



## HYQVIA Patient Start Form

Fax pages 1-4 to **1-866-861-1752** | Phone: **1-866-861-1750** 

Please ensure patient reads and signs pages 3 and 4 for appropriate authorizations.

Name (First, Last):		State License #: ——	
		PTAN #:	
Street Address:	City: -		State: ZIP:
Office Contact:			
Telephone:	Fax:		Email:
2 Patient Information			Male Female
Patient Name (First, Middle Initial, Last): —			
DOB (MM/DD/YYYY):	Last 4 Digits of Social	Security #:	Email:
Street Address:			
City:	State:		ZIP:
Mobile Telephone:		— Home Telephone: —	
Caregiver Name (First, Last):		Relationship to Patie	nt:
Caregiver Telephone:	Caregiver Email:		
3 Insurance Information	Please attach copies of both s medical and prescription insu		Check if patient does not have insurance.
Primary Insurance:	Pharmacy Plan Nan	ne:	Secondary Insurance:
_ Insurance Telephone:	Pharmacy Plan Tele	phone:	Insurance Telephone:
Policy ID #:	Policy ID #:		Policy ID #:
Group ID #:	Group ID #:		Group ID #:
Policy Holder Name:	RX BIN #:		Policy Holder Name:
Policy Holder DOB (MM/DD/YYYY):	RX PCN #:		Policy Holder DOB (MM/DD/YYYY):





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4 Diagnosis/Medical As	sessment	Diagnosis (ICD-10):	
PI		_ CIDP	
_ IgA Level (mg/dL):	Pre-Titer Level (mcg/mL):	EMG/NCS/Nerve Ultrasound (m/sec): NF155 Levels: CN	ITN1 Levels: -
- IgG Level (mg/dL):	IgM Level (mg/dL):	MRI Results:	
Post Titer Level (mcg/mL):			
5 HYQVIA Prescription,	Training Request/Wai	ver, and Prescribing Physician Signature	
Please see Important Safety Information	on on page 6 and click for <u>Full Presc</u>	<u>cribing Information</u> including Boxed Warning regarding Thrombos	is.
– Name (First, Middle Initial, Last): —		DOB (MM/DD/YYYY): Patient is already	on HYQVI
Prescription: HYQVIA® [Immune Glo Choose an indication below and ca	. ,	th Recombinant Human Hyaluronidase] Solution.	
· For PI —————		For CIDP —	
If switching from IVIG (human) treatmer and frequency as the previous IV treatm  If naive to SCIG (human) treatment or sw 300 mg/kg to 600 mg/kg at 3-week or 4-	ent, after the initial dose ramp-up. <sup>1</sup> itching from SCIG, administer HYQVIA at	If switching from IVIG (human) treatment, administer HYQVIA at the and frequency as the previous IV treatment, after the initial dose rail	
	- Soo ramp-up schodulo tables or	page 5 to calculate ramp-up dosage.	
Patient weight (kg) <b>X</b> We	(mg/kg)	grams* Weekly dose X2 for even the work of	ery 3 weeks
Pharmacy to calculate ramp-up per the ramp-up schedule in th Prescribing Information.	ne Full Infusion site(s)†:	sion site(s): 1 2 3 Weekly dose X4 for every 3 Widdle to upper abdomen Thigh(s)	ery 4 weeks
Refills (as allowed bor payer requireme	•		14 mm
To calculate total infusion volume in mL, multiply f 2 infusion sites are used, the infusion sites should	-	es, the sites should be 10 cm apart. Avoid bony prominences or areas that are scarred, in	nflamed, or infe
Prescriber additional instruc	tion:		
No known Patient alle drug allergies (drug and i			
Special instructions:			
Preferred site of care if not self-ad	ministered (check one)	- Has a referral been sent to site of care? Yes No	o
Infusion suite Begin treatment in	clinical setting, then transition to home of	are Prescriber's office Home infusion Hospital ou	utpatient
– Preferred Specialty Pharmacy: ––––		Preferred Infusion Suite/Hospital Outpatient (if applicable):	
Information and will be supervising Patient's to applicable federal and state law regulations, re its agents or contractors, for the purpose of se	reatment. I have received from Patient, or I ferenced medical and/or other patient info eking information related to coverage and/ armacy designated by me, Patient, or Patie	atient identified in this application ("Patient"). I have reviewed the current HYQ\ inis/her personal representative, the necessary authorization to release, in accord rmation relating to HYQVIA therapy to Takeda Pharmaceutical Company Limited for assisting in initiating or continuing HYQVIA therapy. I authorize Takeda Patien ent's plan. I agree that product provided through the Program shall only be used f	dance with I, including It Support to
Prescriber Signature (Required) Stamps not	acceptable		

The prescriber is required to comply with their state-specific prescription requirements such as e-prescribing, state-specific prescription form, fax language, etc. Non-compliance with state-specific requirements could result in delay.



