



### Multiple Sclerosis Drugs

Ocrevus (ocrelizumab) J2350, Copaxone (Glatiramer acetate) J1595, Zenapax (daclizumab) J7513, Briumvi (ublituximab) J2329 are Non-preferred. The preferred products are Part D alternatives including Aubagio and generic glatiramer (See Part D Formulary, no PA required for most preferred Part D alts).

Prior Authorization Step Therapy Request  
Medicare Part B Form

Instructions: \* Indicates required information – Form may be returned if required information is not provided. Please fax this request to the appropriate fax number listed at the bottom of the page.

<input type="checkbox"/>	<b>Standard Request– (72 Hours)</b>	<input type="checkbox"/>	<b>Urgent Request</b> (standard time frame could place the member's life, health or ability in serious jeopardy)
Date Requested _____			
Requestor _____ Clinic name: _____ Phone _____ / Fax _____			

#### MEMBER INFORMATION

\*Name: \_\_\_\_\_ \*ID#: \_\_\_\_\_ \*DOB: \_\_\_\_\_

#### PRESCRIBER INFORMATION

\*Name: \_\_\_\_\_  MD  FNP  DO  NP  PA \*Phone: \_\_\_\_\_

\*Address: \_\_\_\_\_ \*Fax: \_\_\_\_\_

#### DISPENSING PROVIDER / ADMINISTRATION INFORMATION

\*Name: \_\_\_\_\_ Phone: \_\_\_\_\_

\*Address: \_\_\_\_\_ Fax: \_\_\_\_\_

#### PROCEDURE / PRODUCT INFORMATION

HCPC Code	Name of Drug	Dose (Wt: _____ kg Ht: _____ )	Frequency	End Date if known

Self-administered       Provider-administered       Home Infusion

Chart notes attached. Other important information: \_\_\_\_\_

**Diagnosis: ICD10:** \_\_\_\_\_ **Description:** \_\_\_\_\_

Provider attests the diagnosis provided is an FDA-Approved indication for this drug

#### CLINICAL INFORMATION

New Start or Initial Request: (Clinical documentation required for all requests)

**Provider has reviewed the attached “Criteria for Approval” and attests the member meets ALL required PA criteria.**

If not, please provide **clinical rationale** for formulary exception: \_\_\_\_\_

Continuation Requests: (Clinical documentation required for all requests)

**Provider has reviewed the attached “Criteria for Continuation” and attests the member meets ALL required PA Continuation criteria.**

Patient had an adequate response or significant improvement while on this medication.

If not, please provide clinical rationale for continuing this medication: \_\_\_\_\_

**ACKNOWLEDGEMENT**

**Request By (Signature Required):** \_\_\_\_\_ **Date:** \_\_\_\_ / \_\_\_\_ / \_\_\_\_

Any person who knowingly files a request for authorization of coverage of a medical procedure or service with the intent to injure, defraud or deceive any insurance company by providing materially false information or conceals material information for the purpose of misleading, commits a fraudulent insurance act, which is a crime and subjects such person to criminal and civil penalties. **THIS AUTHORIZATION IS NOT A GUARANTEE OF PAYMENT.** PAYMENT IS BASED ON BENEFITS IN EFFECT AT THE TIME OF SERVICE, MEMBER ELIGIBILITY AND MEDICAL NECESSITY.

## Prior Authorization Group – Multiple Sclerosis Drugs PA

### Drug Name(s):

**OCREVUS  
COPAXONE  
ZENAPAX  
BRIUMVI**

**OCRELIZUMAB  
GLATIRAMER ACETATE  
DACLIZUMAB  
UBLITUXIMAB**

### Criteria for approval of Prior Authorization Drug:

1. Prescribed for an approved FDA diagnosis (as listed below):
2. Drug is being used appropriately per MCG GUIDELINES, CMS recognized compendia, authoritative medical literature, evidence-based guidelines and/or accepted standards of medical practice.
3. Member does not have any clinically relevant contraindications, or CMS/Plan exclusions, to the requested drug.
  - If the member meets all these criteria, they may be approved by the Plan for the requested drug.
  - Quantity limits and Tiering will be determined by the Plan.

### Exclusion Criteria:

N/A

### Prescriber Restrictions:

N/A

### Coverage Duration:

Approvals will be for 12 months

### FDA Indications:

#### Ocrevus

- Multiple sclerosis, Relapsing forms
- Primary progressive multiple sclerosis

#### Copaxone, Briumvi

- Multiple sclerosis, Relapsing forms

#### Zenapax

- The worldwide marketing and distribution of Zinbryta(R) (daclizumab) was voluntarily discontinued on March 2, 2018 due to safety concerns

### Off-Label Uses:

N/A

### Age Restrictions:

Safety and effectiveness of ocrelizumab have not been established in pediatric patients

### Other Clinical Considerations:

#### Ocrevus

- Active hepatitis B virus infection

**Resources:**

[https://www.micromedexsolutions.com/micromedex2/librarian/CS/9E37E3/ND\\_PR/evidencexpert/ND\\_P/evidencexpert/DUPLICATIONSHIELDSYNC/5D1C03/ND\\_PG/evidencexpert/ND\\_B/evidencexpert/ND\\_AppProduct/evidencexpert/ND\\_T/evidencexpert/PFActionId/evidencexpert.DoIntegratedSearch?SearchTerm=ocrelizumab&UserSearchTerm=ocrelizumab&SearchFilter=filterNone&navitem=searchGlobal#](https://www.micromedexsolutions.com/micromedex2/librarian/CS/9E37E3/ND_PR/evidencexpert/ND_P/evidencexpert/DUPLICATIONSHIELDSYNC/5D1C03/ND_PG/evidencexpert/ND_B/evidencexpert/ND_AppProduct/evidencexpert/ND_T/evidencexpert/PFActionId/evidencexpert.DoIntegratedSearch?SearchTerm=ocrelizumab&UserSearchTerm=ocrelizumab&SearchFilter=filterNone&navitem=searchGlobal#)

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CLINICAL USE ONLY