Corewell Health

Physician's Orders RITUXIMAB OR BIOSIMILAR,	Physician CSN	
EVERY 14 DAYS TIMES 2 DOSES -		
ADULT, OUTPATIENT, COREWELL HEALTH I	NFUSION CENTER	
Page 1 to 4		

Patient Name

DOB MRN

Anticipated Infusion Date	ICD 10 Code with D	escription	
Height(cm)	Weight(kg) Allergies	S	
Site of Service			
CH Gerber	CH Lemmen Holton (GR)	CH Pennock	CH Greenville
□ CH Helen DeVos (GR)	CH Ludington	CH Reed City	CH Zeeland
CH Blodgett (GR)			
Provider Specialty			
Allergy/Immunology	Infectious Disease	□ OB/GYN	Rheumatology
Cardiology	Internal Med/Family Practice	□ Other	□ Surgery
□ Gastroenterology	Nephrology	Otolaryngology	Urology
Genetics	Neurology	Pulmonary	Wound Care
Treatment Intent			
Conditioning	Curative	□ Mobilization	Supportive
Control	Maintenance	Palliative	

Cycle 1

One dose every 14 days x 2 doses on day 1, 15

Provider Reminder

ONC PROVIDER REMINDER 2

Confirm that the appropriate informed consents have been signed and are in the medical record. Appointment Requests

ONCBCN CALCULATED LENGTH INFUSION APPOINTMENT REQUEST 1

Safety Parameters and Special Instructions

ONC SAFETY PARAMETERS AND SPECIAL INSTRUCTIONS 3

RITUXIMAB DOSING INTERVAL - Rheumatoid arthritis: IV: 1,000 mg on days 1 and 15 (in combination with methotrexate); subsequent courses may be administered every 24 weeks (based on clinical evaluation), if necessary may be repeated no sooner than every 16 weeks

See dosing guidelines for other clinical indications as there variations in dosing interval depending on clinical indication.

Labs

COMPLETE BLOOD COUNT (CBC) W/DIFFERENTIAL

Pregnancy tests recommended for Females aged 12 to 60 with Uterus intact.

Please order as appropriate for clinical presentation

Interval: Once Selection conditions: Patient could become pregnant

HCG, QUANTITATIVE

ONC PROVIDER REMINDER

Review prior hepatitis labs and consider repeat screening as indicated.

HEPATITIS PANEL

Selection conditions: Hides NCCN Hepatitis Screening orders in background if the patient has had a completed hepatitis lab within the last 365 days.

HEP B CORE TOTAL AB

Selection conditions: Hides NCCN Hepatitis Screening orders in background if the patient has had a completed hepatitis lab within the last 365 days.

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NOTE: Epic Treatment/Therapy Plan Orders. To be scanned/attached to the appropriate Infusion Referral Order in Epic.



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Patient Name
DOB
MRN
Physician

RITUXIMAB OR BIOSIMILAR, EVERY 14 DAYS TIMES 2 DOSES -ADULT, OUTPATIENT, COREWELL HEALTH INFUSION CENTER (CONTINUED) Page 2 to 4

