CardinalCare

COMMONWEALTH OF VIRGINIA DEPARTMENT OF MEDICAL ASSISTANCE SERVICES

Service Authorization (SA) Form

Tysabri ® (natalizumab)

If the following information is not complete, correct, or legible, the SA process can be delayed.

Please use one form per member.

Physician Administered Drug: This form is only to be used for members obtaining the medication from a pharmacy through billing the pharmacy benefit at point-of-sale. Please reference <u>Virginia Medicaid Tysabri Clinical Criteria</u> for members/providers that will obtain the medication through the medical benefit.

WEINBER INFORMATION				
Last Name:	First Name:			
Medicaid ID Number:	Date of Birth:			
	Weight in Kilograms:			
PRESCRIBER INFORMATION				
Last Name:	First Name:			
NPI Number:				
Phone Number:	Fax Number:			
DRUG INFORMATION				
Drug Name/Form:				
Strength:				
Dosing Frequency:				
Length of Therapy:				
Quantity per Day:				
(Form continued on next page.)				

Virginia DMAS SA Form: Tysabri ® (natalizumab) Member's Last Name: Member's First Name: DIAGNOSIS AND MEDICAL INFORMATION For an initial request for Multiple Sclerosis, complete the following to receive a 6-month approval: 1. Is the member at least 18 years of age? AND Yes No 2. Has the member prescriber and member enrolled in and meet the conditions of the TOUCH (applicable to Tysabri) or REMS (applicable to Tyruko) programs? AND Yes No 3. Has the member had a JCV antibody test completed within the past 6 months and been counseled on the risks and benefits of treatment? AND Yes No 4. Will the requested product not be used in combination with antineoplastic, immunosuppressant, or immunomodulating agents? AND | Yes No 5. Is the member immunocompetent? AND | Yes l No 6. Will Tysabri be used as a single therapy? AND Yes 7. Does the member have a confirmed diagnosis of multiple sclerosis (MS) as documented by laboratory report (i.e., MRI); AND | Yes No

(Form continued on next page.)

No

Yes

8. Does the member have a diagnosis of a relapsing form of MS [i.e., relapsing-remitting MS (RRMS)*, active

secondary progressive disease (SPMS)**, or clinically isolated syndrome (CIS)***]? OR

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Member's Last Name:

Member's First Name:

Fo	r a renewal request for Multiple Sclerosis, complete the following to receive a 12-month approval:						
Μι	Multiple Sclerosis Renewal Request:						
1.	Does the member continue to meet the relevant criteria identified in the initial criteria? AND						
	Yes No						
2.	Does the member have an absence of unacceptable toxicity from the drug? AND						
	Yes No						
3.	Is the member being continuously monitored for response to therapy indicates a beneficial response?						
	Yes No						
Fo	r an initial request for Crohn's Disease, complete the following questions to receive a 6-month approval:						
1.	Is the member at least 18 years of age? AND						
	☐ Yes ☐ No						
2.	Has the member prescriber and member enrolled in and meet the conditions of the TOUCH (applicable to Tysabri) or REMS (applicable to Tyruko) programs? AND						
	☐ Yes ☐ No						
3.	Does the member have a documented negative JCV antibody ELISA test within the past 6 months? AND						
	☐ Yes ☐ No						
4.	Will the requested product not be used in combination with antineoplastic, immunosuppressant, or immunomodulating agents? AND						
	☐ Yes ☐ No						
5.	Is the member immunocompetent? AND						
	☐ Yes ☐ No						
6.	Does the member have moderate to severe active disease? AND						
	☐ Yes ☐ No						
7.	Has the physician has assessed baseline disease severity utilizing an objective measure and tool? AND						
	☐ Yes ☐ No						
8.	Does the member have a documented trial and failure on ONE oral immunosuppressive therapy for at least 3 months, unless use is contraindicated, such as corticosteroids, methotrexate, azathioprine, and 6-mercaptopurine? AND						
	☐ Yes ☐ No						
	(Form continued on next page.)						

