

PROTOCOL CODE: MYDARLDF (IV Cycle 1)

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<b>Patient</b>	RevAid	#

DOCTOR'S ORDERS Htcm Wt	kg BSAm²			
REMINDER: Please ensure drug allergies and previous bleomycin are documented on the Allergy & Alert Form				
DATE: To be given: Cycle	pe given: Cycle #: 1			
Date of Previous Cycle:  Risk Category: Female of Childbearing Potential (FCBP) Rx valid for 7 days  Risk Category: Male or Female of non-Childbearing Potential (NCBP)  ****Ensure Red Blood Cell Phenotype and Group and Screen for all patients prior to Cycle 1****				
Delay treatment week(s)  CBC & Diff day of treatment  Proceed with all medications as written, if within 96 hours of Day 1: ANC greater than or equal to 1 x 109/L, platelets greater than or equal to 50 x 109/L, and eGFR or creatinine clearance as per protocol  Dose modification for: Hematology: Other Toxicity:  Proceed with treatment based on blood work from				
LENALIDOMIDE One cycle = 28 days  Per physician's clinical judgement, physician to ensure prophylaxis with valACYclovir 500 mg PO daily lenalidomide* mg PO daily, in the evening, on Days 1 to 21 and off for 7 days lenalidomide* mg PO  (*available as 25 mg, 20 mg, 15 mg, 10 mg, 5 mg and 2.5 mg capsules) *Note: Use one capsule strength for the total dose; there are cost implications as costing is per capsule and not weight based  Mitte: FCBP dispense 21 capsules (1 cycle) For Male and Female NCBP: Mitte: 21 capsules (1 cycle).  Physician to ensure DVT prophylaxis in place: ASA, Warfarin, low molecular weight heparin, direct oral anticoagulant or none (select one)	Pharmacy Use for Lenalidomide dispensing:  RevAid confirmation number:  Lenalidomide lot number:  Pharmacist counsel (initial):			
DOCTOR'S SIGNATURE:  Physician Revaid ID:	SIGNATURE: UC:			



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DATE:				
STEROID: RN to use patient's therapeutic steroid as pre-med for daratur	numab - refer to protocol.			
Standard Regimen: daratumumab full dose administered on Cycle 1 Day 1				
☐ dexamethasone ☐ 40 mg or ☐ 20 mg PO before daratumumab on Days 1, 8, 15 and 22 OR				
predniSONE 100 mg PO before daratumumab on Days 1, 8, 15 and 22				
OR				
Alternative Regimen: daratumumab split dose administered on Cycle 1 Day 1	and Day 2			
☐ <b>dexamethasone 20 mg</b> PO before daratumumab on Days 1 and 2, and <b>40 mg</b> before daratumumb on Days 8, 15, 22 <i>OR</i>				
dexamethasone 20 mg PO before daratumumab on Days 1 and 2 and 20 mg before daratumumb on Days 8, 15, 22				
OR ☐ predniSONE 50 mg PO before daratumumab on Days 1 and 2, and predniSONE 100 mg before daratumumb on Days 8, 15, 22				
**Have Hypersensitivity Reaction Tray and Protocol Available**				
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**Have Hypersensitivity Reaction Tray and Protocol  DARATUMUMAB  • Per physician's clinical judgement, physician to ensure prophylaxis with valACYo				
<ul> <li>DARATUMUMAB</li> <li>Per physician's clinical judgement, physician to ensure prophylaxis with valACY</li> <li>DARATUMUMAB PREMEDICATIONS: Patient to take own supply. RN/Pharmacian</li> </ul>	clovir 500 mg PO daily			
Per physician's clinical judgement, physician to ensure prophylaxis with valACY      DARATUMUMAB PREMEDICATIONS: Patient to take own supply. RN/Pharmacidexamethasone as ordered in steroid section.	clovir 500 mg PO daily st to confirm.			
<ul> <li>DARATUMUMAB</li> <li>Per physician's clinical judgement, physician to ensure prophylaxis with valACY</li> <li>DARATUMUMAB PREMEDICATIONS: Patient to take own supply. RN/Pharmacian</li> </ul>	clovir 500 mg PO daily st to confirm.			
Per physician's clinical judgement, physician to ensure prophylaxis with valACY      DARATUMUMAB PREMEDICATIONS: Patient to take own supply. RN/Pharmaci dexamethasone as ordered in steroid section montelukast 10 mg PO prior to daratumumab on Day 1 (and Day 2 if on alternation).	clovir 500 mg PO daily st to confirm. ative regimen)			
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<ul> <li>Per physician's clinical judgement, physician to ensure prophylaxis with valACYo</li> <li>DARATUMUMAB PREMEDICATIONS: Patient to take own supply. RN/Pharmaci dexamethasone as ordered in steroid section montelukast 10 mg PO prior to daratumumab on Day 1 (and Day 2 if on altern montelukast 10 mg PO prior to daratumumab on Days 8, 15 and 22 acetaminophen 650 mg PO prior to each daratumumab. Repeat acetaminophen edded if IV infusion exceeds 4 hours</li> <li>Select one of the following:         <ul> <li>□ loratadine 10 mg PO prior to each daratumumab, then diphenhydrAMINE</li> </ul> </li> <li>OR</li> <li>□ diphenhydrAMINE 50 mg □ PO or □ IV prior to each daratumumab. Repeat</li> </ul>	clovir 500 mg PO daily st to confirm. ative regimen) nen 650 mg PO every 4 hours when 50 mg IV every 4 hours when needed			