	VITAL SIGNS				
	RITUXIMAB - Dose 1 & 2: Vital signs every 15 minutes x 2 hours, then hourly throughout				
	infusion. Subsequent doses: Vital signs with rate changes.				
Nursin	g Orders				
	HYPERSENSITIVITY REACTION ADULT ONCOLOGY PROTOCOL				
	ONC NURSING COMMUNICATION 100				
	May Initiate IV Catheter Patency Adult Protocol				
Treatn	nent Parameters				
	ONC MONITORING AND HOLD PARAMETERS 3				
	May proceed with treatment if hepatitis B core antibody and surface antigen labs have been resulted prior to the first dose, and the results are negative.				
Pre-M	edications				
110 10	Oacetaminophen (Tylenol) tablet				
	Dose: Route: Oral Once for 1 dose				
	□ 325 mg □ 650 mg				
	□ 500 mg □ 1000 mg Instructions: Administer 30 minutes prior to rituximab.				
	OdiphenhydrAMINE (Benadryl) injection				
	Dose: Route: Oral Once for 1 dose				
	□ 25 mg □ 50 mg Instructions: Administer 30 minutes prior to rituximab.				
Pre-M	edications				
	ONC PROVIDER REMINDER 7				
	HISTORY OF INFUSION REACTION: consider adding CORTICOSTEROID pre-medication pric to riTUXimab or biosimilar				
	○ dexAMETHasone (Decadron) injection 10 mg				
	Dose: 10 mg Route: Intravenous Once over 5 Minutes for 1 dose				
	Instructions:				
	For patient with a history of reaction to riTUXimab or biosimilar. Administer 30 minutes prior to start of riTUXimab or biosimilar.				
	O methylPREDNISolone sodium succinate (SOLU-Medrol) injection 125 mg				
	Dose: 125 mg Route: Intravenous Once for 1 dose				
	Instructions:				
	For patient with a history of reaction to riTUXimab or biosimilar. Administer 30 minutes prior to start of riTUXimab or biosimilar.				
	O hydrocortisone sodium succinate (PF) (Solu-CORTEF) injection 100 mg				
	Dose: 100 mg Route: Intravenous Once for 1 dose Instructions:				
	For patient with a history of reaction to riTUXimab or biosimilar. Administer 30 minutes prior to start of riTUXimab or biosimilar.				
Monod	clonal Antibody / Biotherapy				
	riTUXimab-pvvr (RUXIENCE) 1,000 mg in sodium chloride 0.9% 500 mL (2 mg/mL) infusio				
	Dose: 1,000 mg Route: Intravenous Titrate @ 25-200 mL/hr for 1 dose Instructions:				
	Before start of riTUXimab-pvvr, infusion, prime 0.9% sodium chloride 250 mL with primary tubing and have available at bedside in case of hypersensitivity or infusion reaction. Do NOT connect this saline with any other intravenous drugs, fluids, or blood products.				
	Start infusion at 50 mg/br (25 ml /br). For patients tolerating the infusion well, the rate of				
	Start infusion at 50 mg/hr (25 mL/hr). For patients tolerating the infusion well, the rate of				

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RITUXIMAB OR BIOSIMILAR, EVERY 14 DAYS TIMES 2 DOSES -ADULT, OUTPATIENT, COREWELL HEALTH INFUSION CENTER (CONTINUED) Page 3 to 4

infusion may be increased in increments of 50 mg/hr (25 mL/hr) at intervals of 30 minutes to a maximum rate of 400 mg/hr (200 mL/hr).

SUBSEQUENT INFUSIONS: If patient tolerated initial infusion, start at 100 mg/hour (50 mL/hr); if there is no infusion-related reaction, increase the rate by 100 mg/hour (50 mL/hr) increments every 30 minutes, to a maximum rate of 400 mg/hour (200 mL/hr)

riTUXimab-abbs (TRUXIMA) 1,000 mg in sodium chloride 0.9% 500 mL (2 mg/mL) infusion

Dose: 1,000 mgRoute: IntravenousTitrate @ 25-200 mL/hr for 1 doseInstructions:

Before start of riTUXimab-abbs, infusion, prime 0.9% sodium chloride 250 mL with primary tubing and have available at bedside in case of hypersensitivity or infusion reaction. Do NOT connect this saline with any other intravenous drugs, fluids, or blood products.

Start infusion at 50 mg/hr (25 mL/hr). For patients tolerating the infusion well, the rate of infusion may be increased in increments of 50 mg/hr (25 mL/hr) at intervals of 30 minutes to a maximum rate of 400 mg/hr (200 mL/hr).

