DEPARTMENT: Pharmacy,	DOCUMENT NAME:	
Medical Directors, Claims	Idecabtagene Vicleucel (Abecma)	
PAGE: 1 of 6	REPLACES DOCUMENT:	
APPROVED DATE:	RETIRED:	
EFFECTIVE DATE: 10/18/2021	REVIEWED/REVISED:	
PRODUCT TYPE: STAR, STAR	REFERENCE NUMBER: TX.PHAR.94	
Health, STAR Kids, STAR Plus,		
CHIP, CHIP Perinate		

SCOPE:

Superior HealthPlan Pharmacy Department, Medical Directors, Claims

POLICY:

It is the policy of Superior HealthPlan to follow state guidance for medical necessity review of idecabtagene vicleucel (Abecma).

Consistent with the regulation at 42 CFR Section 438.210 and 42 CFR Section 457.1230(d), services covered under managed care contracts, including clinician-administered drugs, must be furnished in an amount, duration, and scope that is no less than the amount, duration, and scope for the same services specified in the state plan.

While MCOs may place appropriate limits on drugs, MCOs may not use a standard for determining medical necessity that is more restrictive than what is used in the state plan, i.e., developed by the Vendor Drug Program. For example, if a member is denied a clinician administered drug in managed care because of the MCO's prior authorization criteria, but would have received the drug under the criteria specified in the state plan, then the MCO's prior authorization criteria would violate the amount, duration, and scope requirements cited above. HHSC intends to amend the Managed Care Contracts at the next opportunity to include this requirement. This same standard applies to CHIP formulary and CAD coverage.

Refer to the Outpatient Drug Services Handbook of the Texas Medicaid Provider Procedure Manual for more details on the clinical policy and prior authorization requirements.

All determinations will be performed by a Superior HealthPlan medical director. A pharmacy clinician will review the prior authorization request and make a recommendation to the medical director but will not make the ultimate determination on any case.

Additionally, this medication is a precision drug. Centene's Precision Drug Action Committee (PDAC) creates a standardized approach for Centene to

i oncy and i loccuuic		
DEPARTMENT: Pharmacy,	DOCUMENT NAME:	
Medical Directors, Claims	Idecabtagene Vicleucel (Abecma)	
PAGE: 2 of 6	REPLACES DOCUMENT:	
APPROVED DATE:	RETIRED:	
EFFECTIVE DATE: 10/18/2021	REVIEWED/REVISED:	
PRODUCT TYPE: STAR, STAR	REFERENCE NUMBER: TX.PHAR.94	
Health, STAR Kids, STAR Plus,		
CHIP, CHIP Perinate		

manage precision drugs and the associated costs for their administration, prior to members presenting with a request for one of these agents. All precision drug requests or potential requests must be reported to the PDAC for tracking, regardless of whether agents are carved out, passed through, etc. All precision drug medical necessity determinations will be supported by PDAC UM recommendation, utilizing specialist input as directed and allowed by turnaround times.

In addition, the procedure code C9081 (used for Abecma) will be limited to once per lifetime, by any provider. If the code is updated in the future it will still be limited to once per lifetime.

Certified healthcare facilities must enroll and comply with the Risk Evaluation and Mitigation Strategies (REMS) requirements for this drug.

BACKGROUND:

Description:

Idecabtagene Vicleucel (Abecma) is an anti-B cell maturation antigen (BCMA) chimeric antigen receptor (CAR) T-cell immunotherapy.

FDA APPROVED INDICATION(S)

Abecma is indicated for the 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.

FORMULATIONS:

Available as a single-dose unit infusion bag: frozen suspension of genetically modified autologous T-cells labeled for the specific recipient.

