



Chemotherapy: Acute Myeloid Leukemia
Mylotarg (gemtuzumab ozogamicin) J9203
Prior Authorization 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.

Form section for NEW START and Continuation information, including fields for Start Date, Date Requested, and Requestor details.

MEMBER INFORMATION

Form section for Member Information, including fields for Name, ID#, and DOB.

PRESCRIBER INFORMATION

Form section for Prescriber Information, including fields for Name, Address, and Phone/Fax.

DISPENSING PROVIDER / ADMINISTRATION INFORMATION

Form section for Dispensing Provider / Administration Information, including fields for Name and Address.

PROCEDURE / PRODUCT INFORMATION

Table with 5 columns: HCPC Code, Name of Drug, Dose (Wt: kg Ht:), Frequency, End Date if known.

Form section for Chart notes and other important information.

Form section for Diagnosis: ICD10 and Description.

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

CLINICAL INFORMATION

Form section for Clinical Information, including New Start or Initial Request criteria.

Form section for Clinical Information, including Continuation Requests criteria.

ACKNOWLEDGEMENT

Form section for Acknowledgement, including Request By (Signature Required) and Date fields, and a disclaimer.

Prior Authorization Group – Oncology: Acute Myeloid Leukemia Drugs PA

Drug Name(s):

MYLOTARG

GEMTUZUMAB OZOGAMICIN

Criteria for approval of Prior Authorization Drug:

1. Prescribed for an approved FDA diagnosis (as listed below):
2. Prescribed by, or in consultation with an oncologist or other cancer specialist related to the diagnosis.
3. Drug is being used appropriately per CMS recognized compendia, authoritative medical literature, evidence-based guidelines and/or accepted standards of medical practice.
4. 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:

Cannot be prescribed for experimental or investigational use.

Prescriber Restrictions:

Oncologist or other cancer specialist

Coverage Duration:

New Start: Approval will be for 6 months

Continuation: Approval will be for 12 months

FDA Indications:

Mylotarg

- Acute myeloid leukemia, Newly-diagnosed, CD33-positive
- Acute myeloid leukemia, Relapsed or refractory, CD33-positive

Off-Label Uses:

Mylotarg

- Acute promyelocytic leukemia, FAB M3

Age Restrictions:

AML, Newly-Diagnosed, CD33-positive: 1 month or older

AML, Relapsed or refractory, CD33-positive: 2 years or older

Other Clinical Considerations:

Cancer diagnoses: Criteria as per NCCN or other FDA-approved cancer related guidelines.

Resources:

https://www.micromedexsolutions.com/micromedex2/librarian/CS/17815A/ND_PR/evidencexpert/ND_P/evidencexpert/DUPLICATIONSHIELDSYNC/BB4E53/ND_PG/evidencexpert/ND_B/evidencexpert/ND_AppProduct/evidencexpert/ND_T/evidencexpert/PFActionId/evidencexpert.GoToDashboard?docId=925216&contentSetId=100&title=Gemtuzumab+Ozogamicin&servicesTitle=Gemtuzumab+Ozogamicin&brandName=Mylotarg&UserMdxSearchTerm=Mylotarg&=null#