




ISSUE DATE November 6, 2024	EFFECTIVE DATE January 6, 2025	NUMBER *See below
SUBJECT Prior Authorization of Multiple Sclerosis Agents – Pharmacy Services		BY  Sally Kozak Deputy Secretary Office of Medical Assistance Programs

IMPORTANT REMINDER: All providers must revalidate the Medical Assistance (MA) enrollment of each service location every 5 years. Providers should log into PROMISE to check the revalidation dates of each service location and submit revalidation applications at least 60 days prior to the revalidation dates. Enrollment (revalidation) applications may be found at: <https://www.pa.gov/en/agencies/dhs/resources/for-providers/provider-enrollment-information/provider-enrollment-documents.html>.

PURPOSE:

The purpose of this bulletin is to issue updated handbook pages that include the requirements for prior authorization and the type of information needed to evaluate the medical necessity of prescriptions for Multiple Sclerosis Agents submitted for prior authorization.

SCOPE:

This bulletin applies to all licensed pharmacies and prescribers enrolled in the Medical Assistance (MA) Program. The guidelines to determine the medical necessity of Multiple Sclerosis Agents will be utilized in the fee-for-service and managed care delivery systems. Providers rendering services to MA beneficiaries in the managed care delivery system should address any questions related to the prior authorization of Multiple Sclerosis Agents to the appropriate managed care organization.

BACKGROUND/DISCUSSION:

The Department of Human Services (Department) is updating the medical necessity guidelines for Multiple Sclerosis Agents to add Briumvi (ublituximab-xiyy) to the list of drugs that require prior authorization, delete Ocrevus (ocrelizumab) from the list of examples of a non-preferred Multiple Sclerosis Agent with a dosing interval exceeding 90 days, and add a guideline to the requests for renewal of the prior authorization section related to requests for non-preferred Multiple Sclerosis Agents with a therapeutically equivalent brand or

*01-25-20	09-25-20	27-25-20	33-25-20
02-25-20	11-25-20	30-25-20	
03-25-20	14-25-20	31-25-20	
08-25-21	24-25-20	32-25-20	

COMMENTS AND QUESTIONS REGARDING THIS BULLETIN SHOULD BE DIRECTED TO:

Fee-for-Service Provider Service Center: 1-800-537-8862

Visit the Office of Medical Assistance Programs website at:

<https://www.pa.gov/en/agencies/dhs/departments-offices/omap-info.html>

generic, interchangeable biosimilar, or brand or unbranded biologic that is preferred on the Preferred Drug List. There are no other changes to the medical necessity guidelines.

The revisions to the guidelines to determine medical necessity of prescriptions for Multiple Sclerosis Agents were subject to public review and comment and subsequently approved for implementation by the Department.

PROCEDURE:

The procedures for prescribers to request prior authorization of Multiple Sclerosis Agents are located in SECTION I of the Prior Authorization of Pharmaceutical Services Handbook. The Department will take into account the elements specified in the clinical review guidelines (which are included in the provider handbook pages in the SECTION II chapter related to Multiple Sclerosis Agents) when reviewing the prior authorization request to determine medical necessity.

As set forth in 55 Pa. Code § 1101.67(a), the procedures described in the handbook pages must be followed to ensure appropriate and timely processing of prior authorization requests for drugs and products that require prior authorization.

ATTACHMENTS:

Prior Authorization of Pharmaceutical Services Handbook - Updated pages

RESOURCES:

Prior Authorization of Pharmaceutical Services Handbook – SECTION I
Pharmacy Prior Authorization General Requirements

<https://www.pa.gov/en/agencies/dhs/resources/pharmacy-services/pharmacy-prior-authorization-general-requirements.html>

Prior Authorization of Pharmaceutical Services Handbook – SECTION II
Pharmacy Prior Authorization Guidelines

<https://www.pa.gov/en/agencies/dhs/resources/pharmacy-services/clinical-guidelines.html>

MEDICAL ASSISTANCE HANDBOOK
PRIOR AUTHORIZATION OF PHARMACEUTICAL SERVICES

I. Requirements for Prior Authorization of Multiple Sclerosis Agents

A. Prescriptions That Require Prior Authorization

Prescriptions for Multiple Sclerosis Agents that meet any of the following conditions must be prior authorized:

1. A non-preferred Multiple Sclerosis Agent. See the Preferred Drug List (PDL) for the list of preferred Multiple Sclerosis Agents at: <https://papdl.com/preferred-drug-list>.
2. A Multiple Sclerosis Agent with a prescribed quantity that exceeds the quantity limit. The list of drugs that are subject to quantity limits, with accompanying quantity limits, is available at: <https://www.pa.gov/en/agencies/dhs/resources/pharmacy-services/quantity-limits-daily-dose-limits.html>.
3. A prescription for Ampyra (dalfampridine ER), Aubagio (teriflunomide), Briumvi (ublituximab-xiiy), Gilenya (fingolimod), Kesimpta (ofatumumab), Ocrevus (ocrelizumab), Tysabri (natalizumab), or Tecfidera (dimethyl fumarate DR).

