Oncology – Lumoxiti™ (moxetumomab pasudotox-tdfk injection for intravenous use – AstraZeneca)

APPROVAL DATE: 09/25/2019

OVERVIEW
Lumoxiti is a CD22-directed cytotoxin. Lumoxiti is produced in E. coli via recombinant DNA technology and consists of a recombinant, murine immunoglobulin variable region fused to a shortened form of the Pseudomonas endotoxin, E38. Lumoxiti binds to the CD22 antigen on the cell surface and is internalized by the B-cell which results in ADP-ribosylation of elongation factor 2, inhibition of protein synthesis, and apoptotic cell death.

Lumoxiti is indicated for the treatment of adult patients with relapsed or refractory hairy cell leukemia who received at least two prior systemic therapies, including treatment with a purine nucleoside analog. Limitations of Use: Lumoxiti is not recommended for use in patients with a creatinine clearance < 29 mL/min.

Dosing: The recommended dose of Lumoxiti is 0.04 mg/kg given as a 30 minute intravenous infusion. Lumoxiti is given on Days 1, 3 and 5 of each 28-day cycle. The recommended maximum duration of Lumoxiti therapy is 6 cycles, or until disease progression or unacceptable toxicity occurs. Dose modifications are recommended for the management of toxicities and are determined by the prescribing physician. Recommendations for dose modifications are provided in the prescribing information, are dependent on the toxicity and severity, and include withholding the dose until the toxicity clears, or discontinuing therapy with Lumoxiti.

Guidelines

POLICY STATEMENT
Prior authorization is recommended for medical benefit coverage of Lumoxiti. Approval is recommended for those who meet the Criteria and Dosing for the listed indication(s). Extended approvals are allowed if the patient continues to meet the Criteria and Dosing. Requests for doses outside of the established dosing documented in this policy will be considered on a case-by-case basis by a clinician (i.e., Medical Director or Pharmacist). All approvals are provided for the duration noted below. In cases where the approval is authorized in months, 1 month is equal to 30 days.

Because of the specialized skills required for evaluation and diagnosis of patients treated with Lumoxiti, as well as the monitoring required for adverse events and long-term efficacy, approval requires Lumoxiti to be prescribed by or in consultation with a physician who specializes in the condition being treated.

RECOMMENDED AUTHORIZATION CRITERIA
Coverage of Lumoxiti is recommended in those who meet the following criteria:
FDA-Approved Indications

1. **Hairy Cell Leukemia.** Approve for 6 months if the patient meets the following criteria (A, B, C, and D):
   
   A) Patient is ≥ 18 years of age; AND
   B) Patient has received ≥ 2 prior systemic therapies, including therapy with a purine analog.
   (Note: Purine analogs include cladribine and pentostatin); AND
   C) Patient has an estimated creatinine clearance > 30 mL/min; AND
   D) Lumoxiti is prescribed by or in consultation with an oncologist.¹³

   **Dosing.** Approve the following dosing regimen: Administer up to 0.04 mg/kg by intravenous infusion up to three times in each 28-day cycle.¹

**Conditions Not Recommended for Approval**

Lumoxiti has not been shown to be effective, or there are limited or preliminary data or potential safety concerns that are not supportive of general approval for the following conditions.

1. Coverage is not recommended for circumstances not listed in the Recommended Authorization Criteria. Criteria will be updated as new published data are available.

**REFERENCES**


**History**

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<th>Approval Date</th>
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<tbody>
<tr>
<td>New policy</td>
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<td>10/03/2018</td>
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<tr>
<td>Annual</td>
<td>No change to criteria. Removed Waste Management Sections.</td>
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