

Title: Tecvayli (teclistamab-cqyv)	Division: Medical Management Department: Utilization Management
Approval Date: 1/31/2023	LOB: Medicaid, HIV SNP, HARP, CHP, Medicare, UltraCare, MetroPlus Gold, Goldcare I&II, Essential Plan, QHP
Effective Date: 1/31/2023	Policy Number: UM-MP345
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1. POLICY DESCRIPTION:

Medical Oncology – Anti CD-3; Anti-BCMA; Bispecific T-Cell Engager; Mab, Tecvayli (teclistamab-cqyv)

2. RESPONSIBLE PARTIES:

Medical Management Administration, Utilization Management, Integrated Care Management, Pharmacy, Claim Department, Providers Contracting.

3. DEFINITIONS:

Tecvayli (teclistamab-cqyv) is an anti-neoplastic therapy that binds to the CD3 receptor on T-cells and B-cell maturation antigen (BCMA) on the surface of multiple myeloma cells. This results in T-cell activation and the release of various inflammatory cytokines, which results in the lysis of BCMA-expressing multiple myeloma cells. Tecvayli is currently used for the treatment of adult patients with multiple myeloma that is refractory or has relapsed after using at least 4 lines of therapy including a proteasome inhibitor, immunomodulatory agent and an anti-CD38 monoclonal antibody.

4. POLICY:

Tecvayli will be considered medically necessary once the following coverage criteria is met:

INITIAL REQUEST:

- A. Member is 18 years of age or older **AND**
- B. Member has a diagnosis of relapsed or refractory multiple myeloma **AND**
- C. The member has received treatment with at least four prior lines of therapy, including at least one drug from each of the following categories:
 - a. Proteasome inhibitor [e.g., bortezomib (Velcade), carfilzomib (Kyprolis)]
 - b. Immunomodulatory agent [e.g., lenalidomide (Revlimid), pomalidomide (Pomalyst), thalidomide (Thalomid)]
 - c. Anti-CD38 monoclonal antibody [e.g., daratumumab (Darzalex)] **AND**
- D. Member has an Eastern Cooperative Oncology Group (ECOG) score < 2 **AND**
- E. Member meets **ALL** of the following laboratory criteria:
 - a. Creatinine clearance ≥ 40 mL/min
 - b. Hemoglobin (Hgb) ≥ 8 g/dL
 - c. Platelets (PLT) ≥ 75 x 10⁹/L
 - d. Absolute Neutrophil Count (ANC) ≥ 1.0 x 10⁹/L
 - e. AST and ALT ≤ 3.0 times upper limit of normal

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- f. Total bilirubin \leq 2.0 times upper limit of normal (unless due to Gilbert disease direct bilirubin must be \leq 1.5 times upper limit of normal)
- g. Corrected Serum Calcium \leq 14mg/dL or free ionized calcium $>$ 6.5 mg/dL **AND**
- F. Member has a negative serum pregnancy test prior to therapy if they are a women of childbearing potential **AND**
- G. Member agrees to use effective contraception during the course of treatment and 5 months after the last dose of Tecvayli **AND**
- H. Member does not have active central nervous system (CNS) involvement including clinical signs of meningeal involvement of multiple myeloma **AND**
- I. Member does not require oxygen supplementation during therapy **AND**
- J. Member does not have a cardiac disease that can adversely affect therapy **AND**
- K. Member does not have an active inflammatory disorder **AND**
- L. Member does not have an active uncontrolled infection including human immunodeficiency virus (HIV), Hepatitis B or C and Cytomegalovirus (CMV) **AND**
- M. Member has not received autologous stem cell transplantation \leq 12 weeks prior to the first dose of Tecvayli **AND**
- N. Member does not have an active autoimmune disease including graft versus host disease requiring to be on immunosuppressive agents **AND**
- O. Member does not have plasma cell leukemia, Waldenström’s macroglobulinemia, POEMS syndrome (polyneuropathy, organomegaly, endocrinopathy, monoclonal protein, and skin changes), or amyloidosis **AND**
- P. Tecvayli will not be given concurrently with live vaccines **AND**
- Q. Member has not used a prior therapy that targets BCMA and/or is a CD3-redirecting therapy including Tecvayli **AND**
- R. Tecvayli will be prescribed through the consultation of a hematologist or oncologist **AND**
- S. Tecvayli will be given based on the FDA approved dosing (See Appendix A and B for dosing guidance) **AND**
- T. Member will receive Tecvayli at a healthcare facility enrolled in the Tecvayli REMS and are aware of how to manage relevant toxicities of Tecvayli (See Appendices C through E)

