BLA 125104

TYSABRI® (natalizumab) Intravenous Injection
Monoclonal Antibody

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RISK EVALUATION AND MITIGATION STRATEGY (REMS)

TYSABRI Outreach: Unified Commitment to Health (TOUCH®) Prescribing Program (MS & CD)
I. GOALS:

The goals of the Tysabri REMS are:

1. To inform prescribers, infusion center healthcare providers, and patients about the risk of progressive multifocal leukoencephalopathy (PML) associated with TYSABRI including the increased risk of PML with longer treatment duration, prior immunosuppressant use and the presence of anti-JCV antibodies.

2. To warn against concurrent use with antineoplastic, immunosuppressant, or immunomodulating agents, and in patients who are immunocompromised.

3. To promote early diagnosis of PML and timely discontinuation of TYSABRI in the event of suspected PML.

II. REMS ELEMENTS:

A. Medication Guide

A Medication Guide (MG) for TYSABRI will be dispensed to each patient prior to each TYSABRI infusion in accordance with 21 CFR 208.24.

Please see the appended Medication Guide.

B. Elements to Assure Safe Use

1. Healthcare providers who prescribe TYSABRI are specially certified.

   a. Biogen Idec will ensure that health care providers who prescribe TYSABRI are specially certified.

   b. To become certified, prescribers will be required to enroll in the TOUCH® Prescribing Program by completing the following requirements:

      i. Review the TYSABRI REMS prescriber educational materials, including the full Prescribing Information.

      ii. Complete and sign the Prescriber/Patient Enrollment Form and acknowledge the following:

         a) I have read and understand the full Prescribing Information for TYSABRI

         b) I understand that TYSABRI increases the risk of progressive multifocal leukoencephalopathy (PML), an opportunistic viral infection of the brain that usually leads to death or severe disability. When initiating and continuing treatment with
TYSABRI, I should consider whether the expected benefit of TYSABRI is sufficient to offset this risk.

c) I am aware that cases of PML have been reported in patients taking TYSABRI who were recently or concomitantly treated with immunomodulators or immunosuppressants, as well as in patients receiving TYSABRI monotherapy.

d) I understand that three risk factors identified thus far that increase the risk of PML in TYSABRI-treated patients are:

- Longer treatment duration, especially beyond 2 years
- Prior treatment with an immunosuppressant (e.g., mitoxantrone, azathioprine, methotrexate, cyclophosphamide, mycophenolate mofetil)
- The presence of anti-JCV antibodies.

These factors should be considered in the context of expected benefit when initiating and continuing treatment with TYSABRI.

e) To my knowledge, this patient has no known contraindications to TYSABRI, including PML.

f) I have instructed this patient to promptly report to me any new or worsening symptoms that persist over several days, especially nervous system symptoms.

g) I understand that this patient should be seen and evaluated 3 months after the first infusion, 6 months after the first infusion, every 6 months thereafter for as long as this patient receives TYSABRI, and for at least 6 months after TYSABRI has been discontinued.

h) I will determine every 6 months whether this patient should continue on TYSABRI and, if so, authorize treatment for another 6 months.

i) I understand that I am required to submit an “Initial Discontinuation Questionnaire” when TYSABRI is discontinued and a “6-Month Discontinuation Questionnaire” following discontinuation of TYSABRI.

j) I should report to Biogen Idec, as soon as possible, cases of PML, hospitalizations due to opportunistic infection, or deaths.
k) I understand that data concerning this patient and me will be entered into the mandatory TOUCH Prescribing Program. Biogen Idec requires my cooperation with periodic data collection. Failure to provide the requested information or otherwise comply with the requirements of the TOUCH Prescribing Program may result in discontinuation of TYSABRI treatment for this patient and termination of my authorization to prescribe TYSABRI.

l) I have received educational materials regarding the benefits and risks of TYSABRI treatment.

m) I have, or another healthcare provider under my direction has, educated this patient on the benefits and risks of treatment with TYSABRI, provided him or her with the Patient Medication Guide and Enrollment Form, instructed him or her to read these materials, and encouraged him or her to ask questions when considering TYSABRI.

Acknowledgments specific to Multiple Sclerosis (MS)

I understand that TYSABRI is indicated as monotherapy for relapsing forms of MS.

n) I understand that this patient has a relapsing form of MS based on clinical and radiological evidence.

o) I understand that TYSABRI increases the risk of progressive multifocal leukoencephalopathy (PML), an opportunistic viral infection of the brain that usually leads to death or severe disability. When initiating and continuing treatment with TYSABRI, I should consider whether the expected benefit of TYSABRI is sufficient to offset this risk.

p) I understand that TYSABRI is not ordinarily recommended for patients who are receiving chronic immunosuppressant or immunomodulatory therapy, or who are significantly immunocompromised from any other cause.

q) I understand that an MRI should be performed prior to initiating therapy with TYSABRI in MS patients.

Acknowledgments specific to Crohn’s Disease (CD)

r) I understand that TYSABRI is indicated for adult patients with moderately to severely active CD with evidence of inflammation who have had an inadequate response to, or are...
unable to tolerate, conventional CD therapies and inhibitors of TNF-α

s) I understand that patients receiving TYSABRI should not take concomitant immunosuppressants (e.g., 6-mercaptopurine, azathioprine, cyclosporine, or methotrexate) or inhibitors of TNF-α

t) I understand that this patient has moderately to severely active CD with evidence of inflammation

u) I have discussed other Crohn’s disease treatments with this patient

v) I understand that TYSABRI should be discontinued if a patient has not experienced a therapeutic benefit by 12 weeks of therapy

w) I understand that patients receiving steroid therapy at the time of TYSABRI initiation must undergo a steroid tapering regimen once a therapeutic response is achieved. If the patient with Crohn’s disease cannot be tapered off of steroids within six months of starting TYSABRI, TYSABRI should be discontinued

c. Biogen Idec will:

  i. Ensure that prescriber enrollment can successfully be completed

  ii. Ensure that prescribers receive the TYSABRI prescribing information and the following materials that are part of the TYSABRI REMS and are appended:

  - TOUCH Prescribing Program Educational Slide Set
  - TOUCH Prescribing Program Enrollment Kits (specific to MS or CD)
    1. TOUCH Prescribing Program Overview
    2. Medication Guide
    3. Prescriber/Patient Enrollment Form (MS or CD)
    4. Pre-Infusion Patient Checklist (combined MS and CD)
    5. Guidance for Evaluation of New Neurologic Symptoms in Patients Receiving TYSABRI (MS)
    6. Understanding PML for Gastroenterologists (CD)
  - TOUCH On-Line (www.TOUCHPROGRAM.com)
  - Change Prescriber Authorization Form
  - 12 Week Questionnaire for Crohn’s Disease
iii. Ensure that enrollment forms are complete before activating a prescriber’s enrollment in the TOUCH Prescribing Program.

iv. Ensure that prescribers are notified when they are successfully enrolled in the TOUCH Prescribing Program, and therefore, are certified to prescribe TYSABRI.

2. **TYSABRI will be dispensed only by pharmacies and infusion sites that are specially certified.**

   a. Biogen Idec will ensure that certified pharmacies that dispense TYSABRI are specially certified.

   b. Pharmacies that dispense TYSABRI to infusion sites must enroll in the Tysabri TOUCH Prescribing Program by submitting a completed enrollment form and designating a person with appropriate authority to acknowledge the following:

      i. The pharmacy has received training and educational materials on the TOUCH Prescribing Program

      ii. I understand that certified pharmacies may dispense TYSABRI only to authorized infusion sites

      iii. I understand that, per the requirements of the TOUCH Prescribing Program, this certified pharmacy’s compliance may be reviewed by the Food and Drug Administration (FDA) and/or audited by Biogen Idec and/or a third party designated by Biogen Idec.

      I understand that noncompliance with the requirements of the TOUCH Prescribing Program may result in my pharmacy no longer being enrolled and termination of our participation in the program.

   c. Biogen Idec will:

      1. Ensure that pharmacies are notified when they are successfully enrolled in the TOUCH Prescribing Program, and therefore, are certified to dispense TYSABRI.
2. Ensure that pharmacies that dispense TYSABRI to authorized infusion sites have been trained on the known risks, potential benefits, and appropriate use of TYSABRI using approved educational materials.

d. The following materials are part of the TYSABRI REMS and are appended:

- TOUCH Prescribing Program Educational Slide Set
- TOUCH Prescribing Program Overview
- Certified Pharmacy Enrollment Form

e. Biogen Idec will ensure that infusion sites where TYSABRI is dispensed and administered are specially certified.

i. Infusion sites that dispense and administer TYSABRI must enroll in the TOUCH Prescribing Program by submitting a completed Infusion Site Enrollment Form and designating a person with appropriate authority to acknowledge the following:

a) The infusion site has received training and educational materials on the TOUCH Prescribing Program

1. I understand that TYSABRI will be administered only to patients who are currently authorized in the TOUCH Prescribing Program. Patient authorization must be confirmed prior to each infusion by:

   1. For TOUCH On-Line infusion sites: Patient Authorization Status must be “Authorized” or
   2. For paper-based infusion sites: Receipt of current Notice of Patient Authorization and verification that no Notice of Patient Discontinuation is on file

c) I understand that each patient will receive a copy of the TYSABRI Patient Medication Guide prior to each infusion

d) I understand that a TYSABRI Pre-infusion Patient Checklist must be completed prior to each infusion. The Pre-infusion Patient Checklist must be submitted to Biogen Idec within 1 business day of the patient visit, regardless of whether or not the patient received the infusion, by:

   1. For paper-based infusion sites: sending a copy of the completed Pre-infusion Patient Checklist to Biogen Idec. A copy must also be placed in the patient’s medical record
2. For TOUCH On-Line infusion sites: The infusion nurse can read, complete and submit the Pre-Infusion Patient Checklist directly in TOUCH On-Line

e) I understand that, per the requirements of the TOUCH Prescribing Program, this infusion site’s compliance with the REMS may be reviewed by FDA and/or audited by Biogen Idec and/or a third party designated by Biogen Idec.

f) I understand that noncompliance with the requirements of the TOUCH Prescribing Program will result in de-enrollment of the infusion site.

f. Biogen Idec will:

i. Ensure that infusion sites are notified when they are successfully enrolled in the TYSABRI REMS Program, and therefore, are certified to dispense and administer TYSABRI.

ii. Ensure that infusion sites that dispense and administer TYSABRI have been trained on the known risks, potential benefits, and appropriate use of TYSABRI using approved educational materials.

g. The following materials are a part of the TYSABRI REMS and are appended:

- TOUCH Prescribing Program Educational Slide Set
- TOUCH Prescribing Program Overview
- Infusion Site Enrollment Form
- Pre-Infusion Patient Checklist
- Medication Guide

3. TYSABRI will be dispensed to patients with evidence or other documentation of safe-use conditions.

a. Biogen Idec will ensure that certified prescribers enroll all patients into the TOUCH Prescribing Program by completing the Prescriber/Patient Enrollment Form for each new patient.

b. A completed and signed Prescriber/Patient Enrollment Form must be submitted to Biogen Idec before the patient may receive an infusion.

The Prescriber/Patient Enrollment Form requires the patient to acknowledge the following:
i. I understand that TYSABRI increases my chance of getting a rare brain infection that usually leads to death or severe disability

a) This infection is called progressive multifocal leukoencephalopathy (PML). PML usually happens in people with weakened immune systems

b) There is no known treatment, prevention, or cure for PML

c) I should call my doctor right away if I get any new or worsening symptoms that last several days, especially nervous system symptoms, while I am taking TYSABRI, and for at least 6 months after I stop taking TYSABRI. Some of these symptoms include a new or sudden change in my thinking, eyesight, balance, or strength, but I should also report other new or worsening symptoms

d) My chance for getting PML increases if I:
   a. Have received TYSABRI for a long time, especially longer than 2 years
   b. Have received certain medicines that can weaken my immune system before I start receiving TYSABRI
   c. Have been exposed to John Cunningham Virus (JCV). JCV is a common virus that is harmless in most people but can cause PML in people who have weakened immune systems, such as people taking TYSABRI. Most people who are exposed to JCV do not know it or have any symptoms. This exposure usually happens in childhood. My doctor may do a blood test to check if I have been exposed to JCV before I start receiving TYSABRI or during my treatment

My risk of getting PML is greatest if I have all 3 risk factors listed above. There may be other risk factors for getting PML during TYSABRI treatment that we do not know about yet. My doctor should discuss the risks and benefits of TYSABRI treatment with me before I decide to receive TYSABRI.

ii. Acknowledgments specific to MS

I understand that TYSABRI is a medicine approved to treat patients with relapsing forms of multiple sclerosis (MS)

a) TYSABRI increases the risk of PML. I understand that when starting and continuing treatment with TYSABRI, I should talk to
my doctor about whether the expected benefit of TYSABRI is enough to outweigh this risk (see important information about PML below)

b) I have talked to my doctor and understand the benefits and risks of TYSABRI treatment

c) My chance for getting PML may be higher if I am also being treated with other medicines that can weaken my immune system, including other MS treatments. Even if I use TYSABRI alone to treat my MS, I can still get PML

iii. Acknowledgments specific to CD

I understand that TYSABRI is a medicine approved to treat patients with moderate to severe Crohn’s Disease who have not been helped enough by, or cannot tolerate, usual Crohn’s disease medicines and medicines called tumor necrosis factor (TNF) inhibitors

a) I have talked to my doctor and understand the benefits and risks of TYSABRI treatment

b) I understand that I should not take certain medicines that weaken the immune system while I am taking TYSABRI

My chance for getting PML may be higher if I am also being treated with other medicines that can weaken my immune system, including other Crohn’s disease treatments. Even if I use TYSABRI alone to treat my Crohn’s disease, I can still get PML.

c. To receive TYSABRI all patients must be enrolled in a special program called the TOUCH Prescribing Program.

i. The TOUCH Prescribing Program is run by the company that makes TYSABRI. Under this program, the company is required to collect information about my health at regular time periods. I cannot receive TYSABRI if I do not agree to follow the requirements of the TOUCH Prescribing Program

ii. The company may use my information to meet the requirements of the TOUCH Prescribing Program, including helping me locate an authorized infusion site
iii. I must notify the TOUCH Prescribing Program if I switch physicians or infusion sites

iv. I have received, read, and understand the Patient Medication Guide

v. I will bring to each TYSABRI infusion a list of all medicines and treatments that I have taken during the last month

The following materials are part of the TYSABRI REMS and are appended:

- Prescriber/Patient Enrollment Forms (specific to MS or CD)
- Medication Guide

Biogen Idec will make these materials available for the enrolled prescribers to provide to their patients.

C. Implementation System

The Implementation system for the TYSABRI REMS includes the following:

a. Biogen Idec must maintain a secure, validated database of all certified prescribers, certified pharmacies, certified infusion sites, and enrolled patients in the TOUCH Prescribing Program. The TOUCH Prescribing Program is an integrated, computerized, validated database that captures enrollment, patient tracking, and drug distribution data.

b. Biogen Idec will monitor the distribution of TYSABRI to ensure that the drug is only delivered to certified pharmacies and infused by certified infusion sites.

c. Biogen Idec will monitor the compliance of certified entities (i.e. pharmacies, infusion sites, and prescribers) as acknowledged in the signed entity specific enrollment form and, if appropriate, institute corrective actions, which could include retraining and de-enrollment.

d. Biogen Idec will maintain TOUCH On-Line, which is an internet-based system that will allow enrolled TOUCH Prescribing Program participants the option of real-time access to view and/or submit required or pertinent patient therapy information collected through the TOUCH Prescribing Program, such as information on the Pre-infusion Patient Checklist, the TYSABRI Patient Status Report and Reauthorization Questionnaire, and the TYSABRI Patient Discontinuation Questionnaire.
e. Biogen Idec will monitor and evaluate the implementation of the elements to assure safe use and take reasonable steps to work to improve implementation of these elements.

D. Timetable for Submission of Assessments of the REMS

Biogen Idec will submit REMS Assessments annually from the date of the initial approval of the REMS (October 7, 2011). To facilitate inclusion of as much information as possible while allowing reasonable time to prepare the submission, the reporting interval covered by each assessment should conclude no earlier than 60 days before the submission date for that assessment. Biogen will submit each assessment so that it will be received by the FDA on or before the due date.
Guidance for evaluation of new neurological symptoms in patients receiving TYSABRI

Please see accompanying full Prescribing Information, including Boxed Warning.
Background information

**Progressive multifocal leukoencephalopathy (PML)**

PML is a demyelinating disease that attacks the central nervous system (CNS). It is caused by a polyomavirus called the JC virus (JCV), which is common and widespread in humans. JCV usually remains latent, typically causing PML only in the setting of immunodeficiency.

The clinical picture of PML or other neurological conditions can be difficult to distinguish from multiple sclerosis (MS), especially early in the disease course. Therefore, this information is intended to offer an overview of some of the key issues regarding the definitive diagnosis of PML, especially as they relate to treatment with TYSABRI. These include:

- Patient monitoring and management
  - Obtaining a pretreatment MRI
  - Performing regular follow-ups
  - Treatment of relapses or other neurological symptoms

- Evaluation of new neurological symptoms in patients receiving TYSABRI
  - Distinguishing PML from MS
  - Suggested diagnostic algorithm
  - Action steps if PML is suspected, including MRI assessment, JCV testing, and plasma exchange (PLEX)
  - Immune Reconstitution Inflammatory Syndrome (IRIS)


Please see accompanying full Prescribing Information, including Boxed Warning.
Patient monitoring and management

Management of patients receiving TYSABRI

Pretreatment MRI

Obtaining a pretreatment brain MRI scan is recommended. It may assist in determining whether MRI lesions noted at the time of new neurological signs or symptoms were preexistent. This may assist in the differential diagnosis between PML and MS activity.

Regular follow-ups

All patients treated with TYSABRI should have regular clinical follow-ups to allow for early detection of changes in neurological status. To that end, Biogen Idec in conjunction with the Food and Drug Administration (FDA), developed a risk management plan for the United States called the TOUCH® Prescribing Program. As part of the TOUCH Prescribing Program:

➢ Physicians evaluate the patient 3 months after the first infusion, 6 months after the first infusion, every 6 months thereafter, and for at least 6 months after discontinuing Tysabri.

