Risk Evaluation and Mitigation Strategy (REMS) Document
BLINCYTO® (blinatumomab) REMS Program

I. Administrative Information

Application Number: BLA 125557
Application Holder: Amgen, Inc.
Initial REMS Approval: 12/2014
Most Recent REMS Update: 05/2019

II. REMS Goals

The goals of the BLINCYTO REMS are to mitigate the risk of cytokine release syndrome which may be life-threatening or fatal; the risk of neurological toxicities which may be severe, life-threatening, or fatal; and the risk of preparation and administration errors associated with use of BLINCYTO by:

1. Informing healthcare providers about the risk of cytokine release syndrome which may be life-threatening or fatal
2. Informing healthcare providers about the risk of neurological toxicities which may be severe, life-threatening, or fatal
3. Informing pharmacists, who will prepare and dispense BLINCYTO, and nurses, who will administer BLINCYTO, about the risk of preparation and administration errors associated with use of BLINCYTO.

III. REMS Requirements

To inform healthcare providers about the REMS Program and the risks and safe use of BLINCYTO, Amgen must disseminate REMS communication materials according to the table below:

<table>
<thead>
<tr>
<th>Target Audience</th>
<th>Communication Materials &amp; Dissemination Plans</th>
</tr>
</thead>
<tbody>
<tr>
<td>Healthcare providers including oncologists, oncology physician assistants, oncology nurse practitioners, hematologists, oncology nurses, home healthcare oncology nurses, and infusion nurses; healthcare providers who have prescribed BLINCYTO within the previous 12 months from the approval of this REMS modification; healthcare providers who are likely to prescribe or administer BLINCYTO</td>
<td>REMS Letter: REFS Letter for Healthcare Provider or REMS Letter for Professional Societies with attachment Fact Sheet for Providers</td>
</tr>
<tr>
<td>1. Email within 60 calendar days of approval of the REMS modification (04/2019) and again 12 months later.</td>
<td></td>
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<tr>
<td>a. Send by mail within 30 calendar days of the date the first email was sent if a healthcare provider’s email address is not available or the email is undeliverable.</td>
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</tr>
<tr>
<td>b. Send a second email within 30 calendar days of the date the first email was sent if the first email is marked as unopened.</td>
<td></td>
</tr>
<tr>
<td>c. Send by mail within 30 calendar days of the date the second email was sent if the second email is marked as unopened.</td>
<td></td>
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<tr>
<td>2. Make available via a link from the BLINCYTO REMS Program Website.</td>
<td></td>
</tr>
<tr>
<td>3. Disseminate through field-based sales and medical representatives for 6 months from approval of the REMS modification (04/2019).</td>
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<tr>
<td>4. Disseminate to professional societies within 60 calendar days of the approval of the REMS modification (04/2019), again 12 months later and request the letter or content be provided to their members.</td>
<td></td>
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<tr>
<td>5. Disseminate at Professional Meetings for 6 months from approval of the REMS modification (04/2019).</td>
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<tr>
<td><strong>Fact Sheet for Providers</strong></td>
<td></td>
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1. Disseminate and prominently display at Professional Meetings where Amgen has a presence for 6 months from approval of the REMS modification (04/2019).
2. Disseminate through field-based sales and medical representatives during the initial or follow-up discussion with healthcare providers for 6 months from approval of the REMS modification (04/2019). Field-based sales or medical representatives to orally review the risk messages contained in the **Fact Sheet for Providers** during the visit with the healthcare provider. |
| **Hospital-based pharmacists and home healthcare pharmacists** | **REMS Letter:** [REMS Letter for Hospital and Home Healthcare Pharmacists](#) or [REMS Letter for Professional Societies](#) with attachment **Fact Sheet for Providers**
1. Email within 60 calendar days of the approval of the REMS modification (04/2019) and again 12 months later.
   a. Send by mail within 30 calendar days of the date the first email was sent if a pharmacist’s email address is not available or the email is undeliverable.
   b. Send a second email within 30 calendar days of the date the first email was sent if the first email is marked as unopened.
   c. Send by mail within 30 calendar days of the date the second email was sent if the second email is marked as unopened.
2. Make available via a link from the BLINCYTO REMS Program Website.
3. Disseminate through field-based sales and medical representatives for 6 months from approval of the REMS modification (04/2019).
4. Disseminate to professional societies within 60 calendar days of the approval of the REMS modification (04/2019), again 12 months later, and request the letter or content be provided to their members.
5. Disseminate at Professional Meetings for 6 months from approval of the REMS modification (04/2019). |
| **Fact Sheet for Providers** | 1. Disseminate and prominently display at Professional Meetings where Amgen has a presence for 6 months from approval of the REMS modification (04/2019).
2. Disseminate through field-based sales and medical representatives during the initial or follow-up discussion with healthcare providers for 6 months from approval of the REMS modification (04/2019). Field-based sales or medical representatives to orally review the risk messages contained in the **Fact Sheet for Providers** during the visit with the healthcare provider. |
| **REMS Program Website** | 1. Include all currently approved REMS materials, Prescribing Information, and Medication Guide.
2. Include a prominent REMS-specific link to the BLINCYTO REMS Program website. The BLINCYTO REMS Program website must not link back to the promotional product website. |
### IV. REMS Assessment Timetable