Initial Duration of Approval: 12 months

RENEWAL REQUEST:

Multiple myeloma (MM) that is refractory or in relapse.

- A. Initial conditions of coverage have been met **AND**
- B. Member has experienced a positive clinical response to Tecvayli and continuation of therapy is deemed clinically appropriate by the prescriber **AND**

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- C. Member has not experienced **ANY** of the following adverse reactions:
- Recurrent grade 3 CRS or grade 3 CRS with duration \geq 48 hours
 - Grade 4 CRS
 - Recurrent grade 3 or grade 4 neurological toxicity
 - Grade 4 infection
 - Grade 4 non-hematological adverse reaction
 - Serious hypersensitivity reaction

Renewal Duration of Approval: 12 months

5. LIMITATIONS/ EXCLUSIONS:

- A. Tecvayli is considered to be experimental and investigational if prescribed for indications other than for the treatment of multiple myeloma that is refractory or in relapse.

6. APPLICABLE PROCEDURE CODES:

CPT	Description
C9399	Unclassified drugs or biologicals
J9999	Not otherwise classified, antineoplastic drugs

7. APPLICABLE DIAGNOSIS CODES:

CODE	Description
C90.00	Multiple myeloma not having achieved remission
C90.02	Multiple myeloma in relapse

8. REFERENCES:

- Tecvayli (teclistamab-cqyv) [prescribing information]. Horsham, PA: Janssen Biotech, Inc; October 2022.
- Moreau P, Garfall AL, van de Donk NWCJ, et al. Teclistamab in Relapsed or Refractory Multiple Myeloma. N Engl J Med. 2022;387(6):495-505. doi:10.1056/NEJMoa2203478

9. Appendix A: Tecvayli Recommended Dosing Schedule

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Dosing Schedule	Day	Dose ^c	
Step-up Dosing Schedule	Day 1	Step-up dose 1	0.06 mg/kg SC
	Day 4 ^a	Step-up dose 2	0.3 mg/kg SC
	Day 7 ^b	First treatment dose	1.5 mg/kg SC
Weekly Dosing Schedule	One week after first treatment dose and weekly thereafter	Subsequent treatment doses	1.5 mg/kg SC once weekly until disease progression or unacceptable toxicity

^aStep-up dose 2 may be administered 2 to 4 days after step-up dose 1 and, if necessary, up to 7 days after step-up dose 1 to allow for resolution of adverse reactions.

^bThe first treatment dose may be administered 2 to 4 days after step-up dose 2 and, if necessary, up to 7 days after step-up dose 2 to allow for resolution of adverse reactions.

^cDose is based on actual body weight

10. Appendix B: Recommendations for Restarting Tecvayli^a After Dose Delay

Last Tecvayli dose administered	Duration of delay from the last Tecvayli dose administered	Action
Step-up dose 1	>7 days	Restart Tecvayli ^a step-up dosing schedule at 0.06 mg/kg (step-up dose 1).
Step-up dose 2	8 to 28 days	Repeat Tecvayli ^b step-up dose 2 (0.3mg/kg) and resume the step-up dosing schedule

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	>28 days ^c	Restart Tecvayli ^b step-up dosing schedule at 0.06 mg/kg (step-up dose 1).
Any treatment dose	8 to 28 days	Continue Tecvayli weekly dosing schedule at 1.5 mg/kg once weekly
	>28 days ^c	Restart Tecvayli ^b step-up dosing schedule at 0.06 mg/kg (step-up dose 1)

^aPatients should be hospitalized for 48 hours after all doses within the teclistamab step-up dosing schedule.