➢ Physicians submit the TYSABRI Patient Status Report and Reauthorization Questionnaire to Biogen Idec 6 months after initiating treatment and every 6 months thereafter, ensuring additional monitoring and reporting by Biogen Idec.

➢ Infusion sites administer the Pre-Infusion Patient Checklist and report to the prescriber any changes in the patient's status prior to infusing

— Infusion sites will not infuse TYSABRI if the patient reports a change in symptoms, unless the prescriber authorizes the infusion

Patient history

Knowing the history and pattern of prior and ongoing MS signs and symptoms can help in the management of patients treated with TYSABRI.
Evaluation of new neurological symptoms in patients receiving TYSABRI

If new neurological symptoms develop, withhold TYSABRI dosing and evaluate the patient

Distinguishing PML from MS

The following information should be considered when undertaking the assessment and management of new or worsening neurological symptoms in MS patients treated with TYSABRI. There are no pathognomonic signs or symptoms that distinguish an MS relapse from PML, but there are certain clinical features that may help differentiate between the 2 conditions (see Table 1).

Table 1. Clinical signs and symptoms typical of MS relapse and PML

<table>
<thead>
<tr>
<th></th>
<th>MS relapse</th>
<th>PML</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>ONSET</strong></td>
<td>Acute</td>
<td>Subacute</td>
</tr>
<tr>
<td><strong>EVOLUTION</strong></td>
<td>Over hours to days</td>
<td>Days to weeks</td>
</tr>
<tr>
<td></td>
<td>Normally stabilize</td>
<td>Progressive</td>
</tr>
<tr>
<td></td>
<td>Resolve spontaneously or with treatment</td>
<td></td>
</tr>
<tr>
<td><strong>CLINICAL</strong></td>
<td>Diplopia</td>
<td>Cortical symptoms/signs</td>
</tr>
<tr>
<td><strong>PRESENTATION</strong></td>
<td>Paresthesia</td>
<td>Behavioral and neuropsychological alteration</td>
</tr>
<tr>
<td></td>
<td>Paraparesis</td>
<td>Retrochiasmal visual deficits</td>
</tr>
<tr>
<td></td>
<td>Optic neuritis</td>
<td>Seizures</td>
</tr>
<tr>
<td></td>
<td>Myelopathy</td>
<td>Hemiparesis</td>
</tr>
</tbody>
</table>

Not intended to be inclusive of all clinical signs and symptoms indicative of MS and PML.

Please see accompanying full Prescribing Information, including Boxed Warning.
**Treatment of MS relapse**

- Relapses should be managed according to usual clinical practice

**If treating with corticosteroids:**

- A single short course of corticosteroids can be considered for cases in which PML is unlikely on clinical grounds
- Progression of symptoms, despite treatment with corticosteroids, should trigger further investigation

- In addition to PML and MS, other medical and CNS conditions including other infections should be considered when evaluating a patient with new neurological symptoms

**New or recurrent neurological symptoms should prompt careful evaluation.**
**Suggested diagnostic algorithm for TYSABRI-treated patients experiencing new neurological symptoms suggestive of non–MS-related disease**

1. **SUSPEND DOSING**
   - New neurological symptoms suggestive of non–MS-related disease
     - If PML is suspected based on clinical presentation and an MRI is not readily available, cerebrospinal fluid (CSF) assessment to exclude PML should be considered prior to MRI.

2. **MRI assessment**
   - PML excluded
   - Cannot exclude PML
     - JCV not detected and low clinical suspicion: Dosing may be resumed
     - JCV not detected and high clinical suspicion: Repeat assessment

3. **CSF assessment**
   - JCV detected: Treat as PML
   - JCV not detected: Monitor for IRIS

**Note:** TYSABRI dosing should only be restarted when the diagnosis of PML is excluded, if necessary, by repeating clinical, MRI, and CSF assessment if clinical suspicion of PML remains.

**CSF assessment for presence of JCV DNA should be performed using a highly sensitive quantitative real-time PCR assay with a limit of quantification (LOQ) of at least 50 copies/mL.**

**For more information, please call Biogen Idec Medical Information at 1-800-456-2255.**

Please see accompanying full Prescribing Information, including Boxed Warning.
**Action steps if PML is suspected**

1. **Suspend dosing**
   TYSABRI dosing should be suspended immediately in all cases in which PML is suspected.

2. **MRI assessment**
   If the clinical presentation is suggestive of PML, further investigation should include brain MRI evaluation as soon as possible.

3. **CSF assessment**
   If MRI evaluation reveals lesions suspicious for PML (see Table 2 on page 9), a lumbar puncture with evaluation of CSF for the detection of JCV-DNA should be undertaken with a highly sensitive quantitative real-time PCR assay.

4. **Repeat testing**
   If clinical suspicion of PML remains despite a negative evaluation, then MRI and CSF assessments should be repeated to exclude a diagnosis of PML.

A definitive diagnosis of PML is made by evaluating clinical and MRI findings plus the identification of JCV in the CNS.

- There is no prevention, FDA-approved treatment, or cure for PML. Rapid recognition of PML and early discontinuation of TYSABRI are key interventions
- PLEX (see page 10) may be considered as a means to accelerate the clearance of TYSABRI
- Healthcare providers should promptly report serious adverse events to Biogen Idec at 1-800-456-2255
  - Biogen Idec can provide additional resources, including suggested next steps

This information is provided as an educational resource for healthcare providers and should be considered current as of the date listed herein. It is not intended to be a substitute for consultation and review of reference materials and medical literature pertaining to individual clinical circumstances. Healthcare providers should make all treatment decisions based on the context of the situation and their own clinical judgment.
MRI assessment

- Although there are no pathognomonic findings that differentiate PML from MS, a brain MRI scan that includes fluid-attenuated inversion recovery (FLAIR) and T1- and T2-weighted sequences, with and without gadolinium (Gd), should be performed to assess patients with neurological changes suggestive of PML (see Table 1 on page 5).
- Comparison with a baseline scan may assist with interpretation of the findings on the new MRI. See Figure 1 and Table 2 for differences in lesion characteristics that may help differentiate between PML and MS.

Figure 1. MRI presentation features of PML

<table>
<thead>
<tr>
<th>T1-weighted imaging</th>
<th>T2-weighted imaging</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image1" alt="T1-weighted image" /></td>
<td><img src="image2" alt="T2-weighted image" /></td>
</tr>
</tbody>
</table>

- Large hypointense lesion in the region of the right occipital lobe (angular gyrus and intraparietal sulcus).
- Typical multifocal affection of PML, with additional lesions in the white matter of the left frontal temporal and occipital lobes. Note sparing of gray matter.


Please see accompanying full Prescribing Information, including Boxed Warning.
Table 2. MRI lesion characteristics typical of PML and MS

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>MS Lesions</th>
<th>PML Lesions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Location</td>
<td>Periventricular perpendicular to ventricles (Dawson’s fingers), deep white matter, isolated U fibers, cerebellum and spinal cord</td>
<td>Subcortical WM in parietal, occipital or frontal lobes May involve precentral or postcentral gyrus (motor/sensory cortex) or insular region Follows WM tracks. Can cross the corpus callosum to contralateral hemisphere (butterfly pattern) or extend through internal capsule Rarely brainstem or cerebellar WM No spinal cord involvement</td>
</tr>
<tr>
<td>Appearance</td>
<td>Well defined borders</td>
<td>Infiltrating, ill-defined, confluent WM lesions which can be multifocal</td>
</tr>
<tr>
<td>Mass effect</td>
<td>Large lesions can have a mass effect</td>
<td>Rare even in large lesions</td>
</tr>
<tr>
<td>FLAIR</td>
<td>Flair = T2</td>
<td>Flair more sensitive for detection of PML lesions in subcortical location</td>
</tr>
<tr>
<td>T1W pre-contrast</td>
<td>Isotense or mildly hypointense to Grey matter</td>
<td>Isointense with progressive hypointensity</td>
</tr>
<tr>
<td>T1 post contrast</td>
<td>Homogeneous or ring-enhancement - resolves in 1-2 months</td>
<td>Patchy, punctate or linear</td>
</tr>
</tbody>
</table>


This information is provided as an educational resource for healthcare providers and should be considered current as of the date listed herein. It is not intended to be a substitute for consultation and review of reference materials and medical literature pertaining to individual clinical circumstances. Healthcare providers should make all treatment decisions based on the context of the situation and their own clinical judgment.
**JCV DNA testing to confirm diagnosis**

➤ Plasma assessment

  Presence of JCV DNA in plasma has not been correlated with the development of PML.
  
  Plasma JCV DNA test positivity is highly variable, so the sensitivity and predictive value of this screening method are unclear.
  
  Plasma JCV DNA testing is not included in the TOUCH Prescribing Program.

➤ CSF assessment

  The detection of JCV DNA in the CSF of a patient with clinical and MRI features suggestive of PML establishes the diagnosis of PML.
  
  If clinical suspicion of PML remains despite a negative CSF testing, a repeat should be considered.
  
  It is recommended to test samples using a validated ultrasensitive quantitative PCR test that has a lower limit of quantification of 50 copies per mL or lower.

➤ Brain biopsy

  If diagnosis remains uncertain and suspicion of PML remains high, a brain biopsy may be considered to establish a definitive diagnosis.

**Note:** TYSABRI dosing should only be resumed if the diagnosis of PML is excluded and if deemed appropriate for the ongoing treatment of MS.

**PLEX**

➤ Three sessions of plasma exchange (PLEX) over 5 to 8 days were shown to accelerate TYSABRI clearance in a study of 12 patients with MS who did not have PML, although in the majority of patients, α4-integrin receptor binding remained high—a potential sign of continued inhibition of α4-integrin–mediated leukocyte activity.

➤ Additional plasma exchanges (up to a total of 5 over a 10-day period) may more consistently reduce TYSABRI plasma concentration and α4-integrin receptor binding to below subtherapeutic levels.

➤ Adverse events that may occur during PLEX include clearance of other medications and volume shifts, which have the potential to lead to hypotension or pulmonary edema.

➤ Although plasma exchange has not been studied in TYSABRI treated patients with PML, it has been used in such patients in the postmarketing setting to remove TYSABRI more quickly from the circulation.


Please see accompanying full Prescribing Information, including Boxed Warning.
Immune Reconstitution Inflammatory Syndrome (IRIS)

- IRIS has been reported in the majority of patients who developed PML and subsequently discontinued TYSABRI.
- In almost all cases, IRIS occurred within days to several weeks after plasma exchange was used to accelerate TYSABRI clearance.
- IRIS usually presents as an unanticipated clinical decline which may be rapid and severe, and may be fatal.
- At the time of IRIS, MRI may show additional changes including Gd enhancement.
- Monitoring for development of IRIS and appropriate treatment of the associated inflammation should be undertaken.

For the latest scientific information about our products or to report an adverse event, contact:

**Biogen Idec Medical Information**

**Telephone:** 1-800-456-2255 (8:30 AM to 8:00 PM ET)

**Fax:** Send Medical Information Request Form to 1-877-462-1530

**Mail:** Biogen Idec Medical Information
14 Cambridge Center
Cambridge, MA 02142

**Web:** http://medinfo.biogenidec.com

**E-mail:** medinfo@biogenidec.com

To report an adverse event, contact:

**Biogen Idec**

**Telephone:** 1-800-456-2255

This information is provided as an educational resource for healthcare providers and should be considered current as of the date listed herein. It is not intended to be a substitute for consultation and review of reference materials and medical literature pertaining to individual clinical circumstances. Healthcare providers should make all treatment decisions based on the context of the situation and their own clinical judgment.
Please see accompanying full Prescribing Information, including Boxed Warning.
**Patient Information**

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*Please attach copies of both sides of patient's insurance and pharmacy card(s).*

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**Patient Authorization to Use/Disclose Health Information**

I understand that I have certain rights related to the collection, use, and disclosure of my medical and health information. This information is called “protected health information” (PHI) and includes demographic information (such as sex, race, date of birth, etc.), the results of physical examinations, clinical tests, blood tests, X-rays, and other diagnostic and medical procedures that may be included in my medical records. Biogen Idec will not use my PHI without my consent.

This Authorization form applies to PHI created or obtained by my prescriber, my infusion site, my pharmacy, and my health insurance company. I understand that by signing this Authorization, I authorize my prescriber, infusion site, pharmacy, and/or health insurance company to disclose the PHI in my medical records to Biogen Idec Inc. and its representatives or agents, including information related to my medical condition, treatment, and health insurance, as well as all information provided on any prescription. I also authorize Biogen Idec to use this information to provide TYSABRI support services, such as investigating insurance coverage for TYSABRI and coordinating delivery of TYSABRI to the prescriber or infusion site administering TYSABRI (which may include forwarding my health information to a pharmacy).

I agree to allow Biogen Idec to ask me about and provide me with these support services, educational kits, and other information related to TYSABRI and/or my medical condition. I understand that, once my PHI has been disclosed to Biogen Idec, federal privacy laws may no longer protect the information. However, Biogen Idec agrees to protect my PHI by using it only for the purposes authorized in this Authorization or as required by law.

I understand that I may refuse to sign this Authorization, and refusing to do so will affect my eligibility to receive these additional services but will not affect my ability to receive TYSABRI. I understand that signing this Authorization will not change how my healthcare providers, health insurance plan, and pharmacies provide my medical treatment or payment for treatment or insurance benefits.

I understand that I may cancel all or a part of this Authorization at any time by mailing a letter requesting such cancellation to TYSABRI Support Services, 5000 Davis Drive, PO Box 13919, Research Triangle Park, NC 27709. If I cancel this Authorization, Biogen Idec will end further use and disclosure of my PHI as soon as possible. This will not affect health information that has already been used or disclosed in reliance upon this Authorization.

I will receive a copy of this signed Authorization. This Authorization expires ten (10) years from the date this Authorization is signed.

**Patient signature** (or personal representative): __________________________ Date: ____________________

Authority of personal representative *(if applicable)*: __________________________

Reference ID: 3753128
Biogen Idec considers patient safety a priority. Read each section below and initial in the space provided if you understand the information. Do not sign this form if there is anything you do not understand about all the information you have received. Ask your doctor about anything you do not understand before you initial and sign this form.

I understand that TYSABRI is a medicine approved to treat patients with relapsing forms of multiple sclerosis (MS).

- I have talked to my doctor and understand the benefits and risks of TYSABRI treatment
- Tysabri increases the risk of PML. I understand that when starting and continuing treatment with Tysabri, I should talk to my doctor about whether the expected benefit of Tysabri is enough to outweigh this risk (see important information about PML below).

Initials:______

I understand that TYSABRI increases my chance of getting a rare brain infection that usually leads to death or severe disability.

- This infection is called progressive multifocal leukoencephalopathy (PML). PML usually happens in people with weakened immune systems
- There is no known treatment, prevention, or cure for PML
- My chance for getting PML may be higher if I am also being treated with other medicines that can weaken my immune system, including other MS treatments. Even if I use TYSABRI alone to treat my MS, I can still get PML
- My chance for getting PML increases if I:
  - Have received TYSABRI for a long time, especially longer than 2 years
  - Have received certain medicines that can weaken my immune system before I start receiving TYSABRI
  - Have been exposed to John Cunningham Virus (JCV). JCV is a common virus that is harmless in most people but can cause PML in people who have weakened immune systems, such as people taking TYSABRI. Most people who are exposed to JCV do not know it or have any symptoms. This exposure usually happens in childhood. My doctor may do a blood test to check if I have been exposed to JCV before I start receiving TYSABRI or during my treatment
- My risk of getting PML is greatest if I have all 3 risk factors listed above. There may be other risk factors for getting PML during TYSABRI treatment that we do not know about yet. My doctor should discuss the risks and benefits of TYSABRI treatment with me before I decide to receive TYSABRI
- I should call my doctor right away if I get any new or worsening symptoms that last several days, especially nervous system symptoms, while I am taking TYSABRI, and for at least 6 months after I stop taking TYSABRI. Some of these symptoms include a new or sudden change in my thinking, eyesight, balance, or strength, but I should also report other new or worsening symptoms

Initials:______

To receive TYSABRI, all patients must be enrolled in a restricted program called the TOUCH Prescribing Program.

- The TOUCH Prescribing Program is run by the company that makes TYSABRI. Under this program, the company is required to collect information about my health at regular time periods. I cannot receive TYSABRI if I do not agree to follow the requirements of the TOUCH Prescribing Program. I understand that the TOUCH Prescribing Program does not require me to sign the Authorization included on page 1 of this form
- The company may use my information to meet the requirements of the TOUCH Prescribing Program, including helping me locate an authorized infusion site
- I must notify the TOUCH Prescribing Program if I switch physicians or infusion sites
- I have received, read, and understand the Patient Medication Guide
- I will bring to each TYSABRI infusion a list of all medicines and treatments that I have taken during the last month

Initials:______

Patient name: ___________________________ Date of birth: __________/________/________ (MM/DD/YYYY)

Patient signature (or personal representative): ___________________________ Date: __________

Authority of personal representative (if applicable): ___________________________
Patient History

at ent name: ___________________________ DOB: __________/________/________ (MM/DD/YYYY)

F rst MI Last  

Date of f rst symptoms: __________/________/________ (MM/DD/YYYY)

Ease nd cate the pat ent’s MOST RECENT therapy for (if pat ent was most recent y on comb nat on therapy, check a that app y). None □

A ONEX® □ PLEGIRIDY □ Beta seron® □ Copaxone® □ eb f® □ YSABRI® □ Extav a® □ G enya™ □

TECFIDERA® □ Aubag o® □ LEMTRADA™ □ Azath opr ne □ Methotrexate □ to anxrone □ Mycopheno ate □

Cyc ophospham de □ Other □

Ease nd cate the start and stop dates of most recent therapy: Start date __________/________/________ Stop date __________/________/________

Has the pat ent ever rece ved YSABRI before? Yes □ No □

Has the pat ent eVeR been prescr bed an immunosuppressant or an ant neop ast c therapy for any cond t on? Yes □ No □

If yes, p ease check a of the fo ow ng that app y:

Azath opr ne □ Cyc ophospham de □ Methotrexate □ to anxrone □ Mycopheno ate □ Other □

Has the pat ent eVeR been tested for the presence of ant-JCV ant bod es? Yes □ No □ nknown □

If yes, has the pat ent eVeR tested POSITIVE for the presence of ant-JCV ant bod es? Yes □ No □ end ng □

Prescription for TYSABRI

Dose: TYSABRI® (natalizumab) 300 mg  Dispense: 1 vial  Refills: 12  Directions: IV infusion per Prescribing Information every 4 weeks

I author ze Bo gen Idec as my des gnated agent and on beha f of my pat ent to (1) use the nformat on on th s form to enro the above-named pat ent n the OUCH Prescr b ng Program, (2) furn sh any nformat on on th s form to the nsurer of the above-named pat ent, (3) forward the nformat on on th s form to the prescr ber or nfus on s te adm n ster ng YSABRI, f app cab e, (4) forward the above prescr pt on by fax or by another mode of de very to a pharmacy, f app cab e, and (5) coord nate de very of YSABRI on beha f of the above-named pat ent.

