



PATIENT INFORMATION (Complete or Fax Existing Chart)		PRESCRIBER INFORMATION	
Name: _____ DOB: _____		Prescriber Name: _____	
Address: _____		State License: _____	
City, State, Zip: _____		NPI #: _____ Tax ID: _____	
Phone: _____ Alt. Phone: _____		Address: _____	
Email: _____ SS#: _____		City, State, Zip: _____	
Gender: ☐ M ☐ F Weight: _____ (lbs) Ht: _____		Phone: _____ Fax: _____	
Allergies: _____		Office Contact: _____ Phone: _____	
INSURANCE INFORMATION – AND – Send a copy of the patient's prescription/insurance cards (front & back)			
Primary Insurance: _____		Secondary Insurance (If Applicable): _____	
Plan #: _____		Plan #: _____	
Group #: _____		Group #: _____	
RX Card (PBM): _____		RX Card (PBM): _____	
BIN: _____ PCN: _____		BIN: _____ PCN: _____	
CLINICAL INFORMATION			
ICD-10 Code(s) & Description: _____			
PI: Attach Labs		CIDP: Attach reports	
IgA Level (mg/dL): _____ Pre-Titer Level (mcg/mL): _____		EMG/NCS/Nerve Ultrasound (m/sec): _____	
IgG Level (mg/dL): _____ IgM Level (mg/dL): _____		NF155 Levels: _____ CNTN1 Levels: _____	
Post Titer Level mcg/mL: _____		MRI Results: _____	
ORDERS			
Prescription type: ☐ New start ☐ Restart ☐ Continued therapy Total Doses Received: _____ Date of Last Injection/Infusion: _____			
Medication	Indication		
☐ HyQvia® [Immune Globulin Infusion 10% (Human) with Recombinant Human Hyaluronidase] Solution.	☐ For PI: ☐ If switching from IVIG (human) treatment, administer HYQVIA at the same dose and frequency as the previous IV treatment, after the initial dose ramp-up ☐ If naive to SCIG (human) treatment or switching from SCIG, administer HYQVIA at 300 mg/kg to 600 mg/kg at 3-week or 4-week intervals, after the initial ramp-up ☐ For CIDP: ☐ If switching from IVIG (human) treatment, administer HYQVIA at the same dose and frequency as the previous IV treatment, after the initial dose ramp-up		
Dose/Frequency			
☐ Infusion provider to calculate ramp-up dose per the ramp-up			
Patient weight (kg): _____ x dose (mg/kg): _____ Frequency: ☐ every 2 weeks ☐ every 3 weeks ☐ every 4 weeks Refills: _____			
Number of infusion site(s): ☐ 1 ☐ 2 ☐ 3 Infusion site(s): ☐ Middle to upper abdomen ☐ Thigh(s)			
High-flow 24G needle length (check one): ☐ 6mm ☐ 9mm ☐ 12mm ☐ 14mm			
Pre-Medication	Dose/Strength	Directions	
☐ Acetaminophen	☐ 500mg	☐ Take 1-2 tablets PO prior to infusion or post-infusion as directed	
☐ Diphenhydramine	☐ 25mg ☐ 50mg	☐ Take 1 tablet PO prior to infusion or as directed OR	
☐ _____	_____	_____	

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INFUSION REACTION ORDERS

Mild reaction protocol:

☒ Diphenhydramine 25mg IV, one time, for pruritus.

If symptoms worsen, see orders for moderate to severe reactions.

Moderate reaction protocol:

☒ Acetaminophen 650mg PO, one time, for pyrexia or rigors

☒ Diphenhydramine 50mg IV, one time, for pruritus or urticaria

☒ Methylprednisolone 125mg IV, one time, for respiratory or neurologic symptoms

If symptoms worsen, see interventions for severe reactions

Severe reaction protocol: (Call 911 if initiated):

☒ Titrate oxygen via continuous flow per nasal cannula or face mask to maintain spO2 of greater than ninety-five percent (>95%)

☒ Diphenhydramine 50mg IV, one time, for respiratory symptoms, edema, or anaphylaxis

☒ Methylprednisolone 125mg IV, one time, for respiratory symptoms, edema, or anaphylaxis

☒ Sodium Chloride 0.9% 500mL IV over 30-60 min, one time, for cardiovascular symptoms

☒ Epinephrine 0.3mg/0.3mL IM into mid-anterolateral aspect of thigh of anaphylaxis, may repeat x1 in 5-15 minutes if symptoms are not resolved or worsen

FLUSHING & LOCKING ORDERS

Flushing Protocol (>66lbs/33kg)

PIV and Midline:

☒ 0.9% Sodium Chloride 2-5mL IV flush before and after each infusion

Implanted Port, PICC, Tunneled Catheter, and Non-tunneled Catheter:

☒ 0.9% Sodium Chloride 5mL IV flush before infusion/lab draw and 10mL IV flush after infusion/lab draw

Locking Protocol (>66lbs/33kg)

PIV and Midline:

☒ Heparin Sodium 10 units/mL 1mL IV final flush post normal saline flush

PICC:

☒ Heparin Sodium 10 units/mL 3mL IV final flush post normal saline flush

Implanted Port, Tunneled Catheter, and Non-tunneled Catheter:

☒ Heparin Sodium 100 units/mL 3-5mL IV final flush post normal saline flush

**** May substitute Dextrose 5% in Water, or alternative, for 0.9% Sodium Chloride, when indicated due to incompatibility with medications being infused**

SIGNATURE

We hereby authorize Talis Healthcare LLC to provide all supplies and additional services (nursing/patient training) required to provide and deliver the medicine as prescribed in this referral.

X _____

Prescriber Signature

Date: _____

To ensure payment by insurance carrier, please include supporting clinical documentation for specified ICD 10 Code, demographic, and insurance information along with faxed order. Initial appointment will be verified upon insurance approval.

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