SUBSEQUENT INFUSIONS: If patient tolerated initial infusion, start at 100 mg/hour (50 mL/hr); if there is no infusion-related reaction, increase the rate by 100 mg/hour (50 mL/hr) increments every 30 minutes, to a maximum rate of 400 mg/hour (200 mL/hr)

riTUXimab (RITUXAN) 1,000 mg in sodium chloride 0.9% 500 mL (2 mg/mL) infusion

Dose: 1,000 mg Route: Intravenous Titrate @ 25-200 mL/hr for 1 dose Instructions:

Before start of riTUXimab, infusion, prime 0.9% sodium chloride 250 mL with primary tubing and have available at bedside in case of hypersensitivity or infusion reaction. Do NOT connect this saline with any other intravenous drugs, fluids, or blood products.

Start infusion at 50 mg/hr (25 mL/hr). For patients tolerating the infusion well, the rate of infusion may be increased in increments of 50 mg/hr (25 mL/hr) at intervals of 30 minutes to a maximum rate of 400 mg/hr (200 mL/hr).

SUBSEQUENT INFUSIONS: If patient tolerated initial infusion, start at 100 mg/hour (50 mL/hr); if there is no infusion-related reaction, increase the rate by 100 mg/hour (50 mL/hr) increments every 30 minutes, to a maximum rate of 400 mg/hour (200 mL/hr)

riTUXimab-arrx (RIABNI) 1,000 mg in sodium chloride 0.9% 500 mL (2 mg/mL) infusion

Dose: 1,000 mg Route: Intravenous Titrate @ 25-200 mL/hr for 1 dose Instructions:

Before start of riTUXimab-arrx infusion, prime 0.9% sodium chloride 250 mL with primary tubing and have available at bedside in case of hypersensitivity or infusion reaction. Do NOT connect this saline with any other intravenous drugs, fluids, or blood products.

Start infusion at 50 mg/hr (25 mL/hr). For patients tolerating the infusion well, the rate of infusion may be increased in increments of 50 mg/hr (25 mL/hr) at intervals of 30 minutes to a maximum rate of 400 mg/hr (200 mL/hr).

SUBSEQUENT INFUSIONS: If patient tolerated initial infusion, start at 100 mg/hour (50 mL/hr); if there is no infusion-related reaction, increase the rate by 100 mg/hour (50 mL/hr) increments every 30 minutes, to a maximum rate of 400 mg/hour (200 mL/hr)

Supportive Care

prochlorperazine (COMPAZINE) tablet 10 mg

Dose: 10 mg Route: Oral Every 6 hours PRN Instructions: Use 1st line for nausea/vomiting for patients able to tolerate oral medications.

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U Health [™]	DOB
RITUXIMAB OR BIOSIMILAR,	Physician CSN
EVERY 14 DAYS TIMES 2 DOSES -	

Patient Name

EVERY 14 DAYS TIMES 2 D ADULT, OUTPATIENT, COREWELL HEALTH INFUSION CENTER (CONTINUED) Page 4 to 4

	prochlorperazine (COMPAZINE) injection 10 mg				
	Dose: 10 mg Route: Intravenous Every 6 hours PRN over 5 Minutes Instructions: Use 1st line for nausea/vomiting for patients unable to tolerate oral medications.				
ondansetron (ZOFRAN-ODT) disintegrating tablet 8 mg					
	Dose: 8 mg Route: Oral Every 8 hours PRN Instructions: Use 2nd line for Nausea/Vomiting. May give IV if unable to tolerate PO.				
	ondansetron (ZOFRAN) injection 8 mg				
	Dose: 8 mg Route: Intravenous Every 8 hours PRN over 5 Minutes Instructions: Use 2nd line for Nausea/Vomiting. May give IV if unable to tolerate PO.				

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Telephone order/Verbal order documented and read-back completed. Practitioner's initials _

NOTE: Unless Order is written DAW (dispense as written), medication may be supplied which is a generic equivalent by nonproprietary name.

TRANSCRIBED:		VALIDATED:		ORDERED:			
TIME	DATE	TIME	DATE	TIME	DATE	Pager #	
			R.N.		Physic	ian	Physician
	Sign		Sign		P	rint	Sign