

Information on this form is a guide only. User will be solely responsible for verifying its currency and accuracy with the corresponding BC Cancer treatment protocols located at www.bccancer.bc.ca and according to acceptable standards of care

PROTOCOL CODE: LYIDELAR

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DOCTOR'S ORDERS Htcm Wtkg	BSA_	m²				
REMINDER: Please ensure drug allergies and previous bleomycin are documented on the Allergy & Alert Form						
DATE: To be given: Cy	/cle #:	of				
Date of Previous Cycle:						
□ Delay treatment week(s) □ CBC & Diff day of treatment May proceed with doses as written if within 96 hours ANC greater than or equal to 0.5 x 109/L, platelets greater than or equal to 25, ALT/AST less than or equal to 5 x ULN, bilirubin less than or equal to 3 x ULN						
Dose modification for:						
Have Hypersensitivity Reaction Tray and Protocol Availa	ble					
For intravenous riTUXimab infusion: diphenhydrAMINE 50 mg PO prior to riTUXimab IV and then q 4 h if IV infusion exceeds 4 h acetaminophen 650 mg to 975 mg PO prior to riTUXimab IV and then q 4 h if IV infusion exceeds 4 h For subcutaneous riTUXimab injection: diphenhydrAMINE 50 mg PO prior to riTUXimab subcutaneous acetaminophen 650 mg to 975 mg PO prior to riTUXimab subcutaneous						
TREATMENT: Physician to ensure antibiotic prophylaxis for PCP/PJP (e.g., cotrimoxazole 1 SS tab daily) is given throughout idelalisib treatment and for a period of 2 to 6 months after discontinuation. Counsel patient to obtain supply of loperamide and take 2 mg PO at first onset of diarrhea and q2h while awake and q4h during the night until diarrhea free x 12 hours						
For Cycle 1 ONLY idelalisib						
DOCTOR'S SIGNATURE:		SIGNATURE:				
		UC:				

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DOCTOR'S ORDERS				
DATE:				
** Have Hypersensitivity Reaction Tray and Protocol Available**				
TREATMENT con For Cycle 1 only (co				
	se) 375 mg/m² x BSA = mg NS on Day 1. Start at 50 mg/hour. After 1 hour, increasity occurs.	se rate by 50 mg/hour every 30	minutes until rate = 400	
Pharmacy to select ri	TUXimab IV brand as per Provincial Systemic Therapy	Policy III-190		
Drug	Brand (Pharmacist to complete. Please print.)	Pharmacist Initial and Dat	te	
riTUXimab				
Mitte: 30 days (dispe	g or ☐ 100 mg (select one) PO BID continuously nse in original container) e 2 and subsequent treatments: a full dose of IV riTUXimab (no severe reactions requiri	ng early termination) and can pr	roceed to subcutaneous	
riTUXimab subcut (RITUXAN SC) 1600 mg (fixed dose in 13.4 mL) subcutaneously into abdomen over 7 minutes. Observe for 15 minutes after administration.				
NB: During treatment possible.	t with subcutaneous riTUXimab, administer other subcu	ıtaneous drugs at alternative inj	jection sites whenever	
See page 3				
DOCTOR'S SIGNA	ATURE:		SIGNATURE:	
			UC:	

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DOCTOR'S ORDERS					
DATE:					
Have Hypersensitivity Reaction Tray and Protocol Available					
TREATM	ENT (con	ntinued):			
		erate a full dose of IV riTUXimab (experienced severe ntinue with IV riTUXimab for this cycle:	reactions requiring early termina	ation) in the previous	
		n² x BSA = mg			
) to 500 mL				
Pharmacy	to select ri	TUXimab IV brand as per Provincial Systemic Therapy	Policy III-190		
Drug		Brand (Pharmacist to complete. Please print.)	Pharmacist Initial and Dat	Pharmacist Initial and Date	
riTUXi	imab				
		mL of 500 mL bag) of the dose over 30 minutes, then usion time = 1 hour 30 min)	infuse the remaining 200 mL (or	r 400 mL of 500 mL bag)	
		rigors, rash, pruritus, vomiting, chest pain, any other no o infusion and page physician. Constant visual observa		ation of any existing	
Cycle 9 an	-	o iniusion and page physician. Constant visual observa	ation is not required.		
_	•	or ☐ 100 mg (<i>select one</i>) PO BID continuously			
		60 days or 90 days (dispense in original contained	er)		
		RETURN APPOINTME	INT ORDERS	ı	
☐ Cycle 1 to 8: Return in <u>four</u> weeks for Doctor and Cycle Book chemo on Day 1					
☐ Cycle 9 and beyond: Return in weeks (maximum 12 weeks) for Doctor					
☐ Last Cycle. Return in week(s).					
		######################################			
Laboratory: Cycles 1-3:					
		s, bilirubin, ALT, CMV-DNA by PCR every two weeks	5		
Cycles 4-6 CBC & Dif		s, bilirubin, ALT, CMV-DNA by PCR monthly			
Cycle 7 an	ıd subseq	uent cycles:			
CBC & Diff, Platelets, bilirubin, ALT, CMV-DNA by PCR monthly <u>OR</u> ☐ every 3 months, as clinically indicated					
☐ Consu		,			
Other t	tests:				
☐ See general orders sheet for additional requests. DOCTOR'S SIGNATURE:					
DOCTOR	'S SIGNA	ATUKE:		SIGNATURE:	
				UC:	

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