

DRUG NAME: Teclistamab

SYNONYM(S): teclistamab-cqyv1

COMMON TRADE NAME(S): TECVAYLI®

CLASSIFICATION: immunotherapy

Special pediatric considerations are noted when applicable, otherwise adult provisions apply.

MECHANISM OF ACTION:

Teclistamab is a bispecific T-cell engager that targets both B cell maturation antigen (BCMA) and CD3 receptors. It is a humanized immunoglobulin G4-proline, alanine, alanine (IgG4-PAA) antibody. Teclistamab binds to CD3 receptors expressed on T-cells and BCMA expressed on the surface of multiple myeloma cells. With its dual binding sites, teclistamab is able to redirect and draw T-cells in close proximity to BCMA-expressing tumour cells leading to T-cell activation, release of cytokines, and subsequent cell lysis.¹⁻³

USES:

Primary uses:

Other uses:

*Multiple myeloma

SPECIAL PRECAUTIONS:

Caution:

- severe *cytokine release syndrome (CRS)* can occur with teclistamab; recommended dosing regimen uses a step-up dosing schedule for initiation of treatment²
- *premedication* with corticosteroid, antihistamine, and antipyretic is recommended prior to all step-up doses and first full treatment dose to reduce the risk of CRS²
- immune effector cell-associated neurotoxicity syndrome (ICANS) has been reported; caution in patients with history of stroke or seizure, or pre-existing neurological problems²
- patients receiving teclistamab are at risk of reduced consciousness due to ICANS; patients should avoid driving or operating heavy machinery during the step-up schedule and for 48 hours after its completion or if experiencing neurologic symptoms²
- teclistamab step-up schedule should not be administered to patients with active infection²
- antimicrobial/antiviral prophylaxis may be required to prevent reactivation of infections such as hepatitis B and herpes zoster in high risk patients²
- vaccine response may be diminished during treatment with teclistamab²
- *immunization with live virus vaccines* is not recommended for at least 4 weeks prior to treatment, during treatment, and for at least 4 weeks after the last dose of teclistamab²

SIDE EFFECTS:

The table includes adverse events that presented during drug treatment but may not necessarily have a causal relationship with the drug. Because clinical trials are conducted under very specific conditions, the adverse event rates observed may not reflect the rates observed in clinical practice. Adverse events are generally included if they were reported in more than 1% of patients in the product monograph or pivotal trials, and/or determined to be clinically important.

BC Cancer Drug Manual[©]. All rights reserved. Page 1 of 6

Teclistamab (interim monograph)

This document may not be reproduced in any form without the express written permission of BC Cancer Provincial Pharmacy.

^{*}Health Canada approved indication



ORGAN SITE	SIDE EFFECT			
Clinically important side effects are in bold, italics				
blood and lymphatic system/ febrile neutropenia	anemia (55%, severe 37%)			
	febrile neutropenia (4%, severe 3%) ⁴			
	leukopenia (18%, severe 7%)			
	lymphopenia (35%, severe 33%)			
	neutropenia (71%, severe 64%)			
	thrombocytopenia (40%, severe 21 %)			
cardiac	arrhythmia (18%, severe 2%)			
gastrointestinal	emetogenic potential: low ^{5,6}			
	constipation (21%)			
	diarrhea (28%, severe 4%)			
	nausea (28%, severe 1%)			
	vomiting (13%, severe 1%)			
general disorders and	extravasation hazard: none ⁷			
administration site conditions	chills (18%)			
Corrainorio	edema, peripheral and facial (14%)			
	fatigue/asthenia (41%, severe 3%)			
	hypersensitivity (1%)			
	injection site reaction (35-38%, severe 1%); see paragraph following Side Effects table			
	pain (21%, severe 2%)			
	<i>pyrexia</i> (79%, severe 3%)			
hepatobiliary	hepatic failure (<1%); fatalities reported			
immune system	cytokine release syndrome (72%, severe 1%); see paragraph following Side Effects table			
	hypogammaglobulinemia (75%, severe 2%)			
infections and	cellulitis (4%)			
infestations	COVID-19 (18%, severe 12%); fatalities reported			
	herpes simplex (2%), herpes zoster (1%)			
	reactivation of viral infection (1%) (e.g., adenovirus, hepatitis B virus, cytomegalovirus, varicella zoster virus)			
	pneumonia (28%, severe 19%)			
	sepsis (8%, severe 7%)			
	upper respiratory tract infection (37%, severe 2%)			
investigations	activated partial thromboplastin time prolonged (8%, severe 1%) ⁴			
	albumin decrease (72%, severe 6%)			
	alkaline phosphatase increase (43%, severe 3%)			