Prescriber signature (stamps not acceptab e): ______________________________ Date: __________________

Prescriber

Prescr ber name: ___________________________ MI  Last

Street address

C ty State ZIP

e p hone □□□□□-□□□□□-□□□□□□□ Fax □□□□□-□□□□□-□□□□□□□

Off ce contact

ax ID #

DEA #

NPI/UPIN/prov der ID # w th pat ent’s nsurer(s)

Continued on next page
Prescriber Acknowledgment

- I have read and understand the full Prescribing Information for TYSABRI
- I understand that TYSABRI is indicated as monotherapy for relapsing forms of MS
- I understand that this patient has a relapsing form of MS based on clinical and radiological evidence
- I understand that TYSABRI increases the risk of progressive multifocal leukoencephalopathy (PML), an opportunistic viral infection of the brain that usually leads to death or severe disability. When initiating and continuing treatment with TYSABRI, I should consider whether the expected benefit of TYSABRI is sufficient to offset this risk
- I am aware that cases of PML have been reported in patients taking TYSABRI who were recently or concomitantly treated with immunomodulators or immunosuppressants, as well as in patients receiving TYSABRI monotherapy
- I understand that three risk factors identified thus far that increase the risk of PML in TYSABRI treated patients are:
  - Longer treatment duration, especially beyond 2 years
  - Prior treatment with an immunosuppressant (e.g., mitoxantrone, azathioprine, methotrexate, cyclophosphamide, mycophenolate mofetil)
  - The presence of anti JCV antibodies
  These factors should be considered in the context of expected benefit when initiating and continuing treatment with TYSABRI
- I understand that TYSABRI is not ordinarily recommended for patients who are receiving chronic immunosuppressant or immunomodulatory therapy, or who are significantly immunocompromised from any other cause
- To my knowledge, this patient has no known contraindications to TYSABRI treatment, including PML
- I understand that an MRI should be performed prior to initiating therapy with TYSABRI in MS patients
- I have instructed this patient to promptly report to me any continuously worsening symptoms that persist over several days, especially nervous system symptoms
- I understand that this patient should be seen and evaluated 3 months after the first infusion, 6 months after the first infusion, every 6 months thereafter, and for at least 6 months after TYSABRI has been discontinued
- I will determine every 6 months whether this patient should continue on TYSABRI and if so, authorize treatment for another 6 months. I understand that I am required to submit an "Initial Discontinuation Questionnaire" when TYSABRI is discontinued and a "6 Month Discontinuation Questionnaire" following discontinuation of TYSABRI
- I should report, as soon as possible, cases of PML, hospitalizations due to opportunistic infection, or deaths to Biogen Idec
- I understand that data concerning this patient and me will be entered into the mandatory TOUCH Prescribing Program. Biogen Idec requires my cooperation with periodic data collection. Failure to provide the requested information or otherwise comply with the requirements of the TOUCH Prescribing Program may result in discontinuation of TYSABRI treatment for this patient and termination of my authorization to prescribe TYSABRI
- I have received educational materials regarding the benefits and risks of TYSABRI treatment
- I have, or another healthcare provider under my direction has, educated this patient on the benefits and risks of treatment with TYSABRI, provided him or her with the Patient Medication Guide and Enrollment Form, instructed him or her to read these materials, and encouraged him or her to ask questions when considering TYSABRI

Patient name: ____________________________ M __________ Last ____________________________
Date of birth: __________ / __________ / ________  (MM/DD/YYYY)

Prescriber signature: ____________________________ Date: __________

Reference ID: 3753128

4 of 5

3-3033-XX
**Patient Information**

Date of birth ___________________ / ___________________ / ___________________<br> (MM/DD/YYYY)<br> Patient name ___________________<br> First _____________ M _____________ Last _____________

In addition, I authorize the sharing of my health information to the person or people I name below. Biogen Idec may contact the people named below to discuss my enrollment in the TOUCH Program.

Designated Individual (patient name): ___________________<br> Relationship: ___________________<br>

**Infusion Site Information**<br><br>**1.** Prescriber administering TYSABRI and request the following services (check one):<br> - [ ] No services requested OR [ ] Forward this prescription to a specialty pharmacy OR [ ] Please conduct insurance research and procurement options for TYSABRI

**2.** Please be efe TYSABRI eae a hee (check one):<br> - [ ] I require assistance in obtaining an infusion site OR [ ] I am referring the patient to the following infusion site or healthcare provider:

Name of administering healthcare provider (first, last)<br> Office contact<br> Street name<br> Street address or state Authorization Number<br> City _____________ State _____________ ZIP _____________<br> Telephone _____________ _____________ _____________ _____________ _____________<br> Fax _____________ _____________ _____________ _____________ _____________

*Note: TYSABRI can only be infused at authorized infusion sites. Biogen Idec will contact you if the infusion site you have indicated is not authorized to infuse TYSABRI.

Please see accompanying prescribing information, including boxed warning, for safety information.
Dear <MD Name>,

Our records indicate that <Patient Name>’s authorization to receive TYSABRI will expire on <MM/DD/YYYY> and he/she will no longer be able to receive TYSABRI. Please submit the completed form to Biogen Idec via TOUCH On-Line (www.touchprogram.com) or fax (1-800-840-1278) by <expiration date> and place a copy in the patient’s record.

A. Is the patient still under <MD name>’s care?
   - Yes
   - No/I don’t know

   If No, please provide name and phone number for new prescriber, if available

B. Is the patient alive?
   - Yes
   - No

Since starting TYSABRI therapy has the patient been diagnosed with any of the following that you have not reported to Biogen Idec:

C. Progressive Multifocal Leuкоencephalopathy (PML)
   - Yes
   - No or Under investigation

D. Opportunistic Infection*
   - For which they have been hospitalized
   - Yes
   - No or Under investigation

    Malignancy
   - Yes
   - No or Under investigation

Since <last authorization>, has the patient been tested for the presence of anti-JCV antibodies?
   - Yes
   - Not performed

   If performed, test result:
   - Positive
   - Negative
   - Pending

G. Is the patient currently receiving or has the patient received intermittent courses of steroids for the treatment of MS relapse in the previous 6 months?
   - Yes
   - No

   If Yes, please circle the number of courses received.
   1 2 3 4 5 6 >6

H. Is the patient currently receiving or has the patient received any IMMUNOMODULATORY or IMMUNOSUPPRESSANT products in the previous 6 months?
   - Yes
   - No

   If Yes, please indicate the type of therapy and number of months of use.

   Months of Use in Last 6 Months

   AVONEX® 1 2 3 4 5 6
   Betaseron® 1 2 3 4 5 6
   Copaxone® 1 2 3 4 5 6
   Rebif® 1 2 3 4 5 6
   Extavia® 1 2 3 4 5 6
   Gilenya™ 1 2 3 4 5 6
   Aubagio® 1 2 3 4 5 6
   PLEGRIDY™ 1 2 3 4 5 6
   TECFIDERA® 1 2 3 4 5 6
   LEMTRADA™ 1 2 3 4 5 6
   Azathioprine 1 2 3 4 5 6
   Methotrexate 1 2 3 4 5 6
   Mitoxantrone 1 2 3 4 5 6
   Mycophenolate 1 2 3 4 5 6
   Cyclophosphamide 1 2 3 4 5 6
   Chronic systemic steroids 1 2 3 4 5 6
   Other immunomodulatory or immunosuppressant therapy 1 2 3 4 5 6

If the patient is still under <MD name>’s care DO YOU AUTHORIZE the continuation of TYSABRI treatment for the next 6 months for the patient?
   - Yes
   - No

   If you answer No, Biogen Idec will contact the patient and the infusion site to STOP TYSABRI TREATMENT will not be eligible to receive TYSABRI treatment, and you will receive a final questionnaire for this patient in 6 monts.

   The patient

   If the patient is still under <MD name>’s care does the patient have any ORGANIZATIONAL or IMMUNOSUPPRESSANT therapy?
   - Yes
   - No or Under investigation

   If Yes, please indicate the type of therapy and number of months of use.

   Months of Use in Last 6 Months

   AVONEX® 1 2 3 4 5 6
   Betaseron® 1 2 3 4 5 6
   Copaxone® 1 2 3 4 5 6
   Rebif® 1 2 3 4 5 6
   Extavia® 1 2 3 4 5 6
   Gilenya™ 1 2 3 4 5 6
   Aubagio® 1 2 3 4 5 6
   PLEGRIDY™ 1 2 3 4 5 6
   TECFIDERA® 1 2 3 4 5 6
   LEMTRADA™ 1 2 3 4 5 6
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   Methotrexate 1 2 3 4 5 6
   Mitoxantrone 1 2 3 4 5 6
   Mycophenolate 1 2 3 4 5 6
   Cyclophosphamide 1 2 3 4 5 6
   Chronic systemic steroids 1 2 3 4 5 6
   Other immunomodulatory or immunosuppressant therapy 1 2 3 4 5 6

Please Note: A TOUCH authorized physician may complete this form on behalf of the Prescriber of record. This questionnaire will be used consistent with the TOUCH Prescriber/Patient Enrollment Form signed by you and your patient and with HIPAA and applicable privacy rules.

Please see full Prescribing Information, including Boxed Warning, at www.TYSABRI.com

Reference ID: 3753128

All other trademarks are the marks of their respective owners.
Is the patient still under <MD name>'s care?

☐ Yes    ☐ No/I don't know

If No, please provide name and phone number for new prescriber, if available

---

Is the patient alive?

☐ Yes    ☐ No

---

Since starting TYSABRI therapy has the patient been diagnosed with any of the following that you have not reported to Biogen Idec:

**PROGRESSIVE MULTIFOCAL LEUKOENCEPHALOPATHY (PML)**

☐ Yes    ☐ No or ☐ Under investigation

**OPPORTUNISTIC INFECTION** for which they have been hospitalized

☐ Yes    ☐ No or ☐ Under investigation

**MALIGNANCY**

☐ Yes    ☐ No or ☐ Under investigation

Since <last authorization>, has the patient been tested for the presence of anti-JCV antibodies?

☐ Yes    ☐ Not performed

If performed, test result:

☐ Positive    ☐ Negative    ☐ Pending

---

Prescriber signature: ___________________________ Date (MM/DD/YYYY): _________ / _________ / _________

(If applicable) Print TOUCH Authorized Prescriber Delegate Name: ___________________________

---

Please Note: A TOUCH authorized physician may complete this form on behalf of the Prescriber of record. This questionnaire will be used consistent with the TOUCH Prescriber/Patient Enrollment Form signed by you and your patient and with HIPAA and applicable privacy rules. If you have questions, or if you need additional information, please call 1-800-456-2255 from 8:30 AM to 8:00 PM (ET).

Please see full Prescribing Information, including Boxed Warning, at www.TYSABRI.com
Is the patient still under <MD name>’s care?

☐ Yes  ☐ No/ I don’t know

If No, please provide name and phone number for new prescriber, if available _______________________________

Is the patient alive?

☐ Yes  ☐ No

Since starting TYSABRI therapy has the patient been diagnosed with any of the following that you have not reported to Biogen Idec:

C PROGRESSIVE MULTIFOCAL LEUKOENCEPHALOPATHY (PML)

☐ Yes  ☐ No or ☐ Under investigation

D OPPORTUNISTIC INFECTION* for which they have been hospitalized

☐ Yes  ☐ No or ☐ Under investigation

E MALIGNANCY

☐ Yes  ☐ No or ☐ Under investigation

Since <last authorization>, has the patient been tested for the presence of anti-JCV antibodies?

☐ Yes  ☐ Not performed

If performed, test result:

☐ Positive  ☐ Negative  ☐ Pending

Prescriber signature: ________________________________ Date (MM/DD/YYYY): __________/_________/__________

(If applicable) Print TOUCH Authorized Prescriber Delegate Name: ________________________________

Please Note: A TOUCH authorized physician may complete this form on behalf of the Prescriber of record. This questionnaire will be used consistent with the TOUCH Prescriber/Patient Enrollment Form signed by you and your patient and with HIPAA and applicable privacy rules. If you have questions, or if you need additional information, please call 1-800-456-2255 from 8:30 AM to 8:00 PM (ET).

Please see full Prescribing Information, including Boxed Warning, at www.TYSABRI.com

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Understanding PML for Gastroenterologists
Understanding PML for Gastroenterologists

The following information should be considered when undertaking the assessment and management of progressive multifocal leukoencephalopathy (PML) in adult patients treated with TYSABRI for moderately to severely active Crohn’s disease (CD). During clinical trials for TYSABRI, 3 cases of PML were identified (2 in multiple sclerosis and 1 in Crohn’s disease). Both multiple sclerosis patients were receiving concomitant immunomodulatory therapy and the Crohn’s disease patient had been treated in the past with immunosuppressive therapy. In the postmarketing setting, additional cases of PML have been reported in multiple sclerosis and Crohn’s disease patients who were receiving no concomitant immunomodulatory therapy.1

About PML

PML is a demyelinating disease that attacks the central nervous system.2 It is an opportunistic infection caused by the JC virus that typically occurs in patients who are immunocompromised.1 The virus removes myelin that surrounds the nerves, and without this protection the nerves cannot transmit signals.3 There are no known interventions that can reliably prevent PML or adequately treat PML if it occurs.1

How to Recognize PML

Typical symptoms associated with PML are diverse, progress over days to weeks, and include3,4:

- Progressive weakness on one side of the body or clumsiness of limbs
- Disturbance of vision
- Changes in thinking, memory, and orientation, leading to confusion and personality changes
- Seizures

The progression of deficits usually leads to death or severe disability over weeks or months.3 Since these symptoms are very different from those of Crohn’s disease, the appearance of any symptom of PML, including those listed above, should be investigated immediately.1 In Crohn’s disease patients, a baseline brain MRI may also be helpful to distinguish pre­existent lesions from newly developed lesions, but brain lesions at baseline that could cause diagnostic difficulty while on TYSABRI therapy are uncommon.1
Action Steps if PML Is Suspected

- TYSABRI dosing should be suspended immediately in all cases in which PML is suspected.
- Immediate referral to a neurologist for assessment, potentially including:
  - A brain MRI to determine if lesions that could be due to PML are present.
  - Cerebrospinal fluid evaluation for the presence of JCV DNA.
- Potential cases of PML should be reported immediately to Biogen Idec at 1-800-456-2255, or to the FDA's MedWatch reporting system at 1-800-FDA-1088, or via the MedWatch Web site at www.fda.gov/medwatch.

Note: TYSABRI dosing should be restored only if the diagnosis of PML is excluded and if deemed appropriate for the ongoing treatment of CD in patients with moderately to severely active Crohn's disease with evidence of inflammation who have had an inadequate response to, or are unable to tolerate, conventional CD therapies and inhibitors of TNF-α, and who are not taking concomitant immunosuppressants (e.g., 6-mercaptopurine, azathioprine, or methotrexate) or concomitant inhibitors of TNF-α.

Indication

TYSABRI is indicated for inducing and maintaining clinical response and remission in adult patients with moderately to severely active Crohn's disease with evidence of inflammation who have had an inadequate response to, or are unable to tolerate, conventional CD therapies and inhibitors of TNF-α. TYSABRI should not be used in combination with immunosuppressants (e.g., 6-mercaptopurine, azathioprine, cyclosporine, or methotrexate) or inhibitors of TNF-α.

Important Safety Information

WARNING

TYSABRI (natalizumab) increases the risk of PML, an opportunistic viral infection of the brain that usually leads to death or severe disability. Risk factors for the development of PML include duration of therapy, prior use of immunosuppressants, and presence of anti-JCV antibodies. These factors should be considered in the context of expected benefit when initiating and continuing treatment with TYSABRI.

Healthcare professionals should monitor patients on TYSABRI for any new sign or symptom that may be suggestive of PML. TYSABRI dosing should be withheld immediately at the first sign or symptom suggestive of PML. For diagnosis, an evaluation including a gadolinium-enhanced MRI scan of the brain and, when indicated, cerebrospinal fluid analysis for JC viral DNA are recommended.

Because of the risk of PML, TYSABRI is available only through a restricted program under a Risk Evaluation and Mitigation Strategy (REMS) called the TOUCH Prescribing Program.

Important Safety Information Continued on next page
Important Safety Information (continued)

Progressive Multifocal Leuкоencephalopathy (PML)

- Infection by the JC Virus is required for the development of PML.
- Anti-JCV antibody testing should not be used to diagnose PML.
- There are no known interventions that can reliably prevent PML or that can adequately treat PML if it occurs. It is not known whether early detection of PML and discontinuation of TYSABRI will mitigate the disease.
- PML has been reported following discontinuation of TYSABRI in patients who did not have findings suggestive of PML at the time of discontinuation. Patients should continue to be monitored for any new signs or symptoms that may be suggestive of PML for approximately six months following discontinuation of TYSABRI.
- In MS patients, an MRI scan should be obtained prior to initiating therapy with TYSABRI. This MRI may be helpful in differentiating subsequent multiple sclerosis symptoms from PML.
- Three sessions of plasma exchange over 5 to 8 days were shown to accelerate TYSABRI clearance in a study of 12 patients with MS who did not have PML, although in the majority of patients, alpha-4 integrin receptor binding remained high. Adverse events which may occur during plasma exchange include clearance of other medications and volume shifts, which have the potential to lead to hypotension or pulmonary edema. Although plasma exchange has not been studied in TYSABRI-treated patients with PML, it has been used in such patients in the postmarketing setting to remove TYSABRI more quickly from the circulation.
- Anti-JCV antibody testing should not be performed during or for at least two weeks following plasma exchange due to the removal of antibodies from the serum.
- Immune reconstitution inflammatory syndrome (IRIS) has been reported in the majority of TYSABRI treated patients who developed PML and subsequently discontinued TYSABRI. In almost all cases, IRIS occurred after plasma exchange was used to eliminate circulating TYSABRI. It presents as a clinical decline in the patient’s condition after TYSABRI removal (and in some cases after apparent clinical improvement) that may be rapid, can lead to serious neurological complications or death and is often associated with characteristic changes in the MRI. TYSABRI has not been associated with IRIS in patients discontinuing treatment with TYSABRI for reasons unrelated to PML. In TYSABRI-treated patients with PML, IRIS has been reported within days to several weeks after plasma exchange. Monitoring for development of IRIS and appropriate treatment of the associated inflammation should be undertaken.

