

## **Pharmacy Prior Authorization Form** Fax completed form to: 877.974.4411 toll free, or 616.942.8206 □ Commercial (Traditional) □ Commercial Individual (Optimized) This form applies to: Medicaid Urgent (life threatening) Non-Urgent (standard review) This request is: Urgent means the standard review time may seriously jeopardize the life or health of the patient or the patient's ability to regain maximum function. Yescarta (axicabtagene ciloleucel) Member Last Name: First Name: DOB: \_\_\_\_\_ Gender: Primary Care Physician: Phys. Phone: Phys. Fax: Requesting Physician: Physician Address: Physician NPI: Contact Name: \_\_\_\_\_ Provider Signature: **Product Information** Dose: \_\_\_\_\_ Dose Frequency:\_\_\_\_ Drug product: ☐ Yescarta intravenous suspension Start date (or date of next dose): Date of last dose (if applicable): Date of next dose: Place of administration: Physician's office ☐ Outpatient infusion NPI:\_\_\_\_\_ Fax:\_\_\_\_ Facility: ☐ Home infusion NPI: \_\_\_\_\_\_ Fax:\_\_\_\_\_ Facility: Billing: ☐ Physician to buy and bill ☐ Facility to buy and bill ☐ Specialty Pharmacy NPI: Fax: Pharmacy: ICD-10 Diagnosis code(s): **Drug cost information** The wholesale acquisition cost for one dose of Yescarta<sup>™</sup> is \$373,000. **Precertification Requirements**

Before this drug is covered, documentation must be submitted to support that the patient meets all of the following requirements:

- 1. Diagnosis of:
  - a. Diffuse Large B-cell Lymphoma (DLBCL) unspecified or from follicular lymphoma
  - b. Primary mediastinal large B-cell lymphoma
  - c. High grade B-cell lymphoma
- 2. Must have previously used two or more lines of systemic therapy



3. Patient has or will receive lymphodepleting chemotherapy with fludarabine and cyclophosphamide intravenously on the fifth, fourth, and third day prior to Yescarta infusion

**NOTE:** Coverage of Yescarta is limited to the dosing listed in the FDA-approved label. Patients greater than 50 kg will be allowed no more than  $2.5 \times 10^8$  CAR-positive viable T cells. Patients weighting 50 kg or less will be authorized for weight-based dosing of  $0.2 - 5 \times 10^6$  CAR-positive viable T cells.

**Note:** Authorization for indications, dosing, or a route of administration not approved by the Food and Drug Administration (FDA) or recognized in CMS-accepted compendia (e.g. DrugDex, AHFS, U.S. Pharmacopeia, and also Clinical Pharmacology for oncology indications only) require supporting evidence for coverage. Please provide two published peer-reviewed literature articles supporting the appropriateness of the drug, the dosing of the drug, or the route of administration to be used for the identified indication.

	☐ Primary media ☐ High grade B-α ☐ Other – the pa	s drug being reques B-cell Lymphoma (DL stinal large B-cell lyn cell lymphoma tient's condition is:	sted for? LBCL) unspecified or from foll	. ,
В.	Has the patient tried two or lines of systemic therapy?			
	☐ Yes <b>Drug</b>	Dose	Dates	Outcome
		or use:		
C.	☐ Yes (please su	bmit documentation	chemotherapy with IV fludation if already completed or treatn	

## **Additional information**

**Note:** Prior to approving coverage of Yescarta<sup>™</sup>, a member's plan documents must include coverage for gene therapies with the applicable rider.