ORGAN SITE	SIDE EFFECT			
Clinically important side effects are in bold, italics				
	ALT increase (35%, severe 4%); can occur with or without concurrent CRS			
	amylase increase (24%, severe 4%)			
	AST increase (41%, severe 3%); can occur with or without concurrent CRS			
	creatinine increase (34%, severe 3%)			
	calcium decrease (35%, severe 2%)			
	calcium increase (28%, severe 4%)			
	gamma-glutamyltransferase increase (38%, severe 9%)			
	INR increase (6%, severe 1%) ⁴			
	lipase increase (25%, severe 5%)			
	potassium decrease (20%, severe 2%)			
	potassium increase (31%, severe 5%)			
	magnesium decrease (28%)			
	phosphate decrease (43%, severe 15%)			
	sodium decrease (36%, severe 12%)			
metabolism and nutrition	appetite decrease (12%, severe 1%)			
musculoskeletal and	bone pain (16%, severe 3%)			
connective tissue	musculoskeletal pain (52%, severe 8%)			
nervous system	encephalopathy (10%)			
(see paragraph following Side Effects table)	Guillain-Barre syndrome; fatalities reported			
	headache (27%, severe <1%)			
	immune effector cell-associated neurotoxicity syndrome (3%)			
	motor dysfunction (19%)			
	peripheral neuropathy (16%, severe <1%)			
renal and urinary	acute kidney injury (11%, severe 4%)			
respiratory, thoracic and mediastinal	cough (24%)			
	<i>dyspnea</i> (13%, severe 2%)			
	hypoxia (20%, severe 4%)			
vascular	hemorrhage (12%, severe 3%); fatalities reported			
	hypertension (13%, severe 5%)			
	hypotension (21%, severe 2%)			

Adapted from standard reference^{1,2,8} unless specified otherwise.

Cytokine release syndrome (CRS) occurs in 72% of patients treated with teclistamab and recurs in more than onethird of patients. Signs and symptoms of CRS may include fever, chills, hypotension, tachycardia, hypoxia, headache, elevated liver enzymes, fatigue, nausea and vomiting. The majority of CRS events are grade 1 or 2. Although grade 3 events are uncommon (0.6%), potentially life-threatening complications of CRS have been

This document may not be reproduced in any form without the express written permission of BC Cancer Provincial

Pharmacy.



reported, including cardiac dysfunction, respiratory distress syndrome, neurologic toxicity, renal and/or hepatic failure, and disseminated intravascular coagulation. Median time to onset of CRS is 2 days after the most recent dose (range 1-6 days), and the median duration is 2 days (range 1-9 days). The step-up dosing schedule and premedication with corticosteroids, antihistamine, and antipyretics are used to reduce the risk of CRS as most patients experience CRS following the initial step-up doses or first treatment dose. Failure to follow the recommended dosing schedule for initiation of therapy and re-initiation of therapy after dose delays may result in increased frequency and severity of CRS. Hold teclistamab until CRS resolves and provide supportive care as needed. Avoid using myeloid growth factors, such as granulocyte macrophage-colony stimulating factor (GM-CSF) during CRS. Permanently discontinue teclistamab for recurrent grade 3 reactions, grade 3 reactions which last longer than 48 hours, and grade 4 reactions. Refer to protocol by which patient is being treated. For further information about the management of CRS, see BC Cancer Protocol SCCRS Cytokine Release Syndrome Management.