- 50 mL infusion bag (NDC: 59572-0515-01)
- 250mL infusion bag (NDC: 59572-0515-02)
- 500mL infusion bag (NDC: 59572-0515-03)

PROCEDURE:

i oncy and i loccuuic		
DEPARTMENT: Pharmacy,	DOCUMENT NAME:	
Medical Directors, Claims	Idecabtagene Vicleucel (Abecma)	
PAGE: 3 of 6	REPLACES DOCUMENT:	
APPROVED DATE:	RETIRED:	
EFFECTIVE DATE: 10/18/2021	REVIEWED/REVISED:	
PRODUCT TYPE: STAR, STAR	REFERENCE NUMBER: TX.PHAR.94	
Health, STAR Kids, STAR Plus,		
CHIP, CHIP Perinate		

Provider <u>must</u> submit documentation (which may include office chart notes and lab results) supporting that member has met all approval criteria.

I. Initial Approval Criteria:

- 1. A medical director is required to review and approve or deny all requests. A pharmacist will make a recommendation on the prior authorization but ultimate determination will be made by the medical director only.
- 2. Medical necessity determinations will be supported by PDAC UM recommendation. The SHP pharmacy clinician will review the UM recommendation with the prior authorization request for clinical appropriateness and make a recommendation to the SHP medical director but will not make the ultimate determination on any case.
- 3. Age \geq 18 years
- 4. The patient must have a histologically confirmed diagnosis relapse or refractory multiple myeloma (diagnosis code: C9000 and C9002)
- 5. The patient must have received four or more prior lines of the following therapies before treatment with idecabtagene vicleucel:
 - An immunomodulatory agent
 - A proteasome inhibitor
 - An anti-CD38 monoclonal antibody
- 6. The patient does not have primary central nervous system lymphoma/disease.
- 7. The patient does not have an active infection or inflammatory disorder.
- 8. Patient has not received prior CAR-T therapy.

Policy and Procedure		
DOCUMENT NAME:		
Idecabtagene Vicleucel (Abecma)		
REPLACES DOCUMENT:		
RETIRED:		
REVIEWED/REVISED:		
REFERENCE NUMBER: TX.PHAR.94		

9. If the facility is non-PAR the medical director will redirect to a PAR provider. On a case by case basis, said medical director may make an exception outside of a PAR provider but will require a single case agreement (SCA). Once the case is determined, the pharmacy team via pharmacy management will work with the SCA team to assist on the SCA. This should be the exception and not the rule as a PAR facility/provider is preferable. The pharmacist supporting the medical director will contact pharmacy management to start the SCA process.

Approval duration: Only 1 dose per lifetime will be provided on this drug regardless of Provider.

II. Dosage and Administration

Indication	Dosing Regimen	Maximum Dose
Multiple myeloma	Single IV infusion; target dose: 300 -	460×10^6 CAR-positive
	460×10^6 CAR-positive T cells	viable T cells

REFERENCES:

- 1. Abecma Prescribing Information. Summit, NJ: Celgene Corporation; March 2021. Available at: https/www.abecma.com
- 2. Outpatient Drug Services Handbook of the Texas Medicaid Provider Procedure Manual, Accessed October 2021

DEFINITIONS/Abbreviations:

MCO: Managed Care Organization HHSC: Health and Human Services Commission SCA: Single Case Agreement PDAC: Precision Drug Action Committee UM: Utilization Management

DEPARTMENT: Pharmacy,	DOCUMENT NAME:	
Medical Directors, Claims	Idecabtagene Vicleucel (Abecma)	
PAGE: 5 of 6	REPLACES DOCUMENT:	
APPROVED DATE:	RETIRED:	
EFFECTIVE DATE: 10/18/2021	REVIEWED/REVISED:	
PRODUCT TYPE: STAR, STAR	REFERENCE NUMBER: TX.PHAR.94	
Health, STAR Kids, STAR Plus,		
CHIP, CHIP Perinate		
	•	

SHP: Superior HealthPlan

REVISION LOG

REVISION	DATE

POLICY AND PROCEDURE APPROVAL

Karen Tadlock, V.P., Pharmacy Operations	Approval on file
Dr. David Harmon, Sr. V.P., Chief Medical Officer	Approval on file
Pharmacy & Therapeutics Committee	Approval on file

NOTE: The electronic approval retained in RSA Archer, Centene's P&P management software, is considered equivalent to a physical signature.