Executive Summary

**Purpose:** Ensure appropriate utilization and control of CAR-T Cell Therapies

**Why Issue Selected:** CAR-T Cell Therapy is a form of immunotherapy where a patient’s T-cells are collected and genetically engineered to produce chimeric antigen receptors (CAR) on the cell surface, allowing the modified T-cells to recognize an antigen on target cancer cells.

Approved by the FDA in August 2017, Kymriah® (tisagenlecleucel) is indicated for the treatment of B-cell precursor acute lymphoblastic leukemia (ALL) that is refractory or in second or later relapse in pediatric and young adult patients (up to 25 years of age); Kymriah is also indicated for the treatment of adult patients with relapsed or refractory large B-cell lymphoma after two or more lines of systemic therapy including diffuse large B-cell lymphoma (DLBCL) not otherwise specified, high grade B-cell lymphoma and DLBCL arising from follicular lymphoma. In May 2022, Kymriah was approved for the treatment of adults with relapsed or refractory follicular lymphoma (FL) after two or more lines of systemic therapy.

FDA approved in October 2017, Yescarta® (axicabtagene ciloleucel) is indicated for the treatment of adult patients with relapsed or refractory large B-cell lymphoma after one or more lines of systemic therapy, including diffuse large B-cell lymphoma (DLBCL) not otherwise specified, primary mediastinal large B-cell lymphoma, high grade B-cell lymphoma, and DLBCL arising from follicular lymphoma. In March 2021, Yescarta received another FDA indication for adult patients with relapsed or refractory follicular lymphoma (FL) after two or more lines of systemic therapy.

FDA approved in July 2020, Tecartus® (brexucabtagene autoleucel) is indicated for the treatment adult patients with relapsed or refractory mantle cell lymphoma (MCL) and for adult patients with relapsed or refractory B-cell precursor acute lymphoblastic leukemia (ALL). Although previous systemic therapy for MCL is not noted in the indication, NCCN Guidelines state Tecartus is recommended for the treatment of adult patients with relapsed or refractory MCL only after chemoimmunotherapy and BTK inhibitor therapy.

Approved by the FDA in February 2021, Breyanzi® (lisocabtagene maraleucel) is indicated for the treatment of adult patients with relapsed or refractory large B-cell
lymphoma after one or more lines of systemic therapy, including diffuse large B-cell lymphoma (DLBCL) not otherwise specified (including DLBCL arising from indolent lymphoma), high-grade B-cell lymphoma, primary mediastinal large B-cell lymphoma, and follicular lymphoma grade 3B.

FDA approved in March 2021, Abecma® (idecabtagene vicleucel) is indicated for treatment of adult patients with relapsed or refractory multiple myeloma after four or more prior lines of therapy, including an immunomodulatory agent, a proteasome inhibitor, and an anti-CD38 monoclonal antibody.

All CAR-T cell therapies have boxed warnings concerning Cytokine Release Syndrome and neurologic toxicities with a REMS program. Abecma has additional boxed warnings for Hemophagic Lymphohistiocytosis/Macrophage Activation Syndrome (HLH/MAS) and prolonged cytopenias.

All requests for therapy will be reviewed by a Clinical Consultant.

Program-Specific Information:

<table>
<thead>
<tr>
<th>Drug</th>
<th>Cost per infusion</th>
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</thead>
<tbody>
<tr>
<td>ABECMA (IDECABTAGENE VICLEUCEL)</td>
<td>$419,500.00 WAC</td>
</tr>
<tr>
<td>BREYANZI (LISOCABTAGENE MARALEUCEL)</td>
<td>$410,300.00 WAC</td>
</tr>
<tr>
<td>KYMRIAH (TISAGENLECLEUCEL) – DLBCL</td>
<td>$373,000.00 WAC</td>
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<tr>
<td>KYMRIAH (TISAGENLECLEUCEL) – PED ALL</td>
<td>$475,000.00 WAC</td>
</tr>
<tr>
<td>TECARTUS (BREXUCABTAGENE AUTOLEUCEL)</td>
<td>$399,000.00 WAC</td>
</tr>
<tr>
<td>YESCARTA (AXICABTAGENE CILOLEUCEL)</td>
<td>$399,000.00 WAC</td>
</tr>
</tbody>
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Type of Criteria: ☒ Increased risk of ADE ☐ Preferred Drug List
☒ Appropriate Indications ☒ Clinical Edit