B. Review of Documentation for Medical Necessity

In evaluating a request for prior authorization of a prescription for a Multiple Sclerosis Agent, the determination of whether the requested prescription is medically necessary will take into account whether the beneficiary:

1. For a natalizumab product, see the prior authorization guideline related to Natalizumab; **OR**
2. For Zeposia (ozanimod) for the treatment of ulcerative colitis, see the prior authorization guideline related to Ulcerative Colitis Agents; **OR**
3. Is being treated for a diagnosis that is indicated in the U.S. Food and Drug Administration (FDA)-approved package labeling or a medically accepted indication; **AND**
4. Is age-appropriate according to FDA-approved package labeling, nationally recognized compendia, or peer-reviewed medical literature; **AND**
5. Is prescribed a dose that is consistent with the FDA-approved package labeling, nationally recognized compendia, or peer-reviewed medical literature; **AND**
6. Is prescribed the Multiple Sclerosis Agent by **one** of the following:
 - a. For Ampyra (dalfampridine ER), a neurologist or physical medicine and rehabilitation (PM&R) specialist
 - b. For all other Multiple Sclerosis Agents, a neurologist;

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AND

7. Does not have a contraindication to the prescribed Multiple Sclerosis Agent; **AND**
8. For a non-preferred Multiple Sclerosis Agent, **one** of the following:
 - a. Has a history of therapeutic failure of or a contraindication or an intolerance to the preferred Multiple Sclerosis Agents approved for the beneficiary's diagnosis
 - b. **One** of the following:
 - i. Has a current prescription (within the past 90 days) for the same non-preferred Multiple Sclerosis Agent (does not apply to non-preferred brands when the therapeutically equivalent generic, interchangeable biosimilar, or unbranded biologic is preferred or to non-preferred generics, interchangeable biosimilars, or unbranded biologics when the therapeutically equivalent brand, interchangeable brand, or brand biologic is preferred)
 - ii. For a non-preferred Multiple Sclerosis Agent with a dosing interval exceeding 90 days (e.g., Lemtrada, Mavenclad), is receiving treatment with the same non-preferred Multiple Sclerosis Agent and will continue therapy at a dosing interval supported by FDA-approved package labeling, nationally recognized compendia, or peer-reviewed medical literature;

AND

9. For Ampyra (dalfampridine ER), has motor dysfunction on a continuous basis that impairs the ability to complete instrumental activities of daily living or activities of daily living; **AND**
10. For Mavenclad (cladribine), has documentation of a recent lymphocyte count within recommended limits according to FDA-approved package labeling before initiating the first treatment course; **AND**
11. If a prescription for a Multiple Sclerosis Agent is for a quantity that exceeds the quantity limit, the determination of whether the prescription is medically necessary will also take into account the guidelines set forth in the Quantity Limits Chapter.

NOTE: If the beneficiary does not meet the clinical review guidelines listed above but, in the professional judgment of the physician reviewer, the services are medically necessary to meet the medical needs of the beneficiary, the request for prior authorization will be approved.

FOR RENEWALS OF PRIOR AUTHORIZATION FOR MULTIPLE SCLEROSIS AGENTS: The determination of medical necessity of a request for renewal of a prior authorization for a Multiple Sclerosis Agent that was previously approved will take into account whether the beneficiary:

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1. Is prescribed a dose that is consistent with FDA-approved package labeling, nationally recognized compendia, or peer-reviewed medical literature; **AND**
2. Is prescribed the Multiple Sclerosis Agent by **one** of the following:
 - a. For Ampyra (dalfampridine ER), a neurologist or PM&R specialist
 - b. For all other Multiple Sclerosis Agents, a neurologist;

AND

3. Does not have a contraindication to the prescribed Multiple Sclerosis Agent; **AND**
4. **One** of the following:
 - a. For Ampyra (dalfampridine ER), has a documented improvement in motor function
 - b. For all other Multiple Sclerosis Agents, **one** of the following:
 - i. For a Multiple Sclerosis Agent prescribed for a diagnosis of a relapsing form of multiple sclerosis, has documented improvement or stabilization of the multiple sclerosis disease course
 - ii. For a Multiple Sclerosis Agent prescribed for a diagnosis of primary progressive multiple sclerosis, continues to benefit from the prescribed Multiple Sclerosis Agent based on the prescriber's assessment;