Amgen must submit REMS Assessments at 18 months, 3 years, 5 years, and 7 years from the date of the initial REMS approval (12/03/2014). 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 calendar days before the submission date for that assessment. Amgen must submit each assessment so that it will be received by the FDA on or before the due date.

### V. REMS Materials

The following materials are part of the BLINCYTO REMS:

#### Communication Materials

1. REMS Letter for Healthcare Provider
2. REMS Letter for Hospital and Home Healthcare Pharmacists
3. REMS Letter for Professional Societies
4. Fact Sheet for Providers

#### Other Materials

5. REMS Program Website
Dear Healthcare Provider:

The Food and Drug Administration (FDA) has required this safety notice as part of the BLINCYTO® REMS (Risk Evaluation and Mitigation Strategy) to highlight new risk information about cytokine release syndrome and neurological toxicities for BLINCYTO.

Please see the non-promotional REMS Fact Sheet for more detailed safety information.

**BOXED WARNING: Cytokine Release Syndrome**
- CRS, which may be life-threatening or fatal, occurred in patients receiving BLINCYTO.
- Monitor patients for signs or symptoms of CRS.
- BLINCYTO has been recently approved for the treatment of minimal residual disease (MRD)-positive B-cell precursor acute lymphoblastic leukemia (ALL) in adults and children.
  - In patients treated for MRD-positive B-cell precursor ALL, hospitalization is recommended for the first 3 days of the first cycle and the first 2 days of the second cycle.
  - In patients treated for relapsed or refractory B-cell precursor ALL, hospitalization is recommended for the first 9 days of the first cycle and the first 2 days of the second cycle.
- Administer corticosteroids for severe or life-threatening CRS.

**BOXED WARNING: Neurological Toxicities**
- In patients with ALL receiving BLINCYTO in clinical studies, neurological toxicities have occurred in approximately 65% of patients.
- Manifestations of neurological toxicity included cranial nerve disorders.

**OTHER SERIOUS RISKS: Preparation and Administration Errors**
- It is very important that the instructions for preparation (including admixing) and administration are strictly followed to minimize medication errors (including underdose and overdose).

Please see the non-promotional REMS Fact Sheet, reviewed by the FDA, and the full Prescribing Information for more detailed safety information. Additional copies of the Fact Sheet and other important information are available at: www.blincytoarms.com.

BLINCYTO is a bispecific CD19-directed CD3 T-cell engager indicated for the treatment of adults and children with 1) B-cell precursor ALL in first or second complete remission with MRD greater than or equal to 0.1% and 2) relapsed or refractory B-cell precursor ALL.

To review the Prescribing Information and Medication Guide, see links below:
- Prescribing Information
- Medication Guide

**REPORTING ADVERSE EVENTS**
You are encouraged to report negative side effects of prescription drugs to the FDA. Visit www.fda.gov/medwatch or call 1-800-FDA-1088. Healthcare Providers should report all suspected adverse events associated with BLINCYTO to the FDA or to Amgen at 1-800-77-AMGEN (1-800-772-6436).

Sincerely,

Lisa L. Bollinger, MD
Vice President, Global Patient Safety, Labeling and Pediatric Regulatory
Dear Healthcare Provider:

The Food and Drug Administration (FDA) has required this safety notice as part of the BLINCYTO® REMS (Risk Evaluation and Mitigation Strategy) to highlight new risk information about cytokine release syndrome and neurological toxicities for BLINCYTO.

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©2018, 2019 Amgen Inc. All right reserved.
Dear Pharmacist:

The Food and Drug Administration (FDA) has required this safety notice as part of the BLINCYTO® REMS (Risk Evaluation and Mitigation Strategy) to remind you of the serious risk of preparation and administration errors and highlight new risk information about cytokine release syndrome and neurological toxicities for BLINCYTO.

