Regimen Reference Order – LYMP – riTUXimab (Weekly X 4)

ARIA: LYMP – [riTUXimab (weekly)]

Planned Course:Weekly for 4 weeksIndication for Use:Non-Hodgkin Lymphoma ORPost-Transplant Lymphoproliferative Disorder OREpstein-Barr Viremia

CVAD: At Provider's Discretion

Proceed with treatment if:

Day 1

• ANC equal to or greater than 1×10^9 /L AND Platelets equal to or greater than 75 x 10^9 /L

Days 8, 15 and 22

• Blood work not required to proceed with treatment

Contact Hematologist if parameters not met

SEQUENCE OF MEDICATION ADMINISTRATION

Pre-treatment Requirements				
Drug	Dose	CCMB Administration Guideline		
allopurinol*	300 mg	Orally once daily for 10 days to begin 3 days prior to Day 1 and at provider's discretion for subsequent days		
		(Self-administered at home)		
		*Only patients at risk of tumor lysis syndrome will be prescribed allopurinol		

Treatment Regimen – LYMP – riTUXimab (Weekly X 4) Establish primary solution 500 mL of: normal saline

Establish primary solution 500 mL of: normal saline			
Drug	Dose	CCMB Administration Guideline	
Day 1			
cetirizine	10 mg	Orally 30 minutes prior to riTUXimab	
acetaminophen	650 mg	Orally 30 minutes prior to riTUXimab	
dexamethasone	20 mg	IV in normal saline 50 mL over 15 minutes	
Wait 30 minutes after	completion of IV pre-mo	edications before starting riTUXimab	



riTUXimab (IV brand name specific)	375 mg/m ²	Slow infusion(if greater than 6 months since last riTUXimab dose or no previous riTUXimab): IV made up to a final concentration of 1 mg/mL in normal saline. Start at 50 mg/hr for 60 minutes and escalate infusion rate in 50 mg/hr
		brackets on prescription label) matches prescribed order *Alert: Pharmacy to ensure final volume on label
Days 8, 15 and 22		Alert. Filamacy to ensure final volume on laber
cetirizine	10 mg	Orally 30 minutes prior to riTUXimab
acetaminophen	650 mg	Orally 30 minutes prior to riTUXimab
		cations before starting riTUXimab
riTUXimab	1400 mg	Subcutaneous: Administer over 5 minutes into abdomen
(Subcutaneous)	(1400 mg = 11.7 mL)	Syringe should be held in hand for 5 minutes to warm up and decrease viscosity
		Use 25G needle *Nursing Alert: Ensure subcutaneous riTUXimab formulation is
		used (riTUXimab-hyaluronidase, human)
	OR	
riTUXimab (IV brand name specific)	375 mg/m ²	Rapid infusion:IV in normal saline over 90 minutes:Infuse 50mL of a 250 mL bag (or 100 mL of a 500 mL bag) over 30minutes, then infuse the remaining 200 mL (or 400 mL of a 500mL bag) over 60 minutes*Alert:Ensure brand name on prescription label (indicated in brackets on prescription label) matches prescribed order*Alert:Pharmacy to ensure final volume on label
	OR	
		Slow infusion: IV made up to a final concentration of 1 mg/mL in normal saline. Start at 100 mg/hr for 30 minutes and escalate infusion rate in 100 mg/hr increments every 30 minutes to maximum of 400 mg/hr
		*Nursing Alert: IV tubing is primed with riTUXimab *Alert: Ensure brand name on prescription label (indicated in
		brackets on prescription label) matches prescribed order

*Alert: Pharmacy to ensure final volume on label

All doses will be automatically rounded that fall within the DSG Approved Dose Bands. See LYMP DSG – Dose Banding document for more information

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

REQUIRED MONITORING

Day 1

• CBC, serum creatinine, urea, electrolytes, liver enzymes, LDH, total bilirubin, uric acid and albumin as per Physician Orders

Days 8, 15 and 22

• No blood work required

INTRAVENOUS riTUXimab

- Full vital signs (temperature, heart rate, respiratory rate, blood pressure and O₂ saturation) at baseline and as clinically indicated
- No observation period is required after riTUXimab administration. Patient can be discharged from treatment room if stable whether they had a reaction or not

SUBCUTANEOUS riTUXimab

- Full vital signs (temperature, heart rate, respiratory rate, blood pressure and O₂ saturation) at baseline, <u>at discharge</u> and as clinically indicated
- 15 minute observation period required after each dose

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

ADDITIONAL INFORMATION

- Administering nurse must document any infusion-related reactions with any dose of riTUXimab
- Ensure there were **no Grade 3 or 4** infusion-related reaction with any previous dose prior to administering riTUXimab via subcutaneous injection or rapid infusion
- Intravenous riTUXimab formulation is available from more than one manufacturer and uses several different brand names. Brand name will be indicated in brackets after riTUXimab. Ensure prescription label matches the brand name on prescribed order for intravenous riTUXimab