Local injection site reactions are reported in approximately one-third of patients. The majority of the reactions are grade 1 in severity. Reported reactions include injection site erythema, bruising, cellulitis, hematoma, induration, inflammation/swelling, discomfort, pruritus, and rash. The volume administered for each subcutaneous injection should not exceed 2 mL and this may require dividing the dosing volume into multiple syringes for administration.²

Neurologic toxicity has been reported in 57% of patients, including headache, motor dysfunction (e.g., dysgraphia, dysphonia, tremour, hypokinesia and gait disturbance), peripheral neuropathy, and encephalopathy. The most frequently reported neurologic toxicity has been headache. Neurologic toxicity can occur days or weeks after the teclistamab injection and initial symptoms may be subtle. The majority of neurologic events are grade 1 or 2; however, serious or life-threatening toxicity such as **Guillain-Barré** and **immune effector cell-associated neurotoxicity syndrome (ICANS)** can also occur. The most frequent clinical manifestations of ICANS are confusional state and dysgraphia. Median time to onset of ICANS is 4 days after the most recent dose of teclistamab (range 2 to 8 days) and the median duration is 3 days (range 1 to 20 days). The onset of ICANS may be concurrent with CRS, follow the resolution of CRS, or it can occur in the absence of CRS. Neurology consult may be required. Hold teclistamab until neurologic toxicity resolves. Symptoms are managed depending on their severity and whether they occur concurrently with CRS. Permanently discontinue teclistamab for recurrent grade 3 and grade 4 events. Due to the potential for ICANS and the risk of reduced consciousness, patients receiving teclistamab should avoid driving or operating heavy machinery during the step-up schedule and for 48 hours after its completion or if experiencing neurologic symptoms.^{1,2}

INTERACTIONS:

The initial release of cytokines associated with teclistamab treatment may suppress CYP450 enzymes. Substrates of CYP450 enzymes with a narrow therapeutic index may require dose adjustment and monitoring for toxicity if given concurrently with teclistamab. The highest risk of interaction is predicted to occur during the teclistamab step-up regimen and up to 7 days after the first treatment dose, or during a CRS event.²

SUPPLY AND STORAGE:

Injection: Janssen Inc. supplies teclistamab as single-use (preservative free) ready-to-use vials in two vial sizes: 30 mg vials in a concentration of 10 mg/mL and 153 mg vials in a concentration of 90 mg/mL. Refrigerate. Store in original carton to protect from light. Do not shake.²

Additional information:

• *Caution:* teclistamab vials are supplied as two different concentrations (10 mg/mL and 90 mg/mL); ensure selection of appropriate vial size for dose preparation

For basic information on the current brand used at BC Cancer, see <u>Chemotherapy Preparation and Stability</u> <u>Chart</u> in Appendix.

BC Cancer Drug Manual[©]. All rights reserved. Page 4 of 6

Teclistamab (interim monograph)

This document may not be reproduced in any form without the express written permission of BC Cancer Provincial



SOLUTION PREPARATION AND COMPATIBILITY:

For basic information on the current brand used at BC Cancer, see <u>Chemotherapy Preparation and Stability</u> <u>Chart</u> in Appendix.

Additional information:

- Caution: teclistamab vials are supplied as two different concentrations (10 mg/mL and 90 mg/mL); ensure selection of appropriate vial size for dose preparation
- injection volumes greater than 2.0 mL should be divided into separate syringes for administration²
- do not use closed system transfer devices (CSTD) for preparation or administration of syringe volumes less than 1 mL¹⁰; filtered venting needles (e.g., Chemo-Vent®) will be used in place of a CSTD for volumes less than 1 mL¹¹

Compatibility: consult detailed reference

PARENTERAL ADMINISTRATION:

BC Cancer administration guideline noted in **bold**, **italics**

	be carried administration gardenie neted in bota, raine	
Subcutaneous ²	injection into abdomen is preferred; may be administered	
	into the thigh ²	
	• up to 2 mL volume can be injected in a single site ²	
	if multiple injections are required, injections should be	
	spaced at least 2 cm apart. ²	
	<u> </u>	
Intramuscular	no information found	
Direct intravenous	do NOT use ²	
Intermittent infusion	do NOT use ²	
Continuous infusion	do NOT use ²	
Intraperitoneal	no information found	
Intrapleural	no information found	
Intrathecal	no information found	
Intra-arterial	no information found	
Intravesical	no information found	

DOSAGE GUIDELINES:

Refer to protocol by which patient is being treated. Numerous dosing schedules exist and depend on disease, response and concomitant therapy. Guidelines for dosing also include consideration of absolute neutrophil count (ANC). Dosage may be reduced, delayed or discontinued in patients with bone marrow depression due to cytotoxic/radiation therapy or with other toxicities.