Please see the non-promotional REMS Fact Sheet for more detailed safety information.

Preparation and Administration Errors

- It is very important that the instructions for preparation (including admixing) and administration are strictly followed to minimize medication errors (including underdose and overdose).
- Please note that the recommended dose for BLINCYTO is by patient weight. Patients greater than or equal to 45 kg receive a fixed-dose. For patients less than 45 kg, the dose is calculated using the patient’s body surface area (BSA).

Special Considerations to Support Accurate Preparation

- Intravenous (IV) Solution Stabilizer is provided and is used to coat the prefilled IV bag prior to addition of reconstituted BLINCYTO.
- Reconstitute BLINCYTO with Sterile Water for Injection, USP, only.
- Aseptic technique must be done in a USP <797> compliant facility and strictly observed when preparing the solution for infusion since BLINCYTO does not contain antimicrobial preservatives.
- Use the specific volumes described in the admixing instructions.
- Please see the full Prescribing Information for important details on preparation and administration, including storage requirements for BLINCYTO.

BOXED WARNING: Cytokine Release Syndrome

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To review the Prescribing Information and Medication Guide, see links below:

  - Prescribing Information
  - Medication Guide

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Sincerely,

Lisa L. Bollinger, MD
Vice President, Global Patient Safety, Labeling and Pediatric Regulatory
Dear [name]:

The Food and Drug Administration (FDA) has required this safety notice as part of the BLINCYTO® REMS (Risk Evaluation and Mitigation Strategy) to be distributed to the [insert Professional Society Name] to highlight new safety information about cytokine release syndrome and neurological toxicities. Amgen requests that you distribute the information to your members, informing them about the serious risks of BLINCYTO.

BOXED WARNING: Cytokine Release Syndrome

- CRS, which may be life-threatening or fatal, occurred in patients receiving BLINCYTO.
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OTHER SERIOUS RISKS: Preparation and Administration Errors

- It is very important that the instructions for preparation (including admixing) and administration are strictly followed to minimize medication errors (including underdose and overdose).

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BLINCYTO is a bispecific CD19-directed CD3 T-cell engager indicated for the treatment of adults and children with 1) B-cell precursor ALL in first or second complete remission with MRD greater than or equal to 0.1% and 2) relapsed or refractory B-cell precursor ALL.

Sincerely,

Lisa L. Bollinger, MD
Vice President, Global Patient Safety, Labeling and Pediatric Regulatory
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**OTHER SERIOUS RISKS: Preparation and Administration Errors**
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Please see the enclosed non-promotional REMS Fact Sheet, reviewed by the FDA, and the full Prescribing Information for more detailed safety information. Additional copies of the Fact Sheet and other important information are available at: [www.blincytoREMS.com](http://www.blincytoREMS.com).
BLINCYTO is a bispecific CD19-directed CD3 T-cell engager indicated for the treatment of adults and children with 1) B-cell precursor ALL in first or second complete remission with MRD greater than or equal to 0.1% and 2) relapsed or refractory B-cell precursor ALL.

Sincerely,

Lisa L. Bollinger, MD
Vice President, Global Patient Safety, Labeling and Pediatric Regulatory
Risk of:
- Cytokine Release Syndrome (CRS), which may be life-threatening or fatal
- Neurological Toxicities, which may be severe, life-threatening or fatal
- Preparation and Administration Errors

**BOXED WARNING**

**Cytokine Release Syndrome**
- CRS, which may be life-threatening or fatal, occurred in patients receiving BLINCYTO®.
- The median time to onset of CRS is 2 days after the start of infusion.
- Manifestations of CRS include fever, headache, nausea, asthenia, hypotension, increased alanine aminotransferase, increased aspartate aminotransferase, increased total bilirubin, and disseminated intravascular coagulation (DIC).
- The manifestations of CRS after treatment with BLINCYTO overlap with those of infusion reactions, capillary leak syndrome (CLS), and hemophagocytic histiocytosis/macrophage activation syndrome (MAS).
- Using all of these terms to define CRS, in clinical trials of BLINCYTO, CRS was reported in 15% of patients with relapsed or refractory acute lymphoblastic leukemia (ALL) and in 7% of patients with minimal residual disease (MRD)-positive ALL.
- Monitor patients for signs or symptoms of these events.
  - In patients treated for MRD-positive B-cell precursor ALL, hospitalization is recommended for the first 3 days of the first cycle and the first 2 days of the second cycle.
  - In patients treated for relapsed or refractory B-cell precursor ALL, hospitalization is recommended for the first 9 days of the first cycle and the first 2 days of the second cycle.
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**Neurological Toxicities**
- In patients with ALL receiving BLINCYTO in clinical studies, neurological toxicities have occurred in approximately 65% of patients.
- Among patients that experienced a neurologic event, the median time to the first event was within the first 2 weeks of BLINCYTO treatment and the majority of events resolved.
- Manifestations of neurological toxicity included cranial nerve disorders.
- Grade 3 or higher (severe, life-threatening or fatal) neurological toxicities following initiation of BLINCYTO administration occurred in approximately 13% of patients and included encephalopathy, convulsions, speech disorders, disturbances in consciousness, confusion and disorientation, and coordination and balance disorders. The majority of events resolved following interruption of BLINCYTO, but some resulted in treatment discontinuation.