Important Safety Information Continued on next page
Contraindications

➤ TYSABRI is contraindicated in patients who have or have had PML.

➤ TYSABRI is contraindicated in patients who have had a hypersensitivity reaction to TYSABRI.

TYSABRI TOUCH Prescribing Program

➤ TYSABRI is available only through a restricted program under a REMS called the TOUCH® Prescribing Program because of the risk of PML.

➤ For prescribers and patients, the TOUCH® Prescribing Program has two components: MS TOUCH® (for patients with multiple sclerosis) and CD TOUCH® (for patients with Crohn's disease).

➤ Prescribers must be certified and comply with the following:

   - Review the TOUCH Prescribing Program prescriber educational materials, including the full prescribing information.
   - Educate patients on the benefits and risks of treatment with TYSABRI, ensure that patients receive the Medication Guide, and encourage them to ask questions.
   - Review, complete, and sign the Patient-Prescriber Enrollment Form.
   - Evaluate patients three months after the first infusion, six months after the first infusion, every six months thereafter, and for at least six months after discontinuing TYSABRI.
   - Determine every six months whether patients should continue on treatment, and if so, authorize treatment for another six months.
   - Submit to Biogen Idec the “TYSABRI Patient Status Report and Reauthorization Questionnaire” six months after initiating treatment and every six months thereafter.
   - Complete an “Initial Discontinuation Questionnaire” when TYSABRI is discontinued and a “6-Month Discontinuation Questionnaire,” following discontinuation of TYSABRI.
   - Report cases of PML, hospitalizations due to opportunistic infections, and deaths to Biogen Idec at 1-800-456-2255 and to the Food and Drug Administration’s MedWatch Program at 1-800-FDA-1088 as soon as possible.

Important Safety Information Continued on next page
Patients must be enrolled in the TOUCH Prescribing Program, read the Medication Guide, understand the risks associated with TYSABRI, and complete and sign the Patient-Prescriber Enrollment Form.

Pharmacies and infusion centers must be specially certified to dispense or infuse TYSABRI.

**Herpes Encephalitis and Meningitis**

- TYSABRI increases the risk of developing encephalitis and meningitis caused by herpes simplex and varicella zoster viruses.
- Serious, life-threatening, and sometimes fatal cases have been reported in the postmarketing setting in multiple sclerosis patients receiving TYSABRI.
- Monitor patients receiving TYSABRI for signs and symptoms of meningitis and encephalitis. If herpes encephalitis or meningitis occurs, TYSABRI should be discontinued, and appropriate treatment for herpes encephalitis/meningitis should be administered.

**Hepatotoxicity**

- Clinically significant liver injury, including acute liver failure requiring transplant, has been reported in patients treated with TYSABRI in the postmarketing setting. In some patients, liver injury recurred upon rechallenge, providing evidence that TYSABRI caused the injury.
- Signs of liver injury, including markedly elevated serum hepatic enzymes and elevated total bilirubin, occurred as early as 6 days after the first dose; and signs of liver injury have also been reported for the first time after multiple doses.
- In some patients, liver injury recurred upon rechallenge, providing evidence that TYSABRI caused the injury.
- The combination of transaminase elevations and elevated bilirubin without evidence of obstruction is generally recognized as an important predictor of severe liver injury that may lead to death or the need for a liver transplant in some patients.
- TYSABRI should be discontinued in patients with jaundice or other evidence of significant liver injury (e.g., laboratory evidence).

**Hypersensitivity/Antibody Formation**

- Hypersensitivity reactions have occurred in patients receiving TYSABRI, including serious systemic reactions (e.g., anaphylaxis), which occurred at an incidence of <1%.

*Important Safety Information Continued on next page*
Reactions usually occur within 2 hours of the start of the infusion. Symptoms associated with these reactions can include urticaria, dizziness, fever, rash, rigors, pruritus, nausea, flushing, hypotension, dyspnea, and chest pain. Generally, these reactions are associated with antibodies to TYSABRI.

If a hypersensitivity reaction occurs, discontinue administration of TYSABRI, and initiate appropriate therapy. Patients who experience a hypersensitivity reaction should not be re-treated with TYSABRI.

Hypersensitivity reactions were more frequent in patients with antibodies to TYSABRI compared to patients who did not develop antibodies to TYSABRI in both MS and CD studies.

Patients who receive TYSABRI after an extended period without treatment may be at higher risk of hypersensitivity reactions.

**Immunosuppression/Infections**

- The immune system effects of TYSABRI may increase the risk for infections.
- In Study MS1, certain types of infections, including pneumonias and urinary tract infections (including serious cases), gastroenteritis, vaginal infections, tooth infections, tonsillitis, and herpes infections, occurred more often in TYSABRI-treated patients than in placebo-treated patients. One opportunistic infection, a cryptosporidial gastroenteritis with a prolonged course, was observed in a patient who received TYSABRI in Study MS1.
- In Studies MS1 and MS2, an increase in infections was seen in patients concurrently receiving short courses of corticosteroids. However, the increase in infections in TYSABRI-treated patients who received steroids was similar to the increase in placebo-treated patients who received steroids.
- Concurrent use of antineoplastic, immunosuppressant, or immunomodulating agents may further increase the risk of infections over the risk observed with use of TYSABRI alone. The safety and efficacy of TYSABRI in combination with antineoplastic, immunosuppressant, or immunomodulating agents have not been established.
- In Study MS1 and Study MS2, the rate of any type of infection was approximately 1.5 per patient-year in both TYSABRI-treated patients and placebo-treated patients.
- In Study MS1, the incidence of serious infections was approximately 3% in TYSABRI-treated patients and in placebo-treated patients. Most patients did not interrupt treatment with TYSABRI during infections.
- In postmarketing experience, serious herpes infections have occurred.

**Important Safety Information Continued on next page**
Laboratory Test Abnormalities

In clinical trials, TYSABRI was observed to induce increases in circulating lymphocytes, monocytes, eosinophils, basophils, and nucleated red blood cells. Observed changes persisted during TYSABRI exposure, but were reversible, returning to baseline levels usually within 16 weeks after the last dose. Elevations of neutrophils were not observed. TYSABRI induces mild decreases in hemoglobin levels that are frequently transient.

Adverse Reactions

The most common adverse reactions reported at an incidence of ≥10% with TYSABRI and ≥2% difference with placebo were headache (38% vs 33%), fatigue (27% vs 21%), infusion reactions (24% vs 18%), urinary tract infections (21% vs 17%), arthralgia (19% vs 14%), depression (19% vs 16%), pain in extremity (16% vs 14%), rash (12% vs 9%), gastroenteritis (11% vs 9%), and vaginitis* (10% vs 6%).

*Percentage based on female patients only.

The most frequently reported serious adverse reactions in Study MS1 were infections (3.2% vs 2.6% placebo), including urinary tract infection (0.8% versus 0.3%) and pneumonia (0.6% versus 0%), acute hypersensitivity reactions (1.1% vs 0.3%, including anaphylaxis/anaphylactoid reaction [0.8% versus 0%]), depression (1.0% vs 1.0%, including suicidal ideation or attempt [0.6% versus 0.3%]), and cholelithiasis (1.0% vs 0.3%).

Based on animal data, TYSABRI may cause fetal harm. TYSABRI should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus.
References:


Please see accompanying full Prescribing Information, including Boxed Warning.
Prescriber/Patient Enrollment Form—Crohn’s Disease

Completion of all pages is required.

**Patient Information**

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**Patient Authorization to Use/ disclosed Health Information**

I understand that I have certain rights related to the collection, use, and disclosure of my medical and health information. This information is called “protected health information” (PHI) and includes demographic information (such as sex, race, date of birth, etc.), the results of physical examinations, clinical tests, blood tests, X-rays, and other diagnostic and medical procedures that may be included in my medical records. Biogen Idec will not use my PHI without my consent.

This Authorization form applies to PHI created or obtained by my prescriber, my infusion site, my pharmacy, and my health insurance company. I understand that by signing this Authorization, I authorize my prescriber, infusion site, pharmacy, and/or health insurance company to disclose the PHI in my medical records to Biogen Idec Inc. and its representatives or agents, including information related to my medical condition, treatment, and health insurance, as well as all information provided on any prescription. I also authorize Biogen Idec to use this information to provide TYSABRI support services, such as investigating insurance coverage for TYSABRI and coordinating delivery of TYSABRI to the prescriber or infusion site administering TYSABRI (which may include forwarding my health information to a pharmacy).

I agree to allow Biogen Idec to ask me about and provide me with these support services, educational kits, and other information related to TYSABRI and/or my medical condition. I understand that, once my PHI has been disclosed to Biogen Idec, federal privacy laws may no longer protect the information. However, Biogen Idec agrees to protect my PHI by using it only for the purposes authorized in this Authorization or as required by law.

I understand that I may refuse to sign this Authorization, and refusing to do so will affect my eligibility to receive these additional services but will not affect my ability to receive TYSABRI. I understand that signing this Authorization will not change how my healthcare providers, health insurance plan, and pharmacies provide my medical treatment or payment for treatment or insurance benefits.

I understand that I may cancel all or a part of this Authorization at any time by mailing a letter requesting such cancellation to TYSABRI Support Services, 5000 Davis Drive, PO Box 13919, Research Triangle Park, NC 27709. If I cancel this Authorization, Biogen Idec will end further use and disclosure of my PHI as soon as possible. This will not affect health information that has already been used or disclosed in reliance upon this Authorization.

I will receive a copy of this signed Authorization. This Authorization expires ten (10) years from the date this Authorization is signed.

**Patient signature** (or personal representative): __________________________Date: __________________________

Reference ID: 3753128
Biogen Idec considers patient safety a priority. Read each section below and initial in the space provided if you understand the information. Do not sign this form if there is anything you do not understand about all the information you have received. Ask your doctor about anything you do not understand before you initial and sign this form.

**understand that** **ABRI** is a medicine approved to treat patients with moderate to severe Crohn’s disease who have not been helped enough by, or cannot tolerate, usual Crohn’s disease medicines and medicines called tumor necrosis factor (TNF) inhibitors.

- I have talked to my doctor and understand the benefits and risks of TYSABRI treatment
- I understand that I should not take certain medicines that weaken the immune system while I am taking TYSABRI

**understand that** **ABRI** increases my chance of getting a rare brain infection that usually leads to death or severe disability.

- This infection is called progressive multifocal leukoencephalopathy (PML). PML usually happens in people with weakened immune systems
- There is no known treatment, prevention, or cure for PML
- My chance for getting PML may be higher if I am also being treated with other medicines that can weaken my immune system, including other Crohn’s disease treatments. Even if I use TYSABRI alone to treat my CD, I can still get PML
- My chance for getting PML increases if I:
  - Have received TYSABRI for a long time, especially longer than 2 years
  - Have received certain medicines that can weaken my immune system before I start receiving TYSABRI
  - Have been exposed to John Cunningham Virus (JCV). JCV is a common virus that is harmless in most people but can cause PML in people who have weakened immune systems, such as people taking TYSABRI. Most people who are exposed to JCV do not know it or have any symptoms. This exposure usually happens in childhood. My doctor may do a blood test to check if I have been exposed to JCV before I start receiving TYSABRI or during my treatment. My risk of getting PML is greatest if I have all 3 risk factors listed above. There may be other risk factors for getting PML during TYSABRI treatment that we do not know about yet. My doctor should discuss the risks and benefits of TYSABRI treatment with me before I decide to receive TYSABRI
- I should call my doctor right away if I get any new or worsening symptoms that last several days, especially nervous system symptoms, while I am taking TYSABRI and for at least 6 months after I stop taking TYSABRI. Some of these symptoms include a new or sudden change in my thinking, eyesight, balance, or strength, but I should also report other new or worsening symptoms

**to receive** **ABRI**, all patients must be enrolled in a restricted program called the TOUCH Prescribing Program.

- The TOUCH Prescribing Program is run by the company that makes TYSABRI. Under this program, the company is required to collect information about my health at regular time periods. cannot receive ABRI if do not agree to follow the requirements of the TOUCH Prescribing Program. I understand that the TOUCH Prescribing Program does not require me to sign the Authorization included on page 1 of this form
- The company may use my information to meet the requirements of the TOUCH Prescribing Program, including helping me locate an authorized infusion site
- I must notify the TOUCH Prescribing Program if I switch physicians or infusion sites
- I have received, read, and understand the Patient Medication Guide
- I will bring to each TYSABRI infusion a list of all medicines and treatments that I have taken during the last month

**Patient Acknowledgment**

**Patient name:** ____________________________ **Date of birth:** __________ / __________ / ________ **(MM/DD/YYYY)**

**Patient signature** (or personal representative): ____________________________ **Date:** __________

Authority of personal representative (if applicable): ____________________________

Reference ID: 3753128
Patient History

Patient name: ___________________________ M ___________________________ Last ___________________________ DOB: ______/_____/______ (MM/DD/YYYY)

Date of first Crohn’s disease symptoms: ___________________________ (MM/YYYY)

Please indicate the patient’s Crohn’s disease therapy(ies) within the past one year AND whether the therapy is ongoing or stopped. Ongoing therapies, except corticosteroids, must be stopped before starting ABRI. (If patient was on multiple therapies, check all that apply.)

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<td>None</td>
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<td>Remicade</td>
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<td>Other immunomodulatory therapy or immunosuppressant therapy (not containing immunomodulators)</td>
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Has the patient had a surgery for Crohn’s disease within the previous year? Yes [ ] No [ ]

Has the patient ever received TYSABRI before? Yes [ ] No [ ]

Has the patient ever been prescribed an immunosuppressant or an antineoplastic therapy for any condition? Yes [ ] No [ ]

If yes, please check all of the following that apply:
- Remicade [ ] Humira [ ] Azathioprine or Mercaptopurine or Thioguanine [ ] Czm a [ ] Methotrexate [ ]
- Systemic steroids [ ] Vedolizumab (Entyvio™) [ ] Other [ ]

Has the patient ever been tested for the presence of anti-JCV antibodies? Yes [ ] No [ ] Unknown [ ]

If yes, has the patient ever tested for the presence of anti-JCV antibodies? Yes [ ] No [ ] Pending [ ]

Prescription for TYSABRI

Dose: ABRI® (natalizumab) 300 mg
Dispense: 1 vial efills: 12
Directions: infusion per rescribing Information every 4 weeks

I authorize Biogen Idec as my designated agent and on behalf of my patient to (1) use the information on this form to enroll the above-named patient in the TOUCH Prescribing Program, (2) furnish any information on this form to the insurer of the above-named patient, (3) forward the information on this form to the prescriber or infusion site administering TYSABRI, if applicable, (4) forward the above prescription by fax or by another mode of delivery to a pharmacy, if applicable, and (5) coordinate delivery of TYSABRI on behalf of the above-named patient.

Resciver signature (stamps not acceptable):

Date:

Prescriber

Prescriber name: ___________________________ M ___________________________ Last ___________________________

Office contact

Tax D #

DEA #

NP /UP N/prov der D # w th pat ent’s nsurer(s)

Continued on next page
I have read and understand the full Prescribing Information for TYSABRI.

I understand that TYSABRI is indicated for adult patients with moderately to severely active Crohn’s disease with evidence of inflammation who have had an inadequate response to, or are unable to tolerate, conventional Crohn’s disease therapies and inhibitors of TNF-α.

I understand that patients receiving TYSABRI should not take concomitant immunosuppressants (e.g., 6-mercaptopurine, azathioprine, cyclosporine, or methotrexate) or inhibitors of TNF-α.

I understand that this patient has moderately to severely active Crohn’s disease with evidence of inflammation.

I understand that TYSABRI increases the risk of progressive multifocal leukoencephalopathy (PML), an opportunistic viral infection of the brain that usually leads to death or severe disability. When initiating and continuing treatment with TYSABRI, I should consider whether the expected benefit of TYSABRI is sufficient to offset this risk.

I am aware that cases of PML have been reported in patients taking TYSABRI who were recently or concomitantly treated with immunomodulators or immunosuppressants, as well as in patients receiving TYSABRI monotherapy.

I understand that three risk factors identified thus far that increase the risk of PML in TYSABRI-treated patients are:
- Longer treatment duration, especially beyond 2 years
- Prior treatment with an immunosuppressant (e.g., mitoxantrone, azathioprine, methotrexate, cyclophosphamide, mycophenolate mofetil)
- The presence of anti-JCV antibodies

These factors should be considered in the context of expected benefit when initiating and continuing treatment with TYSABRI.

I have discussed other Crohn’s disease treatments with this patient.

To my knowledge, this patient has no known contraindications to TYSABRI treatment, including PML.

I have instructed this patient to promptly report to me any continuously worsening symptoms that persist over several days, especially nervous system symptoms.

I understand that this patient should be seen and evaluated 3 months after the first infusion, 6 months after the first infusion, every 6 months thereafter, and for at least 6 months after TYSABRI has been discontinued.

I understand that TYSABRI should be discontinued if the patient has not experienced a therapeutic benefit by 12 weeks of therapy.

I will determine every 6 months whether this patient should continue on TYSABRI and if so, authorize treatment for another 6 months. I understand that I am required to submit an “Initial Discontinuation Questionnaire” when TYSABRI is discontinued and a “6-Month Discontinuation Questionnaire” following discontinuation of TYSABRI.

I understand that patients receiving steroid therapy at the time of TYSABRI initiation must undergo a steroid-tapering regimen once a therapeutic response is achieved. If the patient with Crohn’s disease cannot be tapered off steroids within 6 months of starting TYSABRI, TYSABRI should be discontinued.

I should report, as soon as possible, cases of PML, hospitalizations due to opportunistic infection, and any death to Biogen Idec.