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DATE:					
**Have Hypersensitivity Reaction Tray and Protocol Ava	ailable**				
<b>Standard regimen</b> : daratumumab full dose administered on Cycle 1 Day 1					
CYCLE 1, Day 1:					
daratumumab (First dose) 16 mg/kg x kg = mg IV in 1000 mL NS (use 0.2 micron in-line filter)					
OR					
Alternative regimen: daratumumab split dose administered on Cycle 1 Day 1 and Day 2					
CYCLE 1, Days 1 and 2					
daratumumab 8 mg/kg x kg = mg IV in 500 mL NS (use 0.	.2 micron in-line filter)				
Infusion rate for Day 1, (and Day 2, if Alternative regimen):					
Start at 50 mL/h. If no infusion-related reactions after 60 minutes, increase by 50 mL/h every 60 minutes to a maximum					
rate of 200 mL/h If BP falls to less than 80/50 mmHg or pulse increases to greater than 120 or if flushing	n dyspnea chills rash pruritus				
vomiting, chest pain, throat tightness, cough, wheezing, or any other new acute discon					
infusion and page physician.					
Vitals monitoring:					
Vital signs immediately before the start of infusion, then every 30 minutes x 4, then every					
infusion and at 30 minutes post infusion. Observe patient for 30 minutes after each da	ratumumab infusion				
☐ CYCLE 1, Day 8:					
daratumumab 16 mg/kg x kg = mg IV in 500 mL NS (use 0	0.2 micron in-line filter)				
Infusion rate: Physician to determine rate of infusion					
If no reaction in the previous infusion or reaction is Grade 2 or less:					
☐ Start at 200 mL/h. If no infusion-related reactions after 30 minutes, infuse the remai	inder at 450 mL/h  (Rapid infusion)				
OR					
If reaction in the previous infusion is Grade 3:					
Start at 50 mL/h. If no infusion-related reactions after 60 minutes, increase by 50 mL/h every 60 minutes to a maximum rate of 200 mL/h (Slow infusion)					
Vitals monitoring:					
Vital signs immediately before the start, at the end of the infusion and as needed. Observe patient for 30 minutes after					
infusion.					
DOCTOR'S SIGNATURE.	CICNATURE.				
DOCTOR'S SIGNATURE:	SIGNATURE:				
	UC:				



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DATE:				
**Have Hypersensitivity Reaction Tray and Protocol Available**				
CYCLE 1, Days 15 and 22:				
daratumumab 16 mg/kg x kg = mg IV in 500 mL NS (use 0.2 micron in-line filter)				
Infusion rate for Days 15 and 22: Physician to determine rate of infusion				
If no reaction in the previous infusion or reaction is Grade 2 or less:				
☐ Start at 200 mL/h. If no infusion-related reactions after 30 minutes, infuse the remainder at 450	mL/h (Rapid infusion)			
OR				
If reaction in the previous infusion is Grade 3:				
Start at 100 mL/h. If no infusion-related reactions after 60 minutes, increase by 50 mL/h every 60 minutes to a maximum rate of 200 mL/h. Refer to protocol for modified starting rate if previous infusion reactions were experienced during infusion rate of greater than or equal to 100 mL/h (Slow infusion)				
<b>Vitals monitoring:</b> Vital signs immediately before the start, at the end of the infusion and as needed. Observe patient for 30 minutes after infusion. (Vitals and observation post-infusion not required after 3 treatments with no reaction).				
RETURN APPOINTMENT ORDERS				
☐ STANDARD REGIMEN: For Cycle 1, book chemo on Days 1, 8, 15 and 22				
☐ ALTERNATIVE REGIMEN: For Cycle 1, book chemo on Days 1, 2, 8, 15 and 22				
For Cycle 2 book chemo on Days 1, 8, 15, 22				
Return in <b>four</b> weeks for Doctor and Cycle 2				
CBC & Diff, creatinine, urea, sodium, potassium, total bilirubin, ALT, alkaline phosphatase, calcium, albumin, LDH, random glucose, serum protein electrophoresis <u>and</u> serum free light chain levels every 4 weeks				
TSH every three months (i.e. prior to Cycles 4, 7, 10, 13, 16 etc)  Urine protein electrophoresis every 4 weeks  Immunoglobulin panel (IgA, IgG, IgM) every 4 weeks  Beta-2 microglobulin every 4 weeks  CBC & Diff, platelets Days 8, 15, 22  Creatinine, sodium, potassium Days 8, 15, 22  Total bilirubin, ALT, alkaline phosphatase Days 8, 15, 22  Random glucose Days 8, 15, 22  Calcium, albumin Days 8, 15, 22  Quantitative β-hCG blood test for FCBP 7-14 days and 24 h prior to cycle 1 and every week for 4 weeks during cycle 1  Quantitative β-hCG blood test for FCBP less than or equal to 7 days prior to cycle 2  HBV viral load prior to next cycle  Other tests:  Consults:  See general orders sheet for additional requests				
DOCTOR'S SIGNATURE:	SIGNATURE:			
	UC:			