^bAdminister premedication prior to teclistamab administration and monitor accordingly.

^cConsider risk/benefit of restarting teclistamab if a dose delay of >28 days occurs due to an adverse reaction.

11. Appendix C: CRS Grading and Management Guidance

CRS Grade & Symptoms	Actions
Grade 1 Temperature $\geq 38^{\circ}\text{C}$ (100.4°F)* attributed to CRS.	Withhold teclistamab until CRS resolves. Administer premedication prior to the next teclistamab dose.
Grade 2 Temperature $\geq 38^{\circ}\text{C}$ (100.4°F)* attributed to CRS, with hypotension responsive to fluids and not requiring vasopressors and/or oxygen requirement of low-flow nasal cannula (≤ 6 L/minute) or blow-by.	Withhold teclistamab until CRS resolves. Administer premedication prior to the next teclistamab dose. Patients should be hospitalized for 48 hours following the next teclistamab dose.
Grade 3 Temperature $\geq 38^{\circ}\text{C}$ (100.4°F)* attributed to CRS, with hypotension requiring one vasopressor with or without vasopressin and/or oxygen requirement of high-flow nasal cannula (>6 L/minute), face mask, nonrebreather mask, or Venturi mask.	First occurrence of grade 3 CRS with duration <48 hours: Withhold teclistamab until CRS resolves. Provide supportive therapy as clinically necessary (may include intensive care). Administer premedication prior to the next teclistamab dose. Patients should be hospitalized for 48 hours following the next teclistamab dose.

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	<p>Recurrent grade 3 CRS or grade 3 CRS with duration ≥ 48 hours:</p> <p>Permanently discontinue teclistamab and provide supportive care as clinically necessary (may include intensive care).</p>
<p>Grade 4 Temperature $\geq 38^{\circ}\text{C}$ (100.4°F)* attributed to CRS, with hypotension requiring multiple vasopressors (excluding vasopressin) and/or oxygen requirement of positive pressure (eg, CPAP, BiPAP, intubation, and mechanical ventilation)**.</p>	<p>Permanently discontinue teclistamab and provide supportive care as clinically necessary (may include intensive care).</p>
<p>*Fever may be masked by antipyretics or anticytokine therapy. **CPAP = continuous positive airway pressure; BiPAP = bilevel positive airway pressure.</p>	

12. Appendix D: Tecvayli-Related Neurologic Toxicity Management

Severity Grade (Excluding ICANS)	Actions
Grade 1	Withhold teclistamab until neurologic toxicities/symptoms resolve or stabilize.
Grade 2 or grade 3 (first occurrence)	Withhold teclistamab until neurologic toxicities/symptoms improve to \leq grade 1. Provide supportive therapy as clinically appropriate.
Recurrent grade 3 or grade 4	Permanently discontinue teclistamab. Provide supportive care as clinically appropriate (may include intensive care).
Recommendations for management of Tecvayli-related ICANS	
ICANS Grade ^a & Symptoms ^b	Actions
Grade 1	Withhold teclistamab until ICANS resolves.
ICE score 7 to 9 ^c , or depressed level of consciousness ^d (awakens spontaneously)	Monitor neurologic symptoms and consider consultation with neurologist/other specialists for further evaluation and management (eg, consideration for initiating seizure prophylaxis with non-sedating, antiseizure medication).