I understand that data concerning this patient and me will be entered into the mandatory TOUCH Prescribing Program. Biogen Idec requires my cooperation with periodic data collection. Failure to provide the requested information or otherwise comply with the requirements of the TOUCH Prescribing Program may result in discontinuation of TYSABRI treatment for this patient and termination of my authorization to prescribe TYSABRI.

I have received educational materials regarding the benefits and risks of TYSABRI treatment.

I have, or another healthcare provider under my direction has, educated this patient on the benefits and risks of treatment with TYSABRI, provided him or her with the Patient Medication Guide and Enrollment Form, instructed him or her to read these materials, and encouraged him or her to ask questions when considering TYSABRI.

Patient name: ___________________________  Date of birth: ___________________________

First M Last

Prescriber signature: ___________________________  Date: ___________________________

Reference ID: 3753128
Please submit this form to:
Biogen Idec
www.touchprogram.com
Fax: 1 800 840 1278

Prescriber/Patient Enrollment Form—Crohn’s Disease
Completion of all pages is required.

Patient Information

Date of birth / / (MM/DD/YYYY)
Patient name First M Last

In addition, I allow the sharing of my health information to the person or people I name below. Biogen Idec may contact the people named below to discuss my enrollment in the TOUCH Program.

Designated Individual (print name):
Relationship:

Infusion Site Information*

1. Prescriber will administer ABRI and request the following services (check only one):
   - No services required
   - Forward this prescription to a specialty pharmacy provider to investigate pharmacy coverage and coordinate delivery to prescriber’s office
   - Please conduct insurance research and procurement options for TYSABRI

OR

2. Prescriber will refer AB treatment to another site (check only one):
   - I require assistance in locating an infusion site
   - I am referring the patient to the following infusion site or healthcare provider

   Name of infusion site or healthcare provider (first last)
   Street address or street number
   City State Zip

   Office contact
   Telephone ____________-________-________-________
   Fax ____________-________-________-________

   *Note: TYSABRI can only be infused at authorized infusion sites. Biogen Idec will contact you if the infusion site you have indicated is not authorized to infuse TYSABRI.

Please see accompanying full prescribing information, including Boxed Warning, for important safety information.

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Reference ID: 3753128
Please submit this form to:
Biogen Idec
www.touchprogram.com
TYSABRI
Patient Status Report and Reauthorization Questionnaire—Crohn’s Disease

Re: <Patient Name>
Patient Enrollment Number: <Patient TOUCH ID>
Patient date of birth: <DOB>
Authorization expiration date: <MM/DD/YYY>

Dear <MD Name>,

Our records indicate that <Patient Name>'s authorization to receive TYSABRI will expire on <MM/DD/YY YY> and he/she will no longer be able to receive TYSABRI. Please submit the completed form to Biogen Idec via TOUCH On-Line (www.touchprogram.com) OR fax (1-800-840-1278) and place a copy in the patient’s record.

Is the patient still under <MD name>’s care?
☐ Yes ☐ No/ I don’t know

If No, please provide name and phone number for new prescriber, if available __________________________

Is the patient alive?
☐ Yes ☐ No

Since starting TYSABRI therapy has the patient been diagnosed with any of the following that you have not reported to Biogen Idec:

PROGRESSIVE MULTIFOCAL LEUKOENCEPHALOPATHY (PML)
☐ Yes ☐ No or ☐ Under investigation

OPPORTUNISTIC INFECTION* for which they have been hospitalized
☐ Yes ☐ No or ☐ Under investigation

MALIGNANCY
☐ Yes ☐ No or ☐ Under investigation

Since <last authorization>, has the patient been tested for the presence of anti-JCV antibodies?
☐ Yes ☐ Not performed

If performed, test result:
☐ Positive ☐ Negative ☐ Pending

Is the patient currently receiving or has the patient received systemic steroids for the treatment of Crohn's disease?
☐ Yes ☐ No

Within the past year, and since starting TYSABRI, has the patient received greater than 6 consecutive months of systemic steroids for the treatment of Crohn's disease?
☐ Yes ☐ No

Is the patient currently receiving or has the patient received any IMMUNOMODULATOR, or IMMUNOSUPPRESSANT THERAPIES, in the previous 6 months?
☐ Yes ☐ No

If Yes, please indicate the type of therapy and the number of months of use.

If the patient is still under <MD name>’s care DO YOU AUTHORIZE the continuation of TYSABRI treatment for the next 6 months for the patient?
☐ Yes ☐ No

If you answer No, Biogen Idec will contact the patient and the infusion site to STOP TYSABRI TREATMENT.

The patient will not be eligible to receive TYSABRI treatment, and you will receive a final questionnaire for this patient in 6 months.

*OPPORTUNISTIC INFECTION is defined as an infection due to an organism that generally does not cause disease, or causes only mild or self-limited disease in people with normally functioning immune systems, but causes more significant disease in people with impaired immunity. These infections are frequently severe, prolonged, or disseminated. Examples include esophageal candidiasis, systemic fungal infections, pneumonia, cryptococcal meningitis, mycobacterial infections (including pulmonary and extra-pulmonary tuberculosis), chronic intestinal cryptosporidiosis, and disseminated viral infections (such as disseminated herpes or cytomegalovirus infections).

Prescriber signature: __________________________________________ Date (MM/DD/YYYY): _________ /__________ / __________

(If applicable) Print TOUCH Authorized Prescriber Delegate Name: __________________________________________

Please Note: A TOUCH authorized physician may complete this form on behalf of the Prescriber of record. This questionnaire will be used consistent with the TOUCH Prescriber/Patient Enrollment Form signed by you and your patient and with HIPAA and applicable privacy rules. If you have questions, or if you need additional information, please call 1-800-456-2255 from 8:30 AM to 8:00 PM (ET).

Please see full Prescribing Information, including Boxed Warning, at www.TYSABRI.com

Reference ID: 3753128
Dear <Prescriber Name>,

Our records indicate that it has been 12 weeks since <Patient Name> received his or her first dose of TYSABRI. The Prescribing Information states that if a patient with Crohn’s disease has not experienced a therapeutic benefit by 12 weeks of induction of therapy she/he should be discontinued from TYSABRI treatment.

This questionnaire is necessary to fulfill the tracking requirements of the TOUCH Prescribing Program for Crohn’s disease patients treated with TYSABRI. You may also be contacted for additional information in response to answers provided on this form.

Submit the completed evaluation to Biogen Idec via TOUCH On-Line (www.touchprogram.com) OR fax (1-800-840-1278) and place one copy in the patient’s record.

Please answer Yes or No to the following questions:

1. Has this patient experienced a therapeutic benefit within 12 weeks after starting TYSABRI treatment?
   - Yes
   - No*  
   *TYSABRI should be discontinued if the patient has not experienced a therapeutic benefit by 12 weeks of induction therapy with TYSABRI.

2. Will the patient continue on TYSABRI?
   - Yes
   - No*  
   *If you answer No, Biogen Idec will contact the patient and the infusion site to STOP TYSABRI TREATMENT. The patient will not be eligible to receive TYSABRI treatment, and you will receive a discontinuation questionnaire to complete for this patient.

If you have questions, or if you need additional information, please call 1-800-456-2255 from 8:30 AM to 8:00 PM (ET).

Prescriber signature: __________________________________________ Date (MM/DD/YYYY): _____/_____/_____

(If applicable) Print TOUCH Authorized Prescriber Delegate Name: __________________________________________

Please Note: A TOUCH authorized physician may complete this form on behalf of the TOUCH Prescriber of record. This questionnaire will be used consistent with the TOUCH Prescriber/Patient Enrollment form signed by you and your patient with HIPAA and applicable privacy rules.

For full Prescribing Information, including Boxed Warning, please see www.TYSABRI.com.
TYSABRI Initial Discontinuation Questionnaire—Crohn’s Disease

Re: <Patient Name>
Patient Enrollment Number: <Patient TOUCH ID>
Patient date of birth: <DOB>

Dear <MD Name>,

Our records indicate that <Patient Name> received a final dose of TYSABRI on <MM/DD/YYYY>. You may receive this questionnaire twice, once at discontinuation and a second time 6 months later.

This Initial Discontinuation Questionnaire is necessary to fulfill the tracking requirements of the TOUCH Prescribing Program for Crohn’s disease patients treated with TYSABRI. You may also be contacted for additional information in response to answers provided on this form.

Submit the completed Initial Discontinuation Questionnaire to Biogen Idec via TOUCH On-Line (www.touchprogram.com) OR fax (1-800-840-1278) and place one copy in the patient’s record. This form is mandatory for all discontinued patients.

A Is the patient still under <MD name>’s care?

☐ Yes ☐ No/ I don’t know

If No, please provide name and phone number for new prescriber, if available ____________________________

B Is the patient alive?

☐ Yes ☐ No

Since starting TYSABRI therapy has the patient been diagnosed with any of the following that you have not reported to Biogen Idec:

C PROGRESSIVE MULTIFOCAL LEUKOENCEPHALOPATHY (PML)

☐ Yes ☐ No or ☐ Under investigation

D OPPORTUNISTIC INFECTION* for which they have been hospitalized

☐ Yes ☐ No or ☐ Under investigation

E MALIGNANCY

☐ Yes ☐ No or ☐ Under investigation

F Since <last authorization>, has the patient been tested for the presence of anti-JCV antibodies?

☐ Yes ☐ Not Performed

If performed, test result:

☐ Positive ☐ Negative ☐ Pending

G Since <MM/DD/YYYY> is the patient currently receiving or has the patient received systemic steroids for the treatment of a Crohn’s flare?

☐ Yes ☐ No

If Yes, please circle the number of months of use:

1 2 3 4 5 6

H Within the past year, and since starting TYSABRI, has the patient received greater than 6 consecutive months of systemic steroids for the treatment of Crohn’s disease?

☐ Yes ☐ No

I Since <MM/DD/YYYY> is the patient currently receiving or has the patient received any IMMUNOMODULATORY, or IMMUNOSUPPRESSANT THERAPIES?

☐ Yes ☐ No

If Yes, please indicate the type of therapy and the number of months of use.

<table>
<thead>
<tr>
<th>Therapy</th>
<th>Months of Use in Last 6 Months</th>
</tr>
</thead>
<tbody>
<tr>
<td>Remicade®</td>
<td>1 2 3 4 5 6</td>
</tr>
<tr>
<td>Humira®</td>
<td>1 2 3 4 5 6</td>
</tr>
<tr>
<td>Azathioprine or Mercaptopurine or Thioguanine</td>
<td>1 2 3 4 5 6</td>
</tr>
<tr>
<td>Vedolizumab (Entyvio™)</td>
<td>1 2 3 4 5 6</td>
</tr>
<tr>
<td>Methotrexate</td>
<td>1 2 3 4 5 6</td>
</tr>
<tr>
<td>Cimzia®</td>
<td>1 2 3 4 5 6</td>
</tr>
<tr>
<td>Other immunomodulatory or immunosuppressant therapy†</td>
<td>1 2 3 4 5 6</td>
</tr>
</tbody>
</table>

*OPPORTUNISTIC INFECTION is defined as an infection due to an organism that generally does not cause disease, or causes only mild or self-limited disease in people with normally functioning immune systems, but causes more significant disease in people with impaired immunity. These infections are frequently severe, prolonged, or disseminated. Examples include esophageal candidiasis, systemic fungal infections, pneumocystis carinii pneumonia, mycobacterial infections (including pulmonary and extra-pulmonary tuberculosis), chronic intestinal cryptosporidiosis, and disseminated viral infections (such as disseminated herpes or cytomegalovirus infections).

Prescriber signature: ____________________________ Date (MM/DD/YYYY): __________ / __________ / __________

(If applicable) Print TOUCH Authorized Prescriber Delegate Name: ____________________________

Please Note: A TOUCH authorized physician may complete this form on behalf of the Prescriber of record. This questionnaire will be used consistent with the TOUCH Prescriber/Patient Enrollment Form signed by you and your patient and with HIPAA and applicable privacy rules.

If you have questions, or if you need additional information, please call 1-800-456-2255 from 8:30 AM to 8:00 PM (ET).

Please see full Prescribing Information, including Boxed Warning, at www.TYSABRI.com

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All other trademarks are the marks of their respective owners.

Reference ID: 3753128
Dear <Prescriber Name>,

Our records indicate that <Patient Name> received a final dose of TYSABRI on <MM/DD/YYYY>.

This Discontinuation Questionnaire is necessary to fulfill the tracking requirements of the TOUCH Prescribing Program for Crohn’s disease patients treated with TYSABRI. You may also be contacted for additional information in response to answers provided on this form.

Submit the completed 6-Month Discontinuation Questionnaire to Biogen Idec via TOUCH On-Line (www.touchprogram.com) OR fax (1-800-840-1278) and place a copy in the patient’s record. This form is mandatory for all discontinued patients.

A Is the patient still under <Prescriber Name>’s care?

☐ Yes ☐ No/I don’t know

If No, please provide contact information for new prescriber, if available.
Name and phone of new prescriber: ____________________________________________________________

B Is the patient alive?

☐ Yes ☐ No

Since starting TYSABRI therapy has the patient been diagnosed with any of the following that you have not reported to Biogen Idec:

C PROGRESSIVE MULTIFOCAL LEUKOENCEPHALOPATHY (PML)

☐ Yes ☐ No or ☐ Under investigation

D OPPORTUNISTIC INFECTION* for which they have been hospitalized

☐ Yes ☐ No or ☐ Under investigation

E MALIGNANCY

☐ Yes ☐ No or ☐ Under investigation

*OPPORTUNISTIC INFECTION is defined as an infection due to an organism that generally does not cause disease, or causes only mild or self-limited disease in people with normally functioning immune systems, but causes more significant disease in people with impaired immunity. These infections are frequently severe, prolonged, or disseminated. Examples include esophageal candidiasis, systemic fungal infections, pneumocystis carinii pneumonia, mycobacterial infections (including pulmonary and extra-pulmonary tuberculosis), chronic intestinal cryptosporidiosis, and disseminated viral infections (such as disseminated herpes or cytomegalovirus infections).

Prescriber signature: ____________________________ Date (MM/DD/YYYY): _____ / _____ / ______

(If applicable) Print TOUCH Authorized Prescriber Delegate Name: ____________________________

Please Note: A TOUCH authorized physician may complete this form on behalf of the TOUCH Prescriber of record. This questionnaire will be used consistent with the TOUCH Prescriber/Patient Enrollment Form signed by you and your patient and with HIPAA and applicable privacy rules. If you have questions, or if you need additional information, please call 1-800-456-2255 from 8:30 AM to 8:00 PM (ET).

For full Prescribing Information, including Boxed Warning, please see www.TYSABRI.com.

Reference ID: 3753128

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# Change Prescriber Authorization

**PRESCRIBER AUTHORIZATION REQUESTED**

<table>
<thead>
<tr>
<th>Date:</th>
<th>&lt;Current_Date&gt;</th>
<th>Patient Enrollment Number:</th>
<th>&lt;Alt_ID&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>New Prescriber:</td>
<td>&lt;Phys_First_Name&gt; &lt;Phys_Last_Name&gt;</td>
<td>Patient Name:</td>
<td>&lt;Pat_First_Name&gt; &lt;Pat_Last_Name&gt;</td>
</tr>
<tr>
<td></td>
<td>&lt;MD_Address&gt;</td>
<td>Patient DOB:</td>
<td>&lt;PatDOB&gt;</td>
</tr>
<tr>
<td></td>
<td>&lt;MD_City&gt;, &lt;MD_State&gt; &lt;MD_Zip&gt;</td>
<td>Patient Enrollment Period:</td>
<td>Pat_Auth_Begin&gt; through &lt;Pat_Auth_End&gt;</td>
</tr>
<tr>
<td>Phone:</td>
<td>&lt;MD_Phone&gt;</td>
<td>Infusion Site:</td>
<td></td>
</tr>
<tr>
<td>Fax:</td>
<td>&lt;MD_Fax&gt;</td>
<td>Infusion Site Address</td>
<td></td>
</tr>
<tr>
<td>Prescriber DEA:</td>
<td></td>
<td>Prescriber State License Number:</td>
<td></td>
</tr>
</tbody>
</table>

Our records indicate that <Pat_First_Name> <Pat_Last_Name> will continue his/her TYSABRI (natalizumab) therapy under your care. If you agree to accept this patient, please sign this form and fax it to Biogen Idec at 1-800-840-1278.

If you do not accept this patient or have questions about the TOUCH Prescribing Program Requirements, please call the TOUCH Prescribing Program at 1-800-456-2255. We are available Monday through Friday, 8:30 AM to 8:00 PM (ET).

*I accept <Pat_First_Name> <Pat_Last_Name> under my care for TYSABRI (natalizumab) treatment.*

**Prescription for TYSABRI**

- **Dose:** TYSABRI® (natalizumab) 300 mg
- **Dispense:** 1 vial
- **Refills:** 12
- **Directions:** IV infusion per Prescribing Information every 4 weeks

I authorize Biogen Idec as my designated agent and on behalf of my patient to (1) use the information on this form to continue the enrollment of the above-named patient in the TOUCH Prescribing Program, (2) forward the prescription by fax or by another mode of delivery to a pharmacy, if applicable, and (3) coordinate delivery of TYSABRI on behalf of the above named patient.

---

**Prescriber Signature**

**Date**

**FAX this signed form to 1-800-840-1278**

For full Prescribing Information including Boxed Warning, please see www.TYSABRI.com
The TOUCH Prescribing Program was developed as part of the Biogen Idec commitment to patient safety. Only authorized infusion sites may receive shipments of and infuse TYSABRI. An infusion site may become authorized only after it has taken part in compulsory training conducted by Biogen Idec and faxed a completed Enrollment Form to Biogen Idec. Upon receipt of this Enrollment Form, Biogen Idec will fax and mail an authorization confirmation letter to provide your Site Authorization Number and confirm your Shipping Address.

**Infusion Site Address (address where patient is infused)**

Name of Infusion Site: 
Address 1: 
Address 2: 
City: State: ZIP: 
Contact name: 
Telephone: Fax: 

**Method of acquiring TYSABRI**

Infusion site will acquire TYSABRI directly. If YES, check all that apply: 
- Buy/Bill
- Assignment of Benefits/Specialty Pharmacy

Infusion site will acquire through a certified pharmacy.*

*A certified pharmacy is located within a hospital, group practice, or infusion site and is associated with an infusion site. Retail pharmacies and wholesalers are excluded from holding inventory and dispensing TYSABRI.