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		Virginia Divias sa Forni. Tysabri - (natalizumab)	
Me	emb	er's Last Name: Member's First Name:	
9.		es the member have a documented trial and failure two of the preferred Cytokine and CAM antagonistents for Crohn's Disease (see Cytokine and CAM Antagonists on the PDL)? AND	-
		Yes No	
10.	imi	Il Tysabri be used as single agent therapy [Not used concurrently with another biologic drug or munosuppressant (e.g., 6-mercaptopurine, azathioprine, cyclosporine, methotrexate, etc.) used for other biologic drug or bin's Disease?	
		Yes No	
Foi	ar	enewal request for Crohn's Disease, complete the following questions:	
1.	Init	cial renewal only (6-month approval):	
	a.	Has the member been tapered off of oral corticosteroids within 6 months of starting Tysabri? AND	
		☐ Yes ☐ No	
	b.	Has the disease responded as indicated by improvement in signs and symptoms compared to baseline such as endoscopic activity, number of liquid stools, presence and severity of abdominal pain, presence of abdominal mass, body weight compared to IBW, hematocrit, presence of extra intestinal complications, tapering or discontinuation of corticosteroid therapy, use of anti-diarrheal drugs, and/or an improvement on a disease activity scoring tool?	
		☐ Yes ☐ No	
2.	Sul	osequent renewals (12-month approval):	
	a.	Does the member not require additional steroid use that exceeds 3 months in a calendar year to control their Crohn's disease? AND	
		☐ Yes ☐ No	
	b.	Has the disease responded as indicated by improvement in signs and symptoms compared to baseline such as endoscopic activity, number of liquid stools, presence and severity of abdominal pain, presence of abdominal mass, body weight compared to IBW, hematocrit, presence of extra intestinal complications, tapering or discontinuation of corticosteroid therapy, use of anti-diarrheal drugs,	

(Form continued on next page.)

No

Yes

and/or an improvement on a disease activity scoring tool?

*Definitive diagnosis of MS with a relapsing-remitting course is based upon BOTH dissemination in time and space. Unless contraindicated, MRI should be obtained (even if criteria are met).					
Dissemination In Time	Dissemination In Space				
(development/appearance of new CNS lesions over time)	(development of lesions in distinct anatomical locations				

- ≥ 2 clinical attacks; OR
- 1 clinical attack and one of the following:
 - MRI indicating simultaneous presence of gadolinium-enhancing and non-enhancing lesions at any time or by a new T2- hyperintense or gadolinium-enhancing lesion on follow-up MRI compared to baseline scan CSF-specific oligoclonal bands
- (development of lesions in distinct anatomical locations within the CNS; multifocal)
- 2 lesions; OR
- 1 lesion **and** one of the following:
 - Clear-cut historical evidence of a previous attack involving a lesion in a distinct anatomical location; OR
 - MRI indicating ≥ 1 T2-hyperintense lesions characteristic of MS in ≥ 2 of 4 areas of the CNS (periventricular, r juxtacortical, infratentorial, or spinal cord)

**Active secondary progressive MS (SPMS) is defined as the following:

- Expanded Disability Status Scale (EDSS) score ≥ 3.0; AND
- Disease is progressive ≥ 3 months following an initial relapsing-remitting course (i.e., EDSS score increase by 1.0 in members with EDSS \leq 5.5 or increase by 0.5 in members with EDSS \geq 6); AND
 - ≥ 1 relapse within the previous 2 years; OR
 - Member has gadolinium-enhancing activity or new or unequivocally enlarging T2 contrast-enhancing lesions as evidenced by MRI

***Definitive diagnosis of CIS is based upon ALL of the following:

- A monophasic clinical episode with member-reported symptoms and objective findings reflecting a focal or multifocal inflammatory demyelinating event in the CNS
- Neurologic symptom duration of at least 24 hours, with or without recovery
- Absence of fever or infection; AND
- Member is not known to have multiple sclerosis

(Form continued on next page.)

Member's Last Name:

Member's First Name:

***Definitive diagnosis of CIS is based upon ALL of the following:

- 1 year of disability progression independent of clinical relapse; AND
- Two of the following:
 - ≥ 1 T2-hyperintense lesion characteristic of MS in one or more of the following regions of the CNS:
 periventricular, cortical or juxtacortical, or infratentorial; AND
 - ≥ 2 T2-hyperintense lesions in the spinal cord
- Presence of CSF-specific oligoclonal bands

Proceribor Signature (Poquired)	Date

Prescriber Signature (Required)

By signature, the physician confirms the above information is accurate and verifiable by member records.

Please include ALL requested information; Incomplete forms will delay the SA process.

Submission of documentation does NOT guarantee coverage by the Department of Medical Assistance Services.

The completed form may be: **FAXED TO 800-932-6651**, phoned to 800-932-6648, or mailed to:

Prime Therapeutics State Government Solutions

ATTN: MAP

11013 W. Broad Street Glen Allen, VA 23060