Adults:

BC Cancer usual dose noted in bold, italics

Cycle Length:

4 weeks^{2,12,13}: Subcutaneous:

Cvcle 1:

Dosing Schedule		Day of treatment	Dose (SC)
Step-up dosing schedule	Step-up dose 1	day 1	0.06 mg/kg single dose
	Step-up dose 2	day 3*	0.3 mg/kg single dose
	First treatment dose	day 5**	1.5 mg/kg single dose

^{*} may be given between 2 to 7 days after step-up dose 1

Cycle 2 starting 7 days after the last dose of cycle 1 and onwards: 1.5 mg/kg SC given once weekly on days 1, 8, 15, and 22 (total dose per cycle 6 mg/kg)

minimum of 5 days should be maintained between weekly doses² no dose reductions are recommended2

Following dose delays: refer to protocol by which patient is being treated for instruction as the step-up dose schedule is repeated when restarting teclistamab after a dose delav²

REFERENCES:

- 1. Janssen Biotech Inc. TECVAYLI® full prescribing information. Horsham, PA, USA; Oct 25, 2022
- 2. Janssen Inc. TECVAYLI® product monograph. Toronto, Ontario; July 26, 2023
- 3. Lexi-Drugs® Lexicomp Online (database on the Internet). Teclistamab. Lexi-Comp Inc., Available at: http://online.lexi.com. Accessed August 8, 2023
- 4. Janssen-Cilag. TECVAYLI® product information. Beerse, Antwerp, Belgium; July 21, 2023
- 5. BC Cancer Supportive Care Tumour Group. (SCNAUSEA) BC Cancer Guidelines for Prevention and Treatment of Chemotherapy-Induced Nausea and Vomiting in Adults. Vancouver, British Columbia: BC Cancer; September 1 2022
- 6. Ettinger DS. NCCN Practice Guidelines in Oncology Antiemesis v.2.2023. National Comprehensive Cancer Network, 2023.
- Available at: http://www.nccn.org. Accessed June 26, 2023

 7. BC Cancer Provincial Systemic Therapy Program. Provincial Systemic Therapy Program Policy III-20: Prevention and Management of Extravasation of Chemotherapy. Vancouver, British Columbia: BC Cancer; March 1 2021
- 8. Moreau P, Garfall AL, van de Donk, N. W. C., et al. Teclistamab in relapsed or refractory multiple myeloma. N Engl J Med 2022;387(6):495-505
- 9. Janssen Research & Development LLC. JNJ-64007957 (teclistamab) Investigator's Brochure Edition 7. Beerse, Belgium; May
- 10. Janssen Research & Development LLC. Investigational product preparation and Administration instructions for subcutaneous Administration of 10 mg/mL and 90 mg/mL Teclistamab (JNJ-64007957) For Weight-Based Dosing in mcg/kg for Managed Access Programs - Version 6.0 . Beerse, Belgium; June 9 2023
- 11. de Lemos, M. BC Cancer Provincial Pharmacy Professional Practice Leader and Drug Information Coordinator. SBAR: Teclistamab Subcutaneous Injection for Multiple Myeloma at BC Cancer. (draft). BC Cancer - Systemic Therapy Program 2023 12. Janssen Research & Development LLC. Pre-approval Access (PAA) Named Patient Program (NPP) Treatment Guidelines for Teclistamab (JNJ-64007957) for Treating Physician Use 64007957MMY4001. Version 4.0. Beerse, Belgium, May 12, 2023
- 13. Moreau P, Garfall AL, van de Donk, N. W. C., et al. Teclistamab in relapsed or refractory multiple myeloma (Clinical Protocol). N Engl J Med 2022;387(6):495-505.

^{**} may be given between 2 to 7 days after step-up dose 2 (total dose per cycle 1.86 mg/kg)