

MEDICAL POLICY STATEMENT INDIANA MARKETPLACE					
Policy Name		Policy Number	Date Effective		
CAR-T medications – Kymriah (tisagenlecleucel)		MM-1122	07/01/2021-05/31/2022		
Policy Type					
MEDICAL	Administrative	Pharmacy	Reimbursement		
clinical guidelines, nation standards, and published health care services or su without which the patient a body organ or part, or s area, are the lowest cost necessary services also in	ally recognized utilization and te I MCO clinical policy guidelines. pplies that are proper and nece can be expected to suffer prolo ignificant pain and discomfort. T alternative, and are not provide	chnology assessment guidel Medically necessary services ssary for the diagnosis or trea nged, increased or new morb 'hese services meet the stan d mainly for the convenience a any Evidence of Coverage of	terature based on and supported by ines, other medical management industry s include, but are not limited to, those atment of disease, illness, or injury and idity, impairment of function, dysfunction of dards of good medical practice in the local of the member or provider. Medically documents, Medical Policy Statements,		

Medical Policy Statements prepared by CareSource and its affiliates do not ensure an authorization or payment of services. Please refer to the plan contract (often referred to as the Evidence of Coverage) for the service(s) referenced in the Medical Policy Statement. If there is a conflict between the Medical Policy Statement and the plan contract (i.e., Evidence of Coverage), then the plan contract (i.e., Evidence of Coverage) will be the controlling document used to make the determination.

According to the rules of Mental Health Parity Addiction Equity Act (MHPAEA), coverage for the diagnosis and treatment of a behavioral health disorder will not be subject to any limitations that are less favorable than the limitations that apply to medical conditions as covered under this policy.

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B. Background

Chimeric antigen receptor T cell therapy (CAR-T) is an autologous T-cell immunotherapy. The member's own T lymphocytes are genetically modified with a gene that encodes a CAR-T to the T cells which can then target the lymphoma cells. Once the member's T cells are modifier, the T cells are infused back into the member.

CAR-T is associated with severe complications and may be life-threatening. These complications include cytokine release syndrome and neurological toxicities. Therefore, CAR-T therapy administration should be based on clinical benefits, potential long-term disease control, and toxicity.

This policy is to define medically necessary criteria for Kymriah.

C. Definitions

• **Risk Evaluation and Mitigation Strategy (REMS)** - A drug safety program that the U.S. Food and Drug Administration (FDA) can require for certain medications with serious safety concerns to help ensure the benefits of the medication outweigh its risks. REMS are designed to reinforce medication use behaviors and actions that support the safe use of that medication.

D. Policy

- I. Kymriah
 - A. Kymriah may be approved for 3 months.
 - B. Kymriah will not be reauthorized for continued therapy.
 - C. Kymriah is limited to one infusion per lifetime.
- II. Kymriah is a non-preferred product and will only be considered for coverage under the medical benefit when either of the following medically necessary criteria are met:
 - A. Diagnosis of Acute lymphoblastic Leukemia and all of the following criteria:
 - 1. Member is 1 to 25 years of age;
 - 2. Member has a diagnosis of relapsed or refractory B-cell ALL defined by one of the following:
 - **a.** Second or greater relapse;
 - b. Relapse after allogeneic stem cell transplantation (SCT);
 - **c.** Primary refractory as defined by not achieving a complete remission (CR) after 2 cycles of a standard chemotherapy regimen or chemorefractory as defined by not achieving a CR after 1 cycle of standard chemotherapy for relapsed leukemia;
 - **d.** Philadelphia chromosome positive (Ph+) all and intolerant to or have failed 2 lines of tyrosine kinase inhibitor (TKI) therapy [e.g. imatinib mesylate (Gleevec), dasatinib (Sprycel)];or
 - e. Ineligible for allogeneic SCT;





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- 4. Documentation of CD19 tumor expression;
- Bone marrow with ≥ 5% lymphoblasts by morphologic assessment; Member has been pre-screened for hepatitis B virus (HBV), hepatitis C virus (HCV), and human immunodeficiency virus (HIV) prior to collection of cells (negative results must be submitted);
- 6. Healthcare facility/provider has enrolled in the Kymriah REMS program; and
- 7. Member does not have any of the following:
 - a. Prior gene or CAR-T cell therapy; or
 - b. Life expectancy less than 12 weeks.
- OR
- B. Diagnosis of Large B-cell lymphoma and all of the following criteria:
 - 1. Member is 18 years of age or older;
 - 2. Member has a diagnosis of relapsed or refractory large B-cell lymphoma including one of the following:
 - a. Diffuse large B-cell lymphoma (DLBCL) not otherwise specified;
 - b. High grade B-cell lymphoma; or
 - c. DLBCL arising from follicular lymphoma;
 - 3. Member has received 2 or more lines of chemotherapy, including rituximab and anthracycline, and relapsed following autologous hematopoietic stem cell transplantation (HSCT) or is not eligible for HSCT;
 - 4. Member does not have any of the following:
 - a. Active central nervous system malignancy involvement;
 - b. Prior allogenic HSCT;
 - c. Prior CAR-T therapy (e.g., Yescarta);or
 - d. Life expectancy less than 12 weeks;
 - Member has been pre-screened for hepatitis B virus (HBV), hepatitis C virus (HCV), and human immunodeficiency virus (HIV) prior to collection of cells (negative results must be submitted); and
 - 6. Healthcare facility/provider has enrolled in the Kymriah REMS program.
- III. CareSource considers Kymriah not medically necessary for the treatment of disease states not in this document.
- E. Conditions of Coverage NA
- F. Related Policies/Rules

Evidence of Coverage and Health Insurance Contract Indiana Pharmacy Policy Statement: CAR-T medications CAR-T medications – Yescarta CAR-T medications – Tecartus Pharmacy Policy Statement: Kymriah (tisagenlecleucel)





G. Review/Revision History

	DATE	ACTION
Date Issued	04/14/2021	
Date Revised		
Date Effective	07/01/2021	
Date Archived		This Policy is no longer active and has been archived. Please note that there could be other Policies that may have some of the same rules incorporated and CareSource reserves the right to follow CMS/State/NCCI guidelines without a formal documented Policy.

H. References

- 1. Kymriah [package insert]. East Hanover, NJ; Novartis Pharmaceuticals Corp., May 2018.
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- 3. ClinicalTrials.gov. Identifier NCT02228096. Study of Efficacy and Safety of CTL019 in Pediatric ALL Patients. Retrieved September 22, 2020 from www.clinicaltrials.gov
- 4. Schuster SJ, et al. Primary analysis of Juliet: a global, pivotal, phase 2 trial of CTL019 in adult patients with relapsed or refractory diffuse large B-cell lymphoma. Blood. 2017;130(s1):577 [Abstract 577 from 2017 ASH annual meeting].
- 5. Maude SL, et al. Tisagenlecleucel in Children and Young Adults with B-Cell Lymphoblastic Leukemia. N Engl J Med. 2018;378(5):439-448. [PubMed 29385370]
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The Medical Policy Statement detailed above has received due consideration as defined in the Medical Policy Statement Policy and is approved.

Independent medical review – 08/2020