Data Sources: ☐ Only Administrative Databases ☒ Databases + Prescriber-Supplied

Setting & Population

- Drug class for review: CAR-T Cell Therapies
- Age range: All appropriate MO HealthNet participants

Approval Criteria

- Prescribed by or in consultation with an oncologist, hematologist, or other specialist in the treated disease state AND
- Participant is currently not pregnant AND
- For Breyanzi:
  - Participant is aged ≥ 18 years AND
  - Documented diagnosis of relapsed or refractory large B-cell lymphoma, including diffuse large B-cell lymphoma (DLBCL) not otherwise specified (including DLBCL arising from indolent lymphoma), high-grade B-cell lymphoma, primary mediastinal large B-cell lymphoma, and follicular lymphoma grade 3B AND
  - Documentation of one or more previous lines of therapy for treated diagnosis
- For Kymriah:
  - Participant aged ≤ 25 years AND documented diagnosis of B-cell precursor acute lymphoblastic leukemia (ALL) that is refractory or in second or later relapse OR
  - Participant aged ≥ 18 years AND documented diagnosis of relapsed or refractory (r/r) large B-cell lymphoma after two or more lines of systemic therapy including diffuse large B-cell lymphoma
(DLBCL) not otherwise specified, high grade B-cell lymphoma and DLBCL arising from follicular lymphoma OR
  o Participant aged ≥ 18 years AND documented diagnosis of relapsed or refractory follicular lymphoma (FL) after two or more lines of systemic therapy AND
  o Documentation of two or more previous lines of systemic therapy for treated diagnosis
• For Yescarta:
  o Documented diagnosis of large B-cell lymphoma that is refractory to first-line chemoimmunotherapy or that relapses within 12 months of first-line chemoimmunotherapy OR
  o Documented diagnosis of relapsed or refractory large B-cell lymphoma after two or more lines of systemic therapy, including diffuse large B-cell lymphoma (DLBCL) not otherwise specified, primary mediastinal large B-cell lymphoma, high grade B-cell lymphoma, and DLBCL arising from follicular lymphoma OR
  o Documented diagnosis of relapsed or refractory follicular lymphoma (FL) AND
  o Participant aged ≥ 18 years AND
  o Documentation of one or more previous lines of systemic therapy for treated diagnosis
• For Tecartus:
  o Participant aged ≥ 18 years AND
  o Documented diagnosis of relapsed or refractory mantle cell lymphoma (MCL) AND
  o Documentation of two or more previous lines of systemic therapy for MCL, including chemoimmunotherapy and BTK inhibitor therapy OR
  o Documented diagnosis of relapsed or refractory B-cell precursor acute lymphoblastic leukemia (ALL)
• For Abecma:
  o Participant is aged ≥ 18 years AND
  o Documented diagnosis of relapsed or refractory multiple myeloma AND
  o Documentation of four or more previous lines of therapy, including an immunomodulatory agent, a proteasome inhibitor, and an anti-CD38 monoclonal antibody

Denial Criteria

• Therapy will be denied if all approval criteria are not met
• Previous CAR T Cell Therapy
• Participant has an active infection or inflammatory disorder

Billing Information

• The treating prescriber/facility may choose the method of administration most appropriate for the participant. The options available are:
  o Outpatient FFS Participant: MHD FFS completes Prior Authorization for the drug, claim paid according to the current MHD drug reimbursement. All other expenses are paid for on a FFS basis.
    ▪ The following are items that would be expected to be billed to MHD FFS (this list is used as an example and is not meant to be all inclusive): Apheresis, chemotherapy, cell engineering, antigen receptor T-Cell therapy (CAR-T), infusion, leukapheresis, preparative therapy. If any of these are included in the cost of the drug, they cannot be billed separately to MHD.
    ▪ If the patient should require hospitalization due to cytokine release syndrome (CRS) or neurotoxicities after the CAR-T infusion, inpatient pre-certification is required, and payment will be made at the applicable rate for the facility as determined by established MHD regulations.
  o Outpatient MCO Participant: MHD FFS completes Prior Authorization for the drug, drug claim paid according to the current MHD drug reimbursement. All other expenses are paid for by the MCO.
The following are items that would be expected to be billed to the participant MCO plan (this list is used as an example and is not meant to be all inclusive): Apheresis, chemotherapy, cell engineering, antigen receptor T-Cell therapy (CAR-T), infusion, leukapheresis, preparative therapy. If any of these are included in the cost of the drug, they cannot be billed separately to the MCO.

