



Provincial Health Services Authority

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

DOCTOR'S ORDERS

DATE:

**** Have Hypersensitivity Reaction Tray and Protocol Available****

TREATMENT continued:

For Cycle 1 only (continued):

riTUXimab (first dose) 375 mg/m² x BSA = _____ mg

IV in 250 to 500 mL NS on Day 1. Start at 50 mg/hour. After 1 hour, increase rate by 50 mg/hour every 30 minutes until rate = 400 mg/hour unless toxicity occurs.

Pharmacy to select riTUXimab IV brand as per Provincial Systemic Therapy Policy III-190

Drug	Brand (Pharmacist to complete. Please print.)	Pharmacist Initial and Date
riTUXimab		

For the first dose, patients are to be under constant visual observation during all dose increases and for 30 minutes after infusion completed. If flushing, dyspnea, rigors, rash, pruritus, vomiting, chest pain, any other new acute discomfort or exacerbation of any existing symptoms occur, stop infusion and page physician. Vital signs are not required, unless symptomatic.

Cycle 2, 3, 4, 5, 6, 7 and 8

idelalisib 150 mg or 100 mg (select one) PO BID continuously

Mitte: 30 days (dispense in original container)

riTUXimab for Cycle 2 and subsequent treatments:

Patient tolerated a full dose of IV riTUXimab (no severe reactions requiring early termination) and can proceed to subcutaneous riTUXimab:

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 with subcutaneous riTUXimab, administer other subcutaneous drugs at alternative injection sites whenever possible.

See page 3

DOCTOR'S SIGNATURE:

SIGNATURE:

UC:

PROTOCOL CODE: LYIDELAR

DOCTOR'S ORDERS	
DATE:	
Have Hypersensitivity Reaction Tray and Protocol Available	
TREATMENT (continued):	
<input type="checkbox"/> Patient did not tolerate a full dose of IV riTUXimab (experienced severe reactions requiring early termination) in the previous treatment and will continue with IV riTUXimab for this cycle: riTUXimab 500 mg/m² x BSA = _____ mg IV in 250 to 500 mL NS	
Pharmacy to select riTUXimab IV brand as per Provincial Systemic Therapy Policy III-190	
Drug	Brand (Pharmacist to complete. Please print.)
Pharmacist Initial and Date	
riTUXimab	
Infuse 50 mL (or 100 mL of 500 mL bag) of the dose over 30 minutes, then infuse the remaining 200 mL (or 400 mL of 500 mL bag) over 1 hour. (total infusion time = 1 hour 30 min) If flushing, dyspnea, rigors, rash, pruritus, vomiting, chest pain, any other new acute discomfort or exacerbation of any existing symptoms occur, stop infusion and page physician. Constant visual observation is not required.	
Cycle 9 and beyond	
idelalisib <input type="checkbox"/> 150 mg or <input type="checkbox"/> 100 mg (select one) PO BID continuously Mitte: <input type="checkbox"/> 30 days or <input type="checkbox"/> 60 days or <input type="checkbox"/> 90 days (dispense in original container)	
RETURN APPOINTMENT ORDERS	
<input type="checkbox"/> Cycle 1 to 8: Return in four weeks for Doctor and Cycle _____. Book chemo on Day 1 <input type="checkbox"/> Cycle 9 and beyond: Return in _____ weeks (maximum 12 weeks) for Doctor <input type="checkbox"/> Last Cycle. Return in _____ week(s).	
Laboratory: Cycles 1-3: CBC & Diff, Platelets, bilirubin, ALT, CMV-DNA by PCR every two weeks Cycles 4-6: CBC & Diff, Platelets, bilirubin, ALT, CMV-DNA by PCR monthly Cycle 7 and subsequent cycles: CBC & Diff, Platelets, bilirubin, ALT, CMV-DNA by PCR monthly <u>OR</u> <input type="checkbox"/> every 3 months, as clinically indicated <input type="checkbox"/> Consults: <input type="checkbox"/> Other tests: <input type="checkbox"/> See general orders sheet for additional requests.	
DOCTOR'S SIGNATURE:	SIGNATURE:
	UC: