discontinued of	ther medications used for treating MS (not	including Ampyra (dalfampridine)	)?					
H. ACKNOWLEDGEMENT								
Request Completed By (Signature Require	d):		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.