

Regimen Reference Order – HEME – luspatercept (MDS)

ARIA: HEME - [luspatercept (MDS)]

Planned Course: Every 21 days

Indication for Use: Transfusion-dependent anemia due to Myelodysplastic Syndrome (MDS)

CVAD: Not Required

Proceed with treatment if:

Hemoglobin less than 115 g/L

❖ Contact Hematologist if parameters not met

SEQUENCE OF MEDICATION ADMINISTRATION

Pre-treatment Requirements

Drug	Dose	CCMB Administration Guideline
Not Applicable		

Treatment Regimen – HEME – luspatercept (MDS)

Drug	Dose	CCMB Administration Guideline
luspatercept	1 mg/kg*; maximum dose 168 mg	Subcutaneous once every 3 weeks into the upper arm, thigh, and/or abdomen <i>*Nursing Alert: Maximum volume at each injection site is 1.2 mL. If more than 1 syringe is required, injections should be given using the same anatomical location but on opposite sides of the body (e.g. left thigh and right thigh)</i>
<p><i>*Dose may be increased to 1.33 mg/kg at the hematologist's discretion if patient is not RBC transfusion-free after at least two consecutive doses (6 weeks) at the 1 mg/kg starting dose. Dose may be further increased to 1.75 mg/kg if patient is not RBC transfusion-free after at least two consecutive doses (6 weeks) at the 1.33 mg/kg dose</i></p> <p><i>*If hemoglobin increases by greater than 20 g/L within 3 weeks, in the absence of a transfusion, dose reduction may be required at the hematologist's discretion</i></p> <p>All doses will be automatically rounded that fall within CCMB Approved Dose Bands. See Dose Banding document for more information</p>		

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

REQUIRED MONITORING

All Cycles

- CBC, serum creatinine, urea, liver enzymes and total bilirubin as per Physician Orders
- Blood pressure at each clinic visit
- Full vital signs (temperature, heart rate, respiratory rate, blood pressure and O₂ saturation) prior to each dose and as clinically indicated
- No observation period required. Patient can be discharged from treatment room if stable whether they had a reaction or not

Recommended Support Medications

Drug	Dose	CCMB Administration Guideline
None required		

DISCHARGE INSTRUCTIONS

- Patients should be instructed to contact their cancer team immediately if symptoms of hypersensitivity reactions occur after discharge
- Instruct patient to report to clinic if they are experiencing pain or injection site reactions that are bothersome

ADDITIONAL INFORMATION

- luspatercept may increase blood pressure
- luspatercept is teratogenic