**Shipping Address (address to which drug will be shipped)**

Check here if address is same as above. Please note that this is the ONLY address to which TYSABRI will be shipped.

Name of Infusion Site or Certified Pharmacy: 
Address 1: 
Address 2: 
City: State: ZIP: 
Contact name: 
Telephone: Fax: 

**Infusion Site Acknowledgment**

- The infusion site has received training and educational materials on the TOUCH Prescribing Program for use in the following indication(s): MS ☐ CD ☐ (check all that apply)
- I understand that TYSABRI will be administered only to patients who are currently authorized in the TOUCH Prescribing Program. Patient authorization must be confirmed prior to each infusion by:
  - For TOUCH On-Line infusion sites: Patient Authorization Status must be “Authorized” or
  - For paper-based infusion sites: Receipt of current Notice of Patient Authorization and verification that no Notice of Patient Discontinuation is on file
- I understand that each patient will receive a copy of the TYSABRI Patient Medication Guide prior to each infusion
- I understand that a TYSABRI Pre-infusion Patient Checklist must be completed prior to each infusion. The Pre-infusion Patient Checklist must be submitted to Biogen Idec within 1 business day of the patient visit regardless of whether or not the patient received the infusion by:
  - For paper-based infusion sites: sending a copy of the completed Pre-infusion Patient Checklist to Biogen Idec. A copy must also be placed in the patient’s medical record
  - For TOUCH On-Line infusion sites: the infusion nurse can read, complete and submit the Pre-Infusion Patient Checklist directly in TOUCH On-Line
- I understand that, per the requirements of the TOUCH Prescribing Program, this infusion site’s compliance may be reviewed by the Food and Drug Administration (FDA), and/or audited by Biogen Idec and/or a third party designated by Biogen Idec
- I understand that noncompliance with the requirements of the TOUCH Prescribing Program will result in de-enrollment of the infusion site and termination of the authorization to infuse TYSABRI

**Responsible party acknowledgment:**

Name: __________________________ Title: __________________________ Date: __________________________

Please see full Prescribing Information, including Boxed Warning, at www.TYSABRI.com

Reference ID: 3753128

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3-7237-xx
### TYSABRI

**Pre-infusion Patient Checklist**

**Patient name:**

First  MI  Last

**Patient Enrollment Number:** (Issued by Biogen Idec. Call 1-800-456-2255 or log on to www.touchprogram.com if number is not on file.)

**Site name:**

**Site Authorization Number:**

As a condition of your site's authorization to infuse TYSABRI® (natalizumab), this Pre-infusion Patient Checklist must be completed for each patient prior to each infusion. This page must be submitted on-line (www.touchprogram.com) OR faxed to Biogen Idec (1-800-840-1278) within 1 day of the patient’s visit and a copy retained in the patient’s record whether the patient has been infused or not.

**STEP 1:** Ensure that the patient is currently authorized to receive TYSABRI for MS or Crohn’s disease.

You must refer to the patient’s record prior to every infusion.

- If the patient did not receive his or her previous infusion, and physician clearance was required, you must confirm authorization from the prescriber before providing the current infusion
- Confirm the patient status is listed as “Authorized” on TOUCH On-Line (www.touchprogram.com) OR
- Confirm that there is a current Notice of Patient Authorization on file and that you have not received a Notice of Patient Discontinuation (paper-based process)

**Is the patient currently authorized to receive TYSABRI?**

<table>
<thead>
<tr>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
</table>

**Yes** Continue to next question.

**No** STOP—DO NOT INFUSE. If authorization cannot be verified on-line at www.touchprogram.com OR by calling 1-800-456-2255, the patient must be referred back to the healthcare provider who prescribed TYSABRI.

**STEP 2:** Confirm that the patient has read and understood the Patient Medication Guide.

The patient must read the Patient Medication Guide prior to every infusion. **Has the patient received and read the Patient Medication Guide, including the section “What should I tell my doctor and nurse before each infusion of TYSABRI?”**

<table>
<thead>
<tr>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
</table>

**Yes** Continue to next question.

**No** STOP—provide the Patient Medication Guide. Proceed to the next question after the patient has read it.

**STEP 3:** Read aloud and mark “Yes” or “No” for the patient’s answers to the following questions:

1. Over the past month, have you had any new or worsening medical problems (such as a new or sudden change in your thinking, eyesight, balance, strength, or other problems) that have persisted over several days? **Yes** **No**

2. Do you have a medical condition that can weaken your immune system, such as HIV infection or AIDS, leukemia or lymphoma, or an organ transplant, that may suggest that your body is not able to fight infections well? **Yes** **No**

3. **Crohn’s disease ONLY**
   - In the past month have you taken, or are currently taking, any medicines to treat cancer or MS or any other medicines that may suggest that your body is not able to fight infections well? **Yes** **No**

3. **MS ONLY**
   - In the past month, have you taken medicines to treat cancer or MS or any other medicines that may suggest that your body is not able to fight infections well? **Yes** **No**

**STEP 4:** Record infusion information.

If the patient answered YES to question 1, 2 or 3, **DO NOT INFUSE.** Contact the healthcare provider who prescribed TYSABRI and review the patient’s answers.

- After discussing the patient’s answers, did the prescriber authorize the patient to be infused?

- Check here if you were unable to contact the prescriber. (See next page for further instructions.)

<table>
<thead>
<tr>
<th>Date infused (MM/DD/YYYY):</th>
<th>Not infused</th>
</tr>
</thead>
</table>

If the next infusion has been scheduled, please enter date (MM/DD/YYYY):

**Name and signature of staff completing checklist:**

<table>
<thead>
<tr>
<th>Date</th>
</tr>
</thead>
</table>

**STEP 5:** Submit the Pre-infusion Patient Checklist to Biogen Idec on-line at www.touchprogram.com or fax to 1-800-840-1278.

Reference ID: 3753128
### Examples of Immunosuppressants, Antineoplastics, and Immunomodulators

#### Multiple Sclerosis

<table>
<thead>
<tr>
<th>Approved MS Therapies:</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>D methy umerate (EC DERA®)</td>
<td></td>
</tr>
<tr>
<td>G at ramer acetate (Copaxone®)</td>
<td></td>
</tr>
<tr>
<td>nterferon beta 1a (Reb F® AVONEX®)</td>
<td></td>
</tr>
<tr>
<td>nterferon beta 1b (Betaseron®, Extav a®)</td>
<td></td>
</tr>
<tr>
<td>ngo mod (G enyA®)</td>
<td></td>
</tr>
<tr>
<td>M toxicantrone (Novantrone®)</td>
<td></td>
</tr>
<tr>
<td>Peg nterferon beta 1a (PLEGR Dy®)</td>
<td></td>
</tr>
<tr>
<td>A emtuzumab (LEM RADAm)</td>
<td></td>
</tr>
<tr>
<td>er f unom de (Aubag o®)</td>
<td></td>
</tr>
</tbody>
</table>

#### Crohn’s Disease

<table>
<thead>
<tr>
<th>Approved TNF-α inhibitors for Crohn’s disease:</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Inf x mab (Rem cade®)</td>
<td></td>
</tr>
<tr>
<td>Ada mubab (Hum ra®)</td>
<td></td>
</tr>
</tbody>
</table>

#### Immunosuppressants/Antineoplastics:

- Azath opr ne (muram®, Azasan®)  
- Cadr b ne (Leustat n®)  
- Cyc ophospham de (Cytoxan®, Neosar®)  
- Cyc ospre ne (Sand mmune§, Neora®)  
- udarab ne phosphat e (udara®)  
- Lef unom de (Arava®)  
- Mercaptopur ne (Pur netho §)  
- Methotrexate (Methotrex®, Rheumatrex®, rexa ®)  
- Mycopheno ate mofet (Ce Cept®)  
- Pemtrexed (A mta®)  
- Additional Immumodulators and Immunosupressants: |  |
- Other nterferons (Act mmune®, Infergen®, ntron® A)  
- Pegasys®, PEG ntron® Rebetrone®, Roferon® A)  
- Ada mubab (um ra®)  
- Efacept (Amv®)  
- A emtuzumab (Camaph®)  
- Anak nra (K neret®)  
- Dac zqumab (Zenapax®)  
- Efa zqumab (Rapt va®)  
- Etanercept (Enbre ®)  
- G at ramer acetate (Copaxone®)  
- Intravenous mmunog obu n (IVG)  
- M toxicantrone (Novantrone®)  
- Other nteferons (Act mmune®, Infergen®, Intron® A, Pegasys®, PEG-Intron® , Rebetrone®, Roferon® A)  
- Rux mab (R tuxan®)  
- Trastuzumab (Hercept n®)  
- Vedo zqumab (Enlyv®)  

This list does not include all drugs that can suppress the immune system.

- Patients should consult their prescribing physician regarding drugs that may be taken concurrently with TYSABRI.
  - If there are any questions regarding concurrent therapy, **do not infuse** at this time and consult the healthcare provider who prescribed TYSABRI.

**If you are unable to contact the prescriber:**

Instruct the patient to contact his/her prescriber and to reschedule an infusion as soon as possible. Continue efforts to reach the prescriber to inform him/her of the reason(s) for not infusing this patient. You will need to confirm authorization from the prescriber on the subsequent infusion.

This Pre-infusion Patient Checklist is not intended to replace the infusion site’s general infusion protocol(s). Nor is this Pre-infusion Patient Checklist intended to be a substitute for consultation and review of reference materials and medical literature pertaining to individual clinical circumstances. Healthcare providers should make all treatment decisions based on the context of the situation and their clinical judgment.

Please do not make any extraneous marks on the Pre-infusion Patient Checklist. If there is information that you would like to share with Biogen Idec and the TOUCH Prescribing Program, please contact us at 1-800-456-2255.

Please see accompanying full Prescribing Information, including Boxed Warning, for important safety information.
The TOUCH Prescribing Program was developed as part of the Biogen Idec commitment to patient safety. Only authorized certified pharmacies may dispense to authorized infusion sites. A certified pharmacy may become authorized after it has taken part in compulsory training conducted by Biogen Idec and faxed a completed Enrollment Form to Biogen Idec. Upon receipt of this Enrollment Form, Biogen Idec will fax and mail an Authorization Confirmation Letter to provide your Pharmacy Authorization Number and confirm your Shipping Address. This letter will also provide you with the Site Authorization Numbers of any of your associated infusion sites that have been authorized to infuse TYSABRI.

*A certified pharmacy is located within a hospital, group practice, or infusion site and is associated with an infusion site. Retail pharmacies and wholesalers are excluded from holding inventory and dispensing TYSABRI.

Certified Pharmacy Shipping Address

Please note that this is the ONLY address to which TYSABRI will be shipped.

Name of Certified Pharmacy

Address 1

Address 2

City State ZIP

Contact name

NCPDP

Title/position

Telephone

Fax

Associated Infusion Site Name

Please list all potential infusion sites that your pharmacy supports. If you need additional space, please attach a separate page.

1

2

3

4

Name of Infusion Site

Contact name

Address

City State ZIP

Name of Infusion Site

Contact name

Address

City State ZIP

Name of Infusion Site

Contact name

Address

City State ZIP

Name of Infusion Site

Contact name

Address

City State ZIP

I confirm that the above information is correct. I understand that by signing this form this pharmacy agrees to dispense TYSABRI only to associated infusion sites that have been authorized according to the TOUCH Prescribing Program.

Certified Pharmacy Acknowledgment

The pharmacy has received training and educational materials on the TOUCH Prescribing Program for use in the following indication(s): MS ☐ CD ☐ (check all that apply)

Certified pharmacies may dispense TYSABRI only to authorized infusion sites

➢ I understand that, per the requirements of the TOUCH Prescribing Program, this certified pharmacy’s compliance may be reviewed by the Food and Drug Administration (FDA), and/or audited by Biogen Idec, and/or a third party designated by Biogen Idec.

➢ I understand that noncompliance with the requirements of the TOUCH Prescribing Program may result in my pharmacy no longer being enrolled and termination of our participation in the program

Responsible party acknowledgment: ___________________________________________ Date: ________________________

Name: ___________________________________________ Title: ___________________________________________

Please see accompanying full Prescribing Information.
Please see full Prescribing Information, including Boxed Warning, for important safety information.
Objectives

➢ Provide an overview of important safety information

➢ Provide an overview of the TOUCH Prescribing Program for Multiple Sclerosis (MS) and Crohn’s disease (CD)

➢ Review the process steps to complete TOUCH Prescribing Program components including use of TOUCH On-Line

➢ Review specific MS TOUCH and/or CD TOUCH Prescribing Program materials

➢ Review the responsibilities of each participant in the TOUCH Prescribing Program
TYSABRI® (natalizumab) is indicated as monotherapy for the treatment of patients with relapsing forms of multiple sclerosis.

TYSABRI increases the risk of PML.

When initiating and continuing treatment with TYSABRI, physicians should consider whether the expected benefit of TYSABRI is sufficient to offset this risk.

See Full Prescribing Information regarding the risk of PML with TYSABRI.
TYSABRI® is indicated for inducing and maintaining clinical response and remission in adult patients with moderately to severely active Crohn’s disease with evidence of inflammation who have had an inadequate response to, or are unable to tolerate, conventional CD therapies and inhibitors of TNF-α.

TYSABRI should not be used in combination with immunosuppressants (e.g., 6-mercaptopurine, azathioprine, cyclosporine, or methotrexate) or inhibitors of TNF-α.
TYSABRI® increases the risk of progressive multifocal leukoencephalopathy (PML), an opportunistic viral infection of the brain that usually leads to death or severe disability.

Risk factors for the development of PML include duration of therapy, prior use of immunosuppressants, and presence of anti-JCV antibodies. These factors should be considered in the context of expected benefit when initiating and continuing treatment with TYSABRI.

Healthcare professionals should monitor patients on TYSABRI®, and for six months following discontinuation of TYSABRI, for any new sign or symptom that may be suggestive of PML.

TYSABRI dosing should be withheld immediately at the first sign or symptom that may be suggestive of PML.
Contraindications

➢ TYSABRI is contraindicated in patients who have or have had PML.

➢ TYSABRI is contraindicated in patients who have had a hypersensitivity reaction to TYSABRI.
Warnings and Precautions - PML

Three factors that are known to increase the risk of PML in TYSABRI-treated patients have been identified:

- Longer treatment duration, especially beyond 2 years. There is limited experience in patients who have received more than 6 years of TYSABRI treatment
- Prior treatment with an immunosuppressant (e.g., mitoxantrone, azathioprine, methotrexate, cyclophosphamide, mycophenolate mofetil)
- The presence of anti-JCV antibodies. Patients who are anti-JCV antibody positive have a higher risk for developing PML

These factors should be considered in the context of expected benefit when initiating and continuing treatment with TYSABRI.
Infection by the JC virus is required for the development of PML.

Anti-JCV antibody testing should not be used to diagnose PML.

Anti-JCV antibody negative status indicates that exposure to the JC virus has not been detected.

Patients who are anti-JCV antibody negative have a lower risk of PML than those who are positive. Patients who are anti-JCV antibody negative are still at risk for the development of PML due to the potential for a new JCV infection, or a false negative test result.
The reported rate of seroconversion in patients with MS (changing from anti-JCV antibody negative to positive) is 3 to 8 percent annually. In addition, some patients’ serostatus may change intermittently. Therefore, patients with a negative anti-JCV antibody test result should be retested periodically.

For purposes of risk assessment, a patient with a positive anti-JCV antibody test at any time is considered anti-JCV antibody positive regardless of the results of any prior or subsequent anti-JCV antibody testing. When assessed, anti-JCV antibody status should be determined using an analytically and clinically validated immunoassay.

Anti-JCV antibody testing should not be performed for at least two weeks following plasma exchange due to the removal of antibodies from the serum.
Warnings and Precautions – Herpes Encephalitis and Meningitis

➤ TYSABRI increases the risk of developing encephalitis and meningitis caused by herpes simplex and varicella zoster viruses.

➤ Serious, life-threatening, and sometimes fatal cases have been reported in the postmarketing setting in multiple sclerosis patients receiving TYSABRI.

➤ Monitor patients receiving TYSABRI for signs and symptoms of meningitis and encephalitis. If herpes encephalitis or meningitis occurs, TYSBARI should be discontinued, and appropriate treatment for herpes encephalitis/meningitis should be administered.
Warnings and Precautions – Hepatotoxicity

- Clinically significant liver injury, including acute liver failure requiring transplant, has been reported in patients treated with TYSABRI® in a postmarketing setting.

- Signs of liver injury, including markedly elevated serum hepatic enzymes and elevated total bilirubin, occurred as early as 6 days after the first dose; and signs of liver injury have also been reported for the first time after multiple doses.

- In some patients, liver injury recurred upon rechallenge, providing evidence that TYSABRI caused the injury.

- The combination of transaminase elevations and elevated bilirubin without evidence of obstruction is generally recognized as an important predictor of severe liver injury that may lead to death or the need for a liver transplant in some patients.

- TYSABRI should be discontinued in patients with jaundice or other evidence of significant liver injury (e.g., laboratory evidence).
Warnings and Precautions – Hypersensitivity/Antibody Formation

➢ TYSABRI has been associated with hypersensitivity reactions, including serious systemic reactions (e.g., anaphylaxis), which occurred at an incidence of <1%.

➢ Patients who receive TYSABRI after an extended period without treatment may be at higher risk of hypersensitivity reactions.

➢ If a hypersensitivity reaction occurs, discontinue the use of TYSABRI, and initiate appropriate therapy.

➢ Do not re-treat with TYSABRI.
The immune system effects of TYSABRI® may increase the risk for infections.

Concurrent use of antineoplastic, immunosuppressant, or immunomodulating agents may further increase the risk of infections, including PML and other opportunistic infections, over the risk observed with use of TYSABRI alone.

The safety and efficacy of TYSABRI in combination with antineoplastic, immunosuppressant, or immunomodulating agents have not been established.

For patients with Crohn’s disease who start TYSABRI while on chronic corticosteroids, commence steroid withdrawal as soon as a therapeutic benefit has occurred. If the patient cannot discontinue systemic corticosteroids within 6 months, discontinue TYSABRI.
The most frequently reported serious adverse reactions in the Study MS1 were infections (3.2% vs 2.6% placebo), acute hypersensitivity reactions (1.1% vs 0.3%), depression (1.0% vs 1.0%), and cholelithiasis (1.0% vs 0.3%).