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<p>Grade 2 ICE score 3 to 6^c, or depressed level of consciousness^d (awakens to voice)</p>	<p>Withhold teclistamab until ICANS resolves.</p> <p>Administer dexamethasone 10 mg IV every 6 hours (or equivalent); continue dexamethasone until resolution to ≤ grade 1, then taper.</p> <p>Monitor neurologic symptoms and consider consultation with neurologist/other specialists for further evaluation and management (eg, consideration for initiating seizure prophylaxis with nonsedating, antiseizure medication).</p> <p>Patients should be hospitalized for 48 hours following the next teclistamab dose.</p>
<p>Grade 3 ICE score 0 to 2^c, or depressed level of consciousness^d (awakens only to tactile stimulus), or seizures^d (either any clinical seizure, focal or generalized, that resolves rapidly, or nonconvulsive seizures on EEG that resolve with intervention), or</p> <p>Raised intracranial pressure (focal/local edema on neuroimaging^d)</p>	<p>First occurrence of grade 3 ICANS: Manage as per grade 2 ICANS. Provide supportive therapy as clinically appropriate (may include intensive care).</p> <p>Recurrent grade 3 ICANS: Permanently discontinue teclistamab. Manage as per grade 2 ICANS. Provide supportive therapy as clinically appropriate.</p>
<p>Grade 4 ICE score 0^c, or</p> <p>Depressed level of consciousness^d (either unarousable or requires vigorous/repetitive tactile stimuli to arouse, or stupor or coma), or</p>	<p>Permanently discontinue teclistamab. Manage with dexamethasone as per grade 2 ICANS. Alternatively, consider methylprednisolone 1,000 mg IV daily for ≥2 days.</p> <p>Monitor neurologic symptoms and consider consultation with neurologist/other specialists for further evaluation and management (eg, consideration for initiating seizure prophylaxis with nonsedating, antiseizure medication).</p>

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<p>Seizures^d (either life-threatening prolonged seizure >5 minutes, or repetitive clinical or electrical seizures without return to baseline in between), or</p> <p>Motor findings^d (deep focal motor weakness such as hemiparesis or paraparesis), or</p> <p>Raised intracranial pressure/cerebral edema^d, with signs/symptoms including diffuse cerebral edema on neuroimaging, or decerebrate or decorticate posturing, or cranial nerve VI palsy, or papilledema, or Cushing triad</p>	<p>Provide supportive therapy as clinically appropriate (may include intensive care).</p>
<p>^aBased on American Society for Transplantation and Cellular Therapy (ASTCT) 2019 grading for ICANS.</p> <p>^bManagement is determined by the most severe event (not attributable to any other cause).</p> <p>^cIf patient is arousable and able to perform immune effector cell-associated encephalopathy (ICE) assessment: Orientation (oriented to year, month, city, hospital = 4 points), naming (name 3 objects, eg, point to clock, pen, button = 3 points), following commands (eg, “show me 2 fingers” or “close your eyes and stick out your tongue” = 1 point), writing (ability to write a standard sentence = 1 point), attention (count backwards from 100 by 10 = 1 point). If unarousable and unable to perform ICE assessment (grade 4 ICANS = 0 points).</p> <p>^dNot attributable to any other cause.</p>	

13. Appendix E: Tecvayli Dosage Guidance for Other Adverse Reactions

Adverse Reaction	Severity	Actions
	ANC <500/mm3	Withhold teclistamab until ANC is ≥500/mm3.

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Hematologic toxicity	Febrile neutropenia	Withhold teclistamab until ANC is $\geq 1,000/\text{mm}^3$ and fever resolves.
	Hemoglobin < 8 g/dL	Withhold teclistamab until hemoglobin is ≥ 8 g/dL.
	Platelets $< 25,000/\text{mm}^3$ or platelets 25,000 to 50,000/ mm^3 with bleeding	Withhold teclistamab until platelets are $\geq 25,000/\text{mm}^3$ and no evidence of bleeding.
Hypersensitivity reactions (systemic or local)	Withhold or consider permanently discontinuing teclistamab based on reaction severity.	
Infections	Monitor immunoglobulin levels during treatment; manage according to guidelines, including infection precautions and antibiotic/antiviral prophylaxis.	
	All grades	Withhold teclistamab for active infection during the step-up dosing schedule.
	Grade 3	Withhold subsequent teclistamab treatment doses until infection improves to \leq grade 1.
	Grade 4	Consider permanent discontinuation of teclistamab. If not permanently discontinued, withhold subsequent treatment doses until infection improves to \leq grade 1.
Other nonhematologic adverse reactions	Grade 3	Withhold teclistamab until adverse reaction improves to \leq grade 1.
	Grade 4	Consider permanent discontinuation of teclistamab. If not permanently discontinued, withhold subsequent treatment doses until adverse reaction improves to \leq grade 1.

REVISION LOG:

REVISIONS	DATE
Creation date	1/2023
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