Monitor patients closely for signs and symptoms of these events and interrupt or discontinue BLINCYTO dosing. Advise outpatients on BLINCYTO to contact their healthcare professional if they develop signs or symptoms of these events.
OTHER SERIOUS RISKS:

Preparation and Administration Errors

- Preparation and administration errors have occurred with BLINCYTO treatment.
- It is very important that the instructions for preparation (including admixing) and administration are strictly followed to minimize medication errors (including underdose and overdose).
- Please note that the recommended dose for BLINCYTO is by patient weight. Patients greater than or equal to 45 kg receive a fixed-dose. For patients less than 45 kg, the dose is calculated using the patient’s body surface area (BSA).
- See Dosage and Administration section of Prescribing Information for detailed safety information.

MORE INFORMATION

For detailed information regarding BLINCYTO including storage, preparation, and administration, it is essential that you read the Prescribing Information for BLINCYTO.

INDICATION

BLINCYTO is a bispecific CD19-directed CD3 T-cell engager indicated for the treatment of adults and children with 1) B-cell precursor ALL in first or second complete remission with MRD greater than or equal to 0.1% and 2) relapsed or refractory B-cell precursor ALL.

WHAT IS THE BLINCYTO® REMS?

A REMS (Risk Evaluation and Mitigation Strategy) is a program required by the Food and Drug Administration (FDA) to manage known or potential serious risks associated with a drug product. The FDA has determined that a REMS is necessary to ensure that the benefits of BLINCYTO outweigh its risks. The purpose of the BLINCYTO REMS is to inform Healthcare Providers of the risks of serious neurological toxicities, cytokine release syndrome, and preparation and administration errors. This Fact Sheet is required by the FDA as part of the BLINCYTO REMS program.

Please visit www.blincytorems.com for further information and resources.

This Fact Sheet does not contain the complete safety profile for BLINCYTO. Please refer to the full Prescribing Information, including BOXED WARNINGS and Medication Guide.

REPORTING ADVERSE EVENTS

You are encouraged to report negative side effects of prescription drugs to the FDA. Visit www.fda.gov/medwatch or call 1-800-FDA-1088. Healthcare Providers should report all suspected adverse events associated with BLINCYTO to the FDA or to Amgen at 1-800-77-AMGEN (1-800-772-6436).
BLINCYTO® (blinatumomab) Risk Evaluation and Mitigation Strategy (REMS)

What is the BLINCYTO® REMS?
A Risk Evaluation and Mitigation Strategy (REMS) is a program to manage known or potential serious risks associated with a drug product and is required by the Food and Drug Administration (FDA) to ensure that the benefits of the drug outweigh its risks.

The purpose of the BLINCYTO REMS is to inform Healthcare Providers about the following serious risks:

BOXED WARNING: Cytokine Release Syndrome
- Cytokine Release Syndrome (CRS), which may be life-threatening or fatal, occurred in patients receiving BLINCYTO.
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Monitor patients closely for signs and symptoms of these events and interrupt or discontinue dosing of BLINCYTO. Advise outpatients on BLINCYTO to contact their healthcare professional if they develop signs or symptoms of these events.

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BLINCYTO Fact Sheet:
A non-promotional REMS Fact Sheet reviewed by the FDA, with more detailed information on the serious risks associated with BLINCYTO is available in the "Materials for Healthcare Providers" section above.

INDICATION:
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You are encouraged to report negative side effects of BLINCYTO to Amgen at 1-800-77-AMGEN (1-800-772-6436) and/or the FDA at www.fda.gov/medwatch or call 1-800-FDA-1088

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