The following serious adverse events in the induction Studies CD1 and CD2 were reported more commonly with TYSABRI than placebo and occurred at an incidence of at least 0.3%: intestinal obstruction or stenosis (2% vs. 1% in placebo), acute hypersensitivity reactions (0.5% vs. 0%), abdominal adhesions (0.3% vs. 0%), and cholelithiasis (0.3% vs. 0%).
The most common adverse reactions reported at an incidence of >10% were headache (38% vs 33%), fatigue (27% vs 21%), infusion reactions (24% vs 18%), urinary tract infections (21% vs 17%), arthralgia (19% vs 14%), depression (19% vs 16%), lower respiratory tract infection (17% vs 16%), pain in extremity (16% vs 14%), rash (12% vs 9%), gastroenteritis (11% vs 9%), abdominal discomfort (11% vs 10%), vaginitis* (10% vs 6%), and diarrhea (10% vs 9%).

*Percentage based on female patients only.

Other common adverse reactions (incidence 10%) in the CD population were upper respiratory tract infections and nausea.

Based on animal data, TYSABRI may cause fetal harm. TYSABRI should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus.
Program Overview

What is the TOUCH Prescribing Program?

What tools support the TOUCH Prescribing Program?
- MS TOUCH Educational Materials
- CD TOUCH Educational Materials

What is the enrollment process?

What is the process to infuse TYSABRI®?

How are patients tracked?

What is TOUCH On-Line?
What is the TOUCH Prescribing Program?

A program that makes TYSABRI® available only to prescribers, infusion centers, pharmacies associated with infusion centers, and patients who are enrolled in the program.
What was the TOUCH Prescribing Program designed to do?

- To inform prescribers, infusion center healthcare providers, and patients about the risk of progressive multifocal leukoencephalopathy (PML) associated with TYSABRI® including the increased risk of PML with longer treatment duration, prior immunosuppressant use, and the presence of anti-JCV antibodies.

- To warn against concurrent use with antineoplastic, immunosuppressant, or immunomodulating agents and in patients who are immunocompromised.

- To promote early diagnosis of PML and timely discontinuation of TYSABRI in the event of suspected PML.
What are the program requirements?

**Prescribers**
- Must be registered in and meet all the requirements of the TOUCH Prescribing Program to
  - Prescribe TYSABRI®

**Infusion Sites**
- Infuse TYSABRI

**Pharmacies**
- Dispense TYSABRI

**Patients**
- Must be enrolled in and meet all the requirements of the TOUCH Prescribing Program to
  - Receive TYSABRI
There are 3 main components of the TOUCH Prescribing Program:

- **Enroll**
  - Prescribers and Patients
  - Infusion Sites
  - Pharmacies

- **Infuse**
  - TYSABRI® is only administered to enrolled patients with a current status of ‘Authorized’
  - Pre-infusion Patient Checklist is completed and submitted to the TOUCH Prescribing Program

- **Track**
  - Patients are tracked longitudinally to gather important safety information

**NOTE:** This overview of the TOUCH Prescribing Program components does not include a complete list of the program requirements.
Program Overview

- What is the TOUCH Prescribing Program?

- What tools support the TOUCH Prescribing Program?
  - MS TOUCH Educational Materials
  - CD TOUCH Educational Materials

- What is the enrollment process?

- What is the process to infuse TYSABRI®?

- How are patients tracked?

- What is TOUCH On-Line?
Tools to Support the TOUCH Prescribing Program – MS

Enrollment Forms

Prescriber/Patient

Infusion Site

Certified Pharmacy

Patient Medication Guide

Notice of Patient Authorization

Pre-infusion Patient Checklist

Guidance for Evaluation of New Neurological Symptoms in Patients Receiving TYSABRI®

TOUCH Prescribing Program Overview
Tools to Support the TOUCH Prescribing Program – Crohn’s Disease

Enrollment Forms

- Prescriber/Patient
- Infusion Site
- Certified Pharmacy
- Notice of Patient Authorization
- Patient Medication Guide
- Pre-infusion Patient Checklist
- Understanding PML for Gastroenterologists
- TOUCH Prescribing Program Overview
How Do I Communicate With TOUCH?

WEB
expected: TouchH On-Line
   www.touchprogram.com

PHONE
   1-800-456-2255
   Monday – Friday 8:30AM – 8PM EST

PAPER
   Fax: 1-800-840-1278
Satisfying TOUCH Prescribing Program Requirements

- The TOUCH Prescribing Program has been designed to facilitate appropriate use of TYSABRI®

- In order to assess if the Program is meeting its goals, registered sites and enrolled participant’s compliance may be reviewed by the FDA, and/or audited by Biogen Idec and/or a third party designated by Biogen Idec

- Compliance with the requirements of the TOUCH Prescribing Program is necessary to maintain authorization to prescribe, dispense, infuse, or receive TYSABRI. Failure to comply with these requirements may result in de-enrollment from the TOUCH Prescribing Program and termination of such authorization.
Program Overview

What is the TOUCH Prescribing Program?

What tools support the TOUCH Prescribing Program?
  – MS TOUCH Educational Materials
  – CD TOUCH Educational Materials

What is the enrollment process?

What is the process to infuse TYSABRI®?

How are patients tracked?

What is TOUCH On-Line?
Prescriber/Patient Enrollment
How do prescribers and patients enroll?

**Education**

Prescriber and Patient discuss TYSABRI® as a treatment option.

**Treatment Decision**

Patient reads the Patient Medication Guide and discusses the benefits and risks of TYSABRI with his/her prescriber.

**Enrollment**

Prescriber and patient complete, sign, and fax ALL PAGES of the Prescriber/Patient Enrollment Form to the TOUCH Prescribing Program to initiate therapy.

**Authorization**

TOUCH Case Manager confirms that all paperwork is complete and updates patient status to ‘Authorized’.

OR

TOUCH Case Manager sends a Notice of Patient Authorization and a copy of the Prescriber/Patient Enrollment Form to the authorized Infusion Site.
Enrollment Tools

Enroll
Infuse
Track
Guidance for Evaluation of New Neurological Symptoms in Patients Receiving TYSABRI®

Brochure provided by Biogen Idec
Resource for: Neurology specialists

Key topics include:

➤ Importance of careful evaluation of any new or recurrent symptoms
➤ Differentiating between the signs, symptoms, and lesion characteristics typical of MS and PML
➤ PML diagnostic algorithm incorporating MRI and CSF assessment
➤ Action steps if PML is suspected
➤ Guidance on the treatment of relapse and other neurological symptoms

The information provided in this brochure is an educational resource and is not intended to be a substitute for consultation and review of relevant reference materials and medical literature. Treatment decisions should be made based on the context of the situation and clinical judgment.

Reference ID: 3753128
Enrollment Tools

Enroll
Infuse
Track
Understanding PML

Flashcard provided by Biogen Idec
Resource for: Gastroenterologists, Internists, or other non-Neurology specialists

Key topics include:
- Characteristics of PML
- Guidance on recognizing PML in context of Crohn’s disease
- Action steps if PML is suspected

The information provided in this brochure is an educational resource and is not intended to be a substitute for consultation and review of relevant reference materials and medical literature. Treatment decisions should be made based on the context of the situation and clinical judgment.

Reference ID: 3753128
Infusion Site Enrollment
How does an Infusion Site enroll?

A Biogen Idec representative provides mandatory indication specific TOUCH Prescribing Program training to Infusion Site*

**Infusion Site** completes and faxes the **Infusion Site Enrollment Form** to TOUCH Prescribing Program

**TOUCH Prescribing Program** confirms that all paperwork is complete, assigns a Site Authorization Number, and faxes and mails the Site Authorization Confirmation to the Infusion Site

* NOTE: Infusion Sites must select to receive training and education materials on MS TOUCH, CD TOUCH, or both.

A patient will be matched **ONLY** with Infusion Sites that have been trained on the program materials related to his/her indication.
Certified Pharmacy Enrollment

Enroll

Infuse

Track
How does a Certified Pharmacy* enroll?

A Biogen Idec representative provides training to the Certified Pharmacy regarding the TOUCH Prescribing Program.

A Certified Pharmacy completes and faxes the Certified Pharmacy Enrollment Form to TOUCH Prescribing Program.

TOUCH Prescribing Program confirms that all paperwork is complete, assigns a Site Authorization Number, and faxes and mails the Site Authorization Confirmation to the Certified Pharmacy.

*A Certified Pharmacy is located within a hospital, group practice, or infusion site and is associated with an infusion site.

Reference ID: 3753128
Program Overview

- What is the TOUCH Prescribing Program?
- What tools support the TOUCH Prescribing Program?
  - MS TOUCH Educational Materials
  - CD TOUCH Educational Materials
- What is the enrollment process?
- What is the process to infuse TYSABRI®?
- How are patients tracked?
- What is TOUCH On-Line?
Infusion Overview
What process must be completed in order to infuse TYSABRI®?

TYSABRI should NOT be prepared until the Pre-infusion Patient Checklist has been successfully completed.

Prior to EVERY infusion of TYSABRI:

1. Confirm that the patient is currently ‘Authorized’ to receive TYSABRI on TOUCH On-Line*
2. Provide the patient with the Patient Medication Guide
3. Complete the Pre-infusion Patient Checklist on TOUCH On-Line*


Infusion

ONLY upon successful completion of the Pre-infusion Patient Checklist:

- Start an IV line
- Mix TYSABRI
- Infuse TYSABRI over 1 hour and observe patient for 1 hour post-infusion
- Submit completed Pre-infusion Patient Checklist via TOUCH On-Line* within 1 business day
Checking Patient Authorization Status

Only patients with a status ‘Authorized’ can receive TYSABRI®

- Check patient status as ‘Authorized’ on TOUCH On-Line

Paper process: Notice of Patient Authorization is faxed to both Prescriber and Infusion Site; a copy must be placed in the patient record
Pre-infusion Patient Checklist

All Infusion Sites must complete, sign, and submit the **Pre-infusion Patient Checklist** at every infusion visit.

Submit form within 1 business day of patient’s visit via TOUCH On-Line.

Paper process: Fax page one to 1-800-840-1278 and place original in the patient’s record.

NOTE: Pre-infusion Patient Checklist **must** be completed and submitted whether or not the patient is infused.

Reference ID: 3753128
Program Overview

What is the TOUCH Prescribing Program?

What tools support the TOUCH Prescribing Program?
- MS TOUCH Educational Materials
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What is the enrollment process?

What is the process to infuse TYSABRI®?

How are patients tracked?

What is TOUCH On-Line?
Tracking Overview

Enroll
Infuse
Track
NOTE: Missing or incomplete TOUCH Prescribing Program forms will prompt continued follow-up by a TOUCH Compliance Manager.

Reference ID: 3753128
Prescriber Must Reauthorize the Use of TYSABRI® Every 6 Months

TYSABRI Patient Status Report and Reauthorization Questionnaire

Prescriber will receive a Patient Status Report and Reauthorization Questionnaire every 6 months.

Completion of this form is required as it determines whether the prescriber authorizes the patient to receive TYSABRI for the next 6 months.

OR

Reference ID: 3753128
If a patient discontinues TYSABRI®, important health information is collected and tracked over time.

The prescriber will be sent Discontinuation Questionnaires, which must be completed and submitted to the TOUCH Prescribing Program via TOUCH On-Line.

Paper process: Upon notification of patient discontinuation, the Discontinuation Questionnaire will be faxed to the prescriber.

Fax completed form to 1-800-840-1278 and place original in the patient’s file.

*NOTE: Discontinuation Questionnaires are ONLY sent upon notification of discontinuation and again six months following discontinuation of TYSABRI.
Tracking Tools
The TOUCH Prescribing Program will track all patients over time, so that Biogen Idec can inform the FDA, prescribers, and patients in a timely manner of information regarding the safety of TYSABRI®.

**Pre-infusion Patient Checklist (Every 28 days)**

**TYSABRI Patient Status Report and Reauthorization Questionnaire (Every 6 months)**

**Initial and 6-Month Discontinuation Questionnaire**

*NOTE: Discontinuation Questionnaires are ONLY sent upon notification of discontinuation and again six months following discontinuation of TYSABRI*
Tracking Tools

- Enroll
- Infuse
- Track
The TOUCH Prescribing Program will track all patients over time, so that Biogen Idec can inform the FDA, prescribers, and patients in a timely manner of information regarding the safety of TYSABRI®.

<table>
<thead>
<tr>
<th>Pre-infusion Patient Checklist (Every 28 days)</th>
<th>12-Week Questionnaire (After initial 12 weeks)</th>
<th>Patient Status Report and Reauthorization Questionnaire (Every 6 months)</th>
<th>Initial and 6-Month Discontinuation Questionnaire*</th>
</tr>
</thead>
</table>

*NOTE: Discontinuation Questionnaires are ONLY sent upon notification of discontinuation and again six months following discontinuation of TYSABRI.
Program Overview

What is the TOUCH Prescribing Program?

What tools support the TOUCH Prescribing Program?
- MS TOUCH Educational Materials
- CD TOUCH Educational Materials

What is the enrollment process?

What is the process to infuse TYSABRI®?

How are patients tracked?

What is TOUCH On-Line?

How is TYSABRI acquired?
TOUCH On-Line Overview

- TOUCH On-Line is a Web-based tool designed to:
  - Provide real-time access to TYSABRI® patient data
  - Maintain compliance with the TOUCH Prescribing Program
  - Streamline communication to/from Prescribers and Infusion Sites

- TOUCH On-Line is available only to enrolled TOUCH participants

- TOUCH On-Line is accessed with secure user name and password

Reference ID: 3753128
The TOUCH Prescribing Program makes TYSABRI® available only to prescribers, infusion sites, pharmacies associated with infusion sites, and patients who are enrolled in the program.

There are 3 main components of the program: Enroll – Infuse – Track.

TYSABRI must be administered only to patients who are enrolled in and meet all the conditions of the TOUCH Prescribing Program.

Indication-specific training and educational materials are required for a site to become authorized on MS TOUCH, CD TOUCH or both.

TOUCH On-Line is a web-based tool available only to authorized infusion sites and prescribers enrolled in TOUCH.

Only authorized infusion sites and their associated certified pharmacies may acquire TYSABRI.
Overview

Please see accompanying full Prescribing Information, including Boxed Warning, as well as Important Safety Information on pages 6-9.
INDICATIONS AND USAGE

Multiple Sclerosis (MS)

TYSABRI (natalizumab) is indicated as monotherapy for the treatment of patients with relapsing forms of multiple sclerosis. TYSABRI increases the risk of PML. When initiating and continuing treatment with TYSABRI, physicians should consider whether the expected benefit of TYSABRI is sufficient to offset this risk. See Full Prescribing Information regarding the risk of PML with TYSABRI.

Crohn’s Disease (CD)

TYSABRI is indicated for inducing and maintaining clinical response and remission in adult patients with moderately to severely active Crohn's disease (CD) with evidence of inflammation who have had an inadequate response to, or are unable to tolerate, conventional CD therapies and inhibitors of TNF α. TYSABRI should not be used in combination with immunosuppressants (e.g., 6 mercaptopurine, azathioprine, cyclosporine or methotrexate) or inhibitors of TNF α.

Why the program was developed

Biogen Idec is committed to patient safety. The TOUCH Prescribing Program was designed:

- To inform prescribers, infusion center healthcare providers, and patients about the risk of progressive multifocal leukoencephalopathy (PML) associated with TYSABRI including the increased risk of PML with longer treatment duration, prior immunosuppressant use, and the presence of anti JCV antibodies
- To warn against concurrent use with antineoplastic, immunosuppressant, or immunomodulating agents and in patients who are immunocompromised
- To promote early diagnosis of PML and timely discontinuation of TYSABRI in the event of suspected PML

Prescribers, infusion sites, certified pharmacies associated with infusion sites, and patients must all enroll in the TOUCH Prescribing Program in order to prescribe, infuse, dispense, or receive TYSABRI. All completed Enrollment Forms must be faxed to Biogen Idec at 1 800 840 1278.

Please see accompanying full Prescribing Information, including Boxed Warning, as well as Important Safety Information on pages 6-9.
TOUCH On Line is a web based tool designed to:
- Provide real time access to TYSABRI patient data
- Maintain compliance with the TOUCH Prescribing program
- Reduce administrative burden/paperwork for Prescribers and Infusion Sites

TOUCH On Line is accessed with secure user name and password

**How the program works**

Patients advise prescribers of any new signs and symptoms and keep track of any concurrently used medications

Prescriber and patient discuss the potential benefits and known risks of TYSABRI, including PML, and enroll in TOUCH Prescribing Program

TOUCH Prescribing Program assures that the patient is assigned to an authorized infusion site

Pre-infusion Patient Checklist is administered at every infusion visit to determine whether patient can be infused

Every 6 months, the prescriber determines whether the patient will be reauthorized to receive TYSABRI for the next 6 months

TOUCH Prescribing Program tracks patients over time to monitor the long-term safety of TYSABRI

This Overview serves only as an introduction to the program. For additional details please see the full Prescribing Information, or call 1 800 456 2255.

**Compliance with the requirements of the TOUCH Prescribing Program is necessary to maintain authorization to prescribe, dispense, infuse, or receive TYSABRI. Failure to comply with these requirements may result in de-enrollment from the TOUCH Prescribing Program and termination of such authorization.**

For more information on the TOUCH Prescribing Program or to obtain additional copies of material, please contact your Biogen Idec representative or call 1 800 456 2255, 8:30 AM to 8:00 PM (ET). The Patient Medication Guide is also available online at [www.TYSABRI.com](http://www.TYSABRI.com).
Enrollment

All participants must enroll in the TOUCH Prescribing Program by completing an Enrollment Form.

