

## Regimen Reference Order

### CUTA – tebentafusp (Outpatient CYCLE 2 and Onwards)

ARIA: CUTA - [tebentafusp]

**Planned Course:** Once weekly until disease progression or unacceptable toxicity  
(1 cycle = 21 days)

**Indication for Use:** Uveal Melanoma

**CVAD:** At Provider’s Discretion

**Proceed with treatment if:**

**Day 1**

- *ANC equal to or greater than 1 x 10<sup>9</sup>/L AND Platelets equal to or greater than 50 x 10<sup>9</sup>/L*
- *AST/ALT equal to or less than 3 times upper limit of normal*
- *Total bilirubin equal to or less than 1.5 times the upper limit of normal*
- *Systolic blood pressure greater than 100 mmHg*

**Days 8 and 15**

- *AST/ALT equal to or less than 3 times upper limit of normal*
- *Total bilirubin equal to or less than 1.5 times the upper limit of normal*
- *Systolic blood pressure greater than 100 mmHg*
- ❖ **Contact Physician if parameters not met**

#### SEQUENCE OF MEDICATION ADMINISTRATION

##### Pre-treatment Requirements

Drug	Dose	CCMB Administration Guideline
Not Applicable		

#### Treatment Regimen – CUTA – tebentafusp (Outpatient CYCLE 2 and Onwards)

Establish primary solution 500 mL of: normal saline

Drug	Dose	CCMB Administration Guideline
<b>Cycle 1</b>		
Patients will be admitted to hospital for tebentafusp (Days 1, 8 and 15). Follow inpatient orders		
<b>Cycle 2</b>		
<b>Days 1, 8 and 15</b>		
tebentafusp	68 mcg	IV in normal saline 100 mL over 15 minutes <i>Use 0.2 to 0.22 micron filter</i>

**In the event of an infusion-related hypersensitivity reaction, refer to the ‘Hypersensitivity Reaction Standing Order**

## REQUIRED MONITORING (Outpatient)

### Outpatient Cycle 2 and Onwards

#### All Doses

- Full vital signs (temperature, heart rate, respiratory rate, blood pressure and O<sub>2</sub> saturation) prior to tebentafusp administration and as clinically indicated
- Observe patient for 30 minutes after every tebentafusp infusion. Full vital signs prior to discharge
- Monitor for signs and symptoms of cytokine release syndrome (CRS). Serious adverse events that may be associated with CRS include: pyrexia, headache, nausea, asthenia, hypotension, and elevations in serum aminotransferases and bilirubin
- Skin assessment for rash

#### Day 1

- CBC, serum creatinine, urea, electrolytes, liver enzymes and total bilirubin as per Physician Orders
- Lipase as per Physician Orders

#### Days 8 and 15

- Liver enzymes as per Physician Orders

### Recommended Support Medications

Drug	Dose	CCMB Administration Guideline
None required		

## DISCHARGE INSTRUCTIONS

- Patient to notify clinic if they develop fever, chills, nausea, vomiting, rash and/or headache

## ADDITIONAL INFORMATION

- This Regimen Reference Order (RRO) is intended for Cycle 2 and onwards of tebentafusp. Patients MUST receive inpatient administration of Cycle 1 of tebentafusp prior to outpatient administration
- Cycle 1 of tebentafusp is restricted to inpatient hospital administration (ramp-up dosing) due to the high risk of cytokine release syndrome (CRS)
- Patients should be monitored for CRS throughout therapy. Patients are at highest risk of CRS during the first cycle of tebentafusp
- tebentafusp can cause elevations in lipase and liver enzymes
- tebentafusp can cause skin rash
- tebentafusp preparations contain human albumin (blood product)
- Site restrictions are in place for tebentafusp. tebentafusp must be administered at CCMB MacCharles in Winnipeg