

PATIENT INFORMATION

Patient Name: _____ DOB: _____ Phone: _____ Sex: M F Ht: _____ Wt: _____ lbs kg
 Primary Language: _____ Allergies: _____
 Patient Preferred Location: _____

<ICD 10 CODE REQUIRED> DIAGNOSIS & CLINICAL INFORMATION

ICD 10 Code (PROVIDE COMPLETE CODE)

D80. _____ Hypogammaglobulinemia
 D81. _____ Combined Immunodeficiency
 D82.0 Wiskott-Aldrich Syndrome
 D83. _____ Common Variable Immune Deficiency
 Other: _____

Prescribing Information

For patients previously on another IG treatment, it is recommended to administer the first dose approximately one week after the last infusion of their previous treatment

If applicable:

Previous IG Therapy: _____
 Date of Last Dose: _____

REQUIRED: Demographics & Most Recent: H&P, clinical notes, & medication list. Supporting clinical notes to include any past tried and/or failed therapies, intolerance, outcomes, or contraindications to conventional therapy.
LAB RESULTS: IG Levels

PRESCRIPTION

HYQVIA (Immune Globulin SubQ Infusion)

Subcutaneous Immune Globulin Infusion 10% with Recombinant Human Hyaluronidase

Subcutaneous Administration Only as tolerated. Hyaluronidase to infuse first at 1-2 mL/minute/site

SELECT ONE

Ramp up & Maintenance Dose

Patient is new to therapy, follow ramp up scheduling per chart with the indicated dose, then continue to maintenance as indicated

Maintenance Loading Dose Only

Patient is currently on therapy and will continue as indicated above

Is the patient on any other disease modifying therapy? **Yes No**
 If yes, please note therapy and last dose: _____

TREATMENT INTERVAL	DOSING FREQUENCY Q4 WEEK	DOSING FREQUENCY Q3 WEEK
1st Infusion (week 1)	Grams x 0.25	Grams x 0.33
2nd Infusion (week 2)	Grams x 0.5	Grams x 0.67
3rd Infusion (week 4)	Grams x 0.75	Administer Total Grams
4th Infusion (week 7)	Administer Total Grams	n/a

Post Treatment Observations: The patient is observed for 30 minutes following the first administration.

Adverse Events: In the event of an adverse reaction occurring at a Medix Infusion suite, utilize the Medix Infusion adverse reaction protocol.

Comments:

PRESCRIBER INFORMATION

Prescriber Name: _____ Signature: _____
 Date: _____ NPI #: _____ Specialty: _____
 Supervising Physician: _____ (If Applicable)
 Address: _____ City: _____ State: _____ Zip: _____
 Contact Name: _____ Phone: _____ Fax: _____ Email: _____