If the patient should require hospitalization due to cytokine release syndrome (CRS) or neurotoxicities after the CAR-T infusion, coverage of the hospital stay is the MCO plan responsibility and payment will be made at the applicable MCO rate for the facility as determined by established MCO provider agreement.

- **Inpatient FFS Participant**: MHD FFS completes Prior Authorization of the drug, MHD FFS covers the cost of the drug according to the current MHD drug reimbursement in addition to the hospital per diem. Medical provider costs are billed separately to FFS.
  - Pre-certification of the inpatient hospitalization is required for the CAR-T infusion; should the medical provider feel it is medically necessary to administer chemotherapy inpatient prior to CAR-T infusion, pre-certification is also required for those hospital days.
  - MHD FFS will reimburse the facility at the per diem rate during the participant’s inpatient hospitalization. All other expenses will be reimbursed only if they are performed prior to the participant’s inpatient stay and if they are not included in the cost of the drug. Examples of expenses expected prior to the inpatient hospitalization include, this list is not meant to be considered all inclusive: Apheresis, chemotherapy, cell engineering, leukapheresis, and preparative therapy.
  - If the patient should require subsequent hospitalization after the initial stay involving the therapy due to cytokine release syndrome or neurotoxicities, payment will be made at the applicable rate for the facility as determined by established MHD regulations.

- **Inpatient MCO Participant**: MHD FFS completes Prior Authorization of the drug, MHD FFS covers the cost of the drug according to the current MHD drug reimbursement. The MCO is responsible for the hospital stay and any medical provider costs.
  - The MCO will reimburse the facility at the per diem rate during the participant’s inpatient hospitalization. All expenses, with the exception for the cost of the drug, are the responsibility of the MCO plan.
  - If the patient should require subsequent hospitalization after the initial stay involving the therapy, payment will be made at the applicable MCO per diem rate for the facility as determined by established MCO provider agreement.

**The following apply to all scenarios above:**

- Failure to get the prior authorization for the CAR-T therapy prior to administration will result in no payment for the CAR-T therapy.
- During the prior authorization for the CAR-T therapy the provider must determine if the participant will receive the CAR-T therapy as an inpatient or outpatient service.
- There will be no additional $100,000 reimbursement for hospitalization or treatment in any of the above scenarios. The accompanying treatment will be paid for at the agreed upon provider rates.
- The CAR-T Therapy will only be paid for by MHD when the facility is approved by the manufacturer to administer the CAR-T therapy.
- All pre-therapy assessment and medical services, physician’s charges, and outpatient follow-up care must be billed separately and will be reimbursed at current applicable MHD fee schedule or inpatient per diem rates.
- For experimental use of CAR-T therapy, when the manufacturer has provided the CAR-T drug through compassionate use or as part of a clinical trial, the facility will not be reimbursed for the drug by MHD FFS nor the MCO. All other medically necessary covered services associated with the CAR-T therapy, which are not part of the clinical trial, may be reimbursed according to the process selected by the treating prescriber and outlined above, according to the current hospital manual.
Laboratory Results: [ ] Progress Notes: [ ] MedWatch Form: [ ] Other: [X]

Disposition of Edit

Denial: Exception code “0682” (Clinical Edit)
Rule Type: CE

Default Approval Period

2 months

References

- YESCARTA® (axicabtagene ciloleucel) [package insert]. Santa Monica, CA: Kite Pharma, Inc.; April 2022.
- ABECMA® (idecabtagene vilociecel) [package insert]. Summit, NJ: Celgene Corporation, a Bristol-Myers Squibb Company; March 2021.