AND

5. For Lemtrada (alemtuzumab), received the previous treatment course at least 12 months prior to the requested treatment course with Lemtrada (alemtuzumab); **AND**
6. For Mavenclad (cladribine), **both** of the following:
 - a. Has documentation of a recent lymphocyte count within recommended limits according to FDA-approved package labeling before initiating the second treatment course
 - b. Has not exceeded the recommended total number of treatment courses according to FDA-approved package labeling;

AND

7. For a non-preferred Multiple Sclerosis Agent with a therapeutically equivalent brand or generic, interchangeable biosimilar, or brand or unbranded biologic that is preferred on the PDL, has a history of therapeutic failure of or a contraindication or an intolerance to the preferred therapeutically equivalent brand or generic, interchangeable biosimilar, or brand or unbranded biologic that would not be expected to occur with the requested drug;

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AND

8. If a prescription for a Multiple Sclerosis Agent is for a quantity that exceeds the quantity limit, the determination of whether the prescription is medically necessary will also take into account the guidelines set forth in the Quantity Limits Chapter.

NOTE: If the beneficiary does not meet the clinical review guidelines listed above but, in the professional judgment of the physician reviewer, the services are medically necessary to meet the medical needs of the beneficiary, the request for prior authorization will be approved.

C. Clinical Review Process

Prior authorization personnel will review the request for prior authorization and apply the clinical guidelines in Section B. above to assess the medical necessity of a prescription for a Multiple Sclerosis Agent. If the guidelines in Section B. are met, the reviewer will prior authorize the prescription. If the guidelines are not met, the prior authorization request will be referred to a physician reviewer for a medical necessity determination. Such a request for prior authorization will be approved when, in the professional judgment of the physician reviewer, the services are medically necessary to meet the medical needs of the beneficiary.

D. Dose and Duration of Therapy

Requests for prior authorization of Multiple Sclerosis Agents will be approved as follows:

1. For Ampyra (dalfampridine ER) or Aubagio (teriflunomide):
 - a. Initial requests will be approved for up to three months.
 - b. Renewal requests will be approved for up to six months.
2. For Lemtrada (alemtuzumab):
 - a. Requests for an initial treatment course will be approved for up to five days.
 - b. Requests for subsequent treatment courses will be approved for up to three days.
3. For Mavenclad (cladribine):
 - a. Requests for prior authorization will be approved for a duration of therapy consistent with FDA-approved package labeling.

E. References:

1. Ampyra Package Insert. Pearl River, NY: Acorda Therapeutics, Inc.; June 2022.
2. Aubagio Package Insert. Cambridge, MA: Genzyme Corporation; December 2022.
3. Bafiertam Package Insert. High Point, NC: Banner Life Sciences; January 2023.

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4. Briumvi Package Insert. Morrisville, NC: TG Therapeutics, Inc.; December 2022.
5. Clinical Resource, Multiple Sclerosis Treatments, The Pharmacists Letter/Prescriber's Letter. September 2017.
6. Gilenya Package Insert. East Hanover, New Jersey: Novartis Pharmaceuticals Corporation; July 2022.
7. Hauser SL, Bar-Or A, Comi G, et al. Ocrelizumab versus Interferon Beta-1a in Relapsing Multiple Sclerosis. *New England Journal of Medicine*. January 19, 2017; 376:221-234.
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10. Mavenclad Package Insert. Rockland, MA: EMD Serono, Inc.; September 2022.
11. Mayzent Package Insert. East Hanover, NJ: Novartis Pharmaceuticals Corporation; January 2023.
12. Montalban X, Hauser SL, Kappos L, et al. Ocrelizumab versus Placebo in Primary Progressive Multiple Sclerosis. *New England Journal of Medicine*. January 19, 2017. 376:209-220.
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18. Ponvory Package Insert. Titusville, NJ: Janssen Pharmaceuticals, Inc.; September 2022.
19. Rae-Grant A, Day GS, Marrie RA, et al. Practice guideline recommendations summary: Disease-modifying therapies for adults with multiple sclerosis: Report of the Guideline Development, Dissemination, and Implementation Subcommittee of the American Academy of Neurology. *Neurology* 2018; 90:777.
20. Tascenso ODT Package Insert. Cambridge, United Kingdom: Cycle Pharmaceuticals Ltd; December 2022.
21. Tecfidera Package Insert. Cambridge, MA: Biogen Inc.; February 2023.
22. Vumerity Package Insert. Cambridge, MA: Biogen Inc.; February 2023.
23. Zeposia Package Insert. Princeton, NJ: Bristol-Myers Squibb Company; August 2023.