**Prescribers and Patients**
Prior to enrollment, prescribers must receive and review the full Prescribing Information and educational materials relating to the TOUCH Prescribing Program. Before completing and signing a Prescriber/Patient Enrollment Form, prescribers and patients are required to:

- Understand and discuss the benefits and risks of treatment with TYSABRI, including PML and other opportunistic infections
- Understand and acknowledge their respective program responsibilities as outlined in the Enrollment Kit

Patients should be fully counseled by either the enrolled prescriber or a healthcare provider under that prescriber’s direction before an initial prescription is written. A copy of the completed Prescriber/Patient Enrollment Form should be retained in the patient’s medical record. Upon receipt of a properly completed Enrollment Form:

- A Patient Enrollment Number will be assigned
- A Biogen Idec Case Manager will be assigned to assure that the patient is assigned to an authorized infusion site

**Infusion Sites and Certified Pharmacies**
Before completing and signing their respective Enrollment Forms, infusion sites and associated certified pharmacies must receive training from a Biogen Idec representative.

*A certified pharmacy is a pharmacy that is part of a hospital, group practice, or infusion site, and is affiliated with one or more infusion sites. Retail pharmacies, wholesalers, and specialty distributors are excluded from holding inventory and dispensing TYSABRI.*

Please see accompanying full Prescribing Information, including Boxed Warning, as well as Important Safety Information on pages 6-9.
Program Overview

**Infusion**

Only infusion sites authorized by the TOUCH Prescribing Program can infuse TYSABRI. They are required to:

- Confirm that the patient is currently authorized to receive TYSABRI
- Provide the patient with a copy of the TYSABRI Patient Medication Guide prior to each infusion
- Administer the Pre Infusion Patient Checklist to every patient prior to each infusion and submit it to Biogen Idec within 1 business day, regardless of whether the patient is infused or not

Authorized infusion sites must use the Authorization Number that is provided upon enrollment to order and receive shipments of TYSABRI.* Certified pharmacies may only dispense TYSABRI to authorized, associated infusion sites.

*The TOUCH Prescribing Program utilizes a closed distribution system that restricts all product shipments. This system includes a single distributor, specialty pharmacies under contract with Biogen Idec and authorized certified pharmacies associated with authorized infusion sites.

**Tracking**

The TOUCH Prescribing Program will track all patients over time so that Biogen Idec can inform the FDA, prescribers, and patients in a timely manner of information regarding the safety of TYSABRI. Prescribers are required to report any case of PML, serious opportunistic infection, or death in TYSABRI treated patients to Biogen Idec or the FDA. Furthermore, prescribers are also required to cooperate in the investigation of potential adverse events including providing relevant information upon request. The primary tracking tools include:

- Pre infusion Patient Checklist
- Patient Status Report and Reauthorization Questionnaire
- Initial and 6 month Discontinuation Questionnaires

Missing or incomplete forms will prompt TOUCH Case Managers to follow up with infusion sites, patients, and/or prescribers to obtain such information in compliance with program requirements. Prescribers, infusion sites, and certified pharmacies may be audited by the FDA, Biogen Idec, and/or a third party authorized by Biogen Idec.

Reference ID: 3753128
IMPORTANT SAFETY INFORMATION

Indication
TYSABRI (natalizumab) is indicated as monotherapy for the treatment of patients with relapsing forms of multiple sclerosis. TYSABRI increases the risk of PML. When initiating and continuing treatment with TYSABRI, physicians should consider whether the expected benefit of TYSABRI is sufficient to offset this risk. See important information regarding the risk of PML with TYSABRI.

WARNING
TYSABRI (natalizumab) increases the risk of PML, an opportunistic viral infection of the brain that usually leads to death or severe disability. Risk factors for the development of PML include duration of therapy, prior use of immunosuppressants, and presence of anti-JCV antibodies. These factors should be considered in the context of expected benefit when initiating and continuing treatment with TYSABRI.

Healthcare professionals should monitor patients on TYSABRI for any new sign or symptom that may be suggestive of PML. TYSABRI dosing should be withheld immediately at the first sign or symptom suggestive of PML. For diagnosis, an evaluation including a gadolinium-enhanced MRI scan of the brain and, when indicated, cerebrospinal fluid analysis for JC viral DNA are recommended.

Because of the risk of PML, TYSABRI is available only through a restricted program under a Risk Evaluation and Mitigation Strategy (REMS) called the TOUCH Prescribing Program.

Progressive Multifocal Leukoencephalopathy (PML)
- Infection by the JC Virus is required for the development of PML.
- Anti JCV antibody testing should not be used to diagnose PML.
- There are no known interventions that can reliably prevent PML or that can adequately treat PML if it occurs. It is not known whether early detection of PML and discontinuation of TYSABRI will mitigate the disease.
- PML has been reported following discontinuation of TYSABRI in patients who did not have findings suggestive of PML at the time of discontinuation. Patients should continue to be monitored for any new signs or symptoms that may be suggestive of PML for approximately six months following discontinuation of TYSABRI.
- In MS patients, an MRI scan should be obtained prior to initiating therapy with TYSABRI. This MRI may be helpful in differentiating subsequent multiple sclerosis symptoms from PML.
- Three sessions of plasma exchange over 5 to 8 days were shown to accelerate TYSABRI clearance in a study of 12 patients with MS who did not have PML, although in the majority of patients, alpha 4 integrin receptor binding remained high. Adverse events which may occur during plasma exchange include clearance of other medications and volume shifts, which have the potential to lead to hypotension or pulmonary edema. Although plasma exchange has not been studied in TYSABRI treated patients with PML, it has been used in such patients in the postmarketing setting to remove TYSABRI more quickly from the circulation.
- Anti JCV antibody testing should not be performed during or for at least two weeks following plasma exchange due to the removal of antibodies from the serum.

Please see accompanying full Prescribing Information, including Boxed Warning.
Immune reconstitution inflammatory syndrome (IRIS) has been reported in the majority of TYSABRI treated patients who developed PML and subsequently discontinued TYSABRI. In almost all cases, IRIS occurred after plasma exchange was used to eliminate circulating TYSABRI. It presents as a clinical decline in the patient’s condition after TYSABRI removal (and in some cases after apparent clinical improvement) that may be rapid, can lead to serious neurological complications or death and is often associated with characteristic changes in the MRI. TYSABRI has not been associated with IRIS in patients discontinuing treatment with TYSABRI for reasons unrelated to PML. In TYSABRI treated patients with PML, IRIS has been reported within days to several weeks after plasma exchange. Monitoring for development of IRIS and appropriate treatment of the associated inflammation should be undertaken.

**Contraindications**

- TYSABRI is contraindicated in patients who have or have had PML.
- TYSABRI is contraindicated in patients who have had a hypersensitivity reaction to TYSABRI.

**TYSABRI TOUCH Prescribing Program**

- TYSABRI is available only through a restricted program under a REMS called the TOUCH® Prescribing Program because of the risk of PML.
- For prescribers and patients, the TOUCH® Prescribing Program has two components: MS TOUCH® (for patients with multiple sclerosis) and CD TOUCH® (for patients with Crohn's disease).
- Prescribers must be certified and comply with the following:
  - Review the TOUCH Prescribing Program prescriber educational materials, including the full prescribing information.
  - Educate patients on the benefits and risks of treatment with TYSABRI, ensure that patients receive the Medication Guide, and encourage them to ask questions.
  - Review, complete, and sign the Patient Prescriber Enrollment Form.
  - Evaluate patients three months after the first infusion, six months after the first infusion, every six months thereafter, and for at least six months after discontinuing TYSABRI.
  - Determine every six months whether patients should continue on treatment, and if so, authorize treatment for another six months.
  - Submit to Biogen Idec the “TYSABRI Patient Status Report and Reauthorization Questionnaire” six months after initiating treatment and every six months thereafter.
  - Complete an “Initial Discontinuation Questionnaire” when TYSABRI is discontinued and a “6 Month Discontinuation Questionnaire,” following discontinuation of TYSABRI.
  - Report cases of PML, hospitalizations due to opportunistic infections, and deaths to Biogen Idec at 1 800 456 2255 and to the Food and Drug Administration’s MedWatch Program at 1 800 FDA 1088 as soon as possible.
- Patients must be enrolled in the TOUCH Prescribing Program, read the Medication Guide, understand the risks associated with TYSABRI, and complete and sign the Patient Prescriber Enrollment Form.
- Pharmacies and infusion centers must be specially certified to dispense or infuse TYSABRI.
Herpes Encephalitis and Meningitis

- Tysabri increases the risk of developing encephalitis and meningitis caused by herpes simplex and varicella zoster viruses.
- Serious, life threatening, and sometimes fatal cases have been reported in the postmarketing setting in multiple sclerosis patients receiving Tysabri.
- Monitor patients receiving Tysabri for signs and symptoms of meningitis and encephalitis. If herpes encephalitis or meningitis occurs, TYSABRI should be discontinued, and appropriate treatment for herpes encephalitis/meningitis should be administered.

Hepatotoxicity

- Clinically significant liver injury, including acute liver failure requiring transplant, has been reported in patients treated with TYSABRI in the postmarketing setting. In some patients, liver injury recurred upon rechallenge, providing evidence that TYSABRI caused the injury.
- Signs of liver injury, including markedly elevated serum hepatic enzymes and elevated total bilirubin, occurred as early as 6 days after the first dose; and signs of liver injury have also been reported for the first time after multiple doses.
- In some patients, liver injury recurred upon rechallenge, providing evidence that TYSABRI caused the injury.
- The combination of transaminase elevations and elevated bilirubin without evidence of obstruction is generally recognized as an important predictor of severe liver injury that may lead to death or the need for a liver transplant in some patients.
- TYSABRI should be discontinued in patients with jaundice or other evidence of significant liver injury (e.g., laboratory evidence).

Hypersensitivity/Antibody Formation

- Hypersensitivity reactions have occurred in patients receiving TYSABRI, including serious systemic reactions (e.g., anaphylaxis), which occurred at an incidence of <1%.
- Reactions usually occur within 2 hours of the start of the infusion. Symptoms associated with these reactions can include urticaria, dizziness, fever, rash, rigors, pruritus, nausea, flushing, hypotension, dyspnea, and chest pain. Generally, these reactions are associated with antibodies to TYSABRI.
- If a hypersensitivity reaction occurs, discontinue administration of TYSABRI, and initiate appropriate therapy. Patients who experience a hypersensitivity reaction should not be retreated with TYSABRI.
- Hypersensitivity reactions were more frequent in patients with antibodies to TYSABRI compared with patients who did not develop antibodies to TYSABRI in both MS and CD studies.
- Patients who receive TYSABRI after an extended period without treatment may be at higher risk of hypersensitivity reactions.

Immunosuppression/Infections

- The immune system effects of TYSABRI may increase the risk for infections.

Please see accompanying full Prescribing Information, including Boxed Warning.
In Study MS1, certain types of infections, including pneumonias and urinary tract infections (including serious cases), gastroenteritis, vaginal infections, tooth infections, tonsillitis, and herpes infections, occurred more often in TYSABRI treated patients than in placebo treated patients. One opportunistic infection, a cryptosporidial gastroenteritis with a prolonged course, was observed in a patient who received TYSABRI in Study MS1.

In Studies MS1 and MS2, an increase in infections was seen in patients concurrently receiving short courses of corticosteroids. However, the increase in infections in TYSABRI treated patients who received steroids was similar to the increase in placebo treated patients who received steroids.

Concurrent use of antineoplastic, immunosuppressant, or immunomodulating agents may further increase the risk of infections over the risk observed with use of TYSABRI alone. The safety and efficacy of TYSABRI in combination with antineoplastic, immunosuppressant, or immunomodulating agents have not been established.

In Study MS1 and Study MS2, the rate of any type of infection was approximately 1.5 per patient year in both TYSABRI treated patients and placebo treated patients.

In Study MS1, the incidence of serious infections was approximately 3% in TYSABRI treated patients and in placebo treated patients. Most patients did not interrupt treatment with TYSABRI during infections.

In postmarketing experience, serious herpes infections have occurred.

**Laboratory Test Abnormalities**

In clinical trials, TYSABRI was observed to induce increases in circulating lymphocytes, monocytes, eosinophils, basophils, and nucleated red blood cells. Observed changes persisted during TYSABRI exposure, but were reversible, returning to baseline levels usually within 16 weeks after the last dose. Elevations of neutrophils were not observed. TYSABRI induces mild decreases in hemoglobin levels that are frequently transient.

**Adverse Reactions**

The most common adverse reactions reported at an incidence of $\leq 10\%$ with TYSABRI and $\geq 2\%$ difference with placebo were headache (38% vs 33%), fatigue (27% vs 21%), infusion reactions (24% vs 18%), urinary tract infections (21% vs 17%), arthralgia (19% vs 14%), depression (19% vs 16%), pain in extremity (16% vs 14%), rash (12% vs 9%), gastroenteritis (11% vs 9%), and vaginitis* (10% vs 6%).

*Percentage based on female patients only.

The most frequently reported serious adverse reactions in Study MS1 were infections (3.2% vs 2.6% placebo), including urinary tract infection (0.8% versus 0.3%) and pneumonia (0.6% versus 0%), acute hypersensitivity reactions (1.1% vs 0.3%, including anaphylaxis/anaphylactoid reaction [0.8% versus 0%]), depression (1.0% vs 1.0%, including suicidal ideation or attempt [0.6% versus 0.3%]), and cholelithiasis (1.0% vs 0.3%).

Based on animal data, TYSABRI may cause fetal harm. TYSABRI should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus.

Please see accompanying full Prescribing Information, including Boxed Warning.
**Important Responsibilities**

**PRESCRIBERS**—Among the important responsibilities of prescribers in the TOUCH Prescribing Program are the following:

- Acknowledge that TYSABRI should only be prescribed in accordance with the FDA label
- Educate the patient on the benefits and risks of treatment with TYSABRI by using the Patient Medication Guide
- Evaluate the patient 3 and 6 months after the first infusion, and every 6 months thereafter, and for 6 months after TYSABRI has been discontinued
- Determine every 6 months whether the patient should continue on treatment, and if so, reauthorize treatment
- Submit to Biogen Idec the TYSABRI Patient Status Report and Reauthorization Questionnaire 6 months after initiating treatment and every 6 months thereafter
- Report serious opportunistic infections and atypical infections with TYSABRI to Biogen Idec at 1 800 456 2255 and to the Food and Drug Administration’s MedWatch program at 1 800 FDA 1088

**PATIENTS**—Among the important responsibilities of patients in the TOUCH Prescribing Program are the following:

- Bring to each infusion a list of all medicines and treatments they have taken during the last month
- Read the Patient Medication Guide before starting TYSABRI and before each TYSABRI infusion
- Promptly report any continuously worsening symptoms that persist over several days to their prescriber
- Inform all of their physicians that they are receiving TYSABRI
- Plan to see their prescriber 3 and 6 months after the first infusion, and at least as frequently as every 6 months thereafter

**INFUSION SITES**—Among the important responsibilities of infusion sites in the TOUCH Prescribing Program are the following:

- Confirm that the patient is currently authorized to receive TYSABRI
- Provide the patient with a copy of the TYSABRI Patient Medication Guide
- Administer the Pre infusion Patient Checklist to every patient prior to each infusion and submit to Biogen Idec within 1 business day, regardless of whether the patient is infused or not

Please see accompanying full Prescribing Information, including Boxed Warning, as well as Important Safety Information on pages 6-9.
YTSABRI (raltegravir) is available only through the TOUCH Prescribing Program, which stands for TSAR-Unified Intravenous and Oral Regimen. The TOUCH Prescribing Program is a nationwide distribution program focused on safety and adherence with the intent of reducing HIV drug resistance. Identification of TSAR patients and distribution of YTSABRI are monitored and tracked. TSAR and QD are pre-authorized prescription programs that must be certified by patients' prescribers. TSAR patients must be confirmed as intravenous patients on TSAR by the TOUCH Prescribing Program. Patients who switch to the TSAR program from QD, or vice versa, must be confirmed as intravenous patients on TSAR by the TOUCH Prescribing Program.

The TOUCH Prescribing Program is designed to:

- Ensure patient safety
- Enhance treatment adherence
- Reduce the risk of HIV drug resistance
- Increase access to raltegravir

Healthcare providers are required to follow the TOUCH Prescribing Program as it relates to the use of YTSABRI. Providers are required to refer to the TOUCH Prescribing Program website for up-to-date information and to ensure that patients are enrolled in the program.

TOUCH Prescribing Program participants in fulfilling their TOUCH Prescribing Program Requirements

- Complete the TOUCH On-Line Log-In
- Complete the TOUCH On-Line Log-In
- Complete the TOUCH On-Line Log-In
- Complete the TOUCH On-Line Log-In
- Complete the TOUCH On-Line Log-In

Not a TOUCH On-Line Log-In

[Details of the TOUCH On-Line Log-In process]

Indications

YTSABRI (raltegravir) is indicated as part of a combination antiretroviral therapy regimen for the treatment of adult patients with documented failure of an antiretroviral regimen containing two or more reverse transcriptase inhibitors. It is not indicated for use in patients with documented or suspected resistance to raltegravir.

Contraindications

YTSABRI is contraindicated in patients with a history of hypersensitivity reactions with raltegravir, IRBIVAN (raltegravir), or any component of the formulation.

Warnings

- Hepatic Transaminase Increases: YTSABRI is contraindicated in patients with severe hepatic impairment (Child-Pugh class C) or with total bilirubin levels greater than 3 mg/dL.
- Drug-Drug Interactions: YTSABRI is contraindicated in patients taking medications with a narrow therapeutic index (e.g., digoxin, digoxin-digoxigenin [Digoxin-Digoxigenin]).

Adverse Reactions

The most common adverse reactions associated with YTSABRI are nausea, diarrhea, vomiting, and abdominal pain. These adverse reactions are generally mild to moderate in severity and are often transient. In clinical trials, the rate of diarrhea was higher in patients taking YTSABRI compared to those taking placebo.

Drug Interactions

YTSABRI is generally well tolerated and has a low incidence of drug interactions. However, it should be used with caution in patients taking medications with a narrow therapeutic index or those with a history of liver disease.

Dosage and Administration

- The recommended dose of YTSABRI is 400 mg once daily, taken with food.
- YTSABRI should be administered with food to minimize the risk of stomach upset.
- YTSABRI is not recommended for use in patients with severe hepatic impairment (Child-Pugh class C) or with total bilirubin levels greater than 3 mg/dL.

Administration

- YTSABRI is administered orally as a single tablet.
- YTSABRI should be taken consistently with meals to ensure optimal absorption.

Patient Counseling

- Patients should be advised to take YTSABRI consistently with meals to ensure optimal absorption.
- Patients should be advised to consult their healthcare provider if they experience serious adverse reactions.

References

[List of references]

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