HyQvia [Immune Globulin Infusion 10% (Human) with Recombinant Human Hyaluronidase]

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6 Patient HIPAA Authorization	
Patient Name (First, Middle Initial, Last):	DOB (MM/DD/YYYY):

By signing the Patient Authorization section on the third page of this Takeda Patient Support Ig **Enrollment Form, I authorize my physician, health insurance, and pharmacy providers (including** any specialty pharmacy that receives my prescription) to disclose my protected health information, including, but not limited to, information relating to my medical condition, treatment, care management, and health insurance, as well as all information provided on this form ("Protected Health Information"), to Takeda Pharmaceuticals U.S.A., Inc. and its present or future affiliates, including the affiliates and service providers that work on Takeda's behalf in connection with the Takeda Patient Support, Ig Patient Support Program (the "Companies"). The Companies will use my Protected Health Information for the purpose of facilitating the provision of the Takeda Patient Support, Ig Patient Support Program products, supplies, or services as selected by me or my physician and may include (but not be limited to) verification of insurance benefits and drug coverage, prior authorization education, financial assistance with co-pays, patient assistance programs, and other related programs. Specifically, I authorize the Companies to 1) receive, use, and disclose my Protected Health Information in order to enroll me in Takeda Patient Support, Ig and contact me, and/or the person legally authorized to sign on my behalf, about Takeda Patient Support, Ig. 2) provide me, and/or the person legally authorized to sign on my behalf, with educational materials, information, and services related to Takeda Patient Support, Iq; 3) verify, investigate, and provide information about my coverage for HYQVIA, including but not limited to communicating with my insurer, specialty pharmacies, and others involved in processing my pharmacy claims to verify my coverage; 4) coordinate prescription fulfillment; and 5) use my information to conduct internal analyses. I understand that employees of the Companies only use my Protected Health Information for the purposes described herein, to administer the Takeda Patient Support, Iq Patient Support Program or as otherwise required or allowed under the law, unless information that specifically identifies me is removed. Further, I understand that my physician, health insurance, and pharmacy providers may receive financial remuneration from the Companies for providing Protected Health Information, which may be used for marketing purposes. I understand that Protected Health Information disclosed under this Authorization may no longer be protected by federal privacy law. I understand that I am entitled to a copy of this Authorization. I understand that I may revoke this Authorization and that instructions for doing so are contained in Takeda's Website Privacy Notice available at www.takeda.com/privacy-notice/ or I may revoke this Authorization at any time by sending written notice of revocation to Takeda Patient Services 610 Crescent Executive Court, Suite 200 Lake Mary, FL 32746. I understand that such revocation will not apply to any information already used or disclosed through this Authorization. This Authorization will expire within five (5) years from the date it is signed and provided on the first page of this enrollment form, unless a shorter period is provided for by state law. I understand that I may refuse to sign this Authorization and that refusing to sign this Authorization will not change the way my physician, health insurance, and pharmacy providers treat me. I also understand that if I do not sign this Authorization, I will not be able to receive Takeda Patient Support, Ig Patient Support Program products, supplies, or services.

Signature of Patient (Required)		*Legal Representative Name:	Date
*Legal Representative Signature	Date	*Relationship to Patient:	

<sup>\*</sup>Required only if applicable.



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atient Name (First, Middle Initial, Last):	DOB (MM/DD/YYYY):
ditional services	
litional services and infusion training are available.	
eck the box next to any of the services below. Check	the last box if the patient opts out.
Pharmacy to provide needles, syringes, durable medical equipment, and other ancillary supplies needed for infusion	Training  If HYQVIA is intended for self-administration or administration by a caregiver, the
Pharmacy to provide anaphylactic kit:	patient or caregiver should be trained by a healthcare professional. Takeda Patient Support provides free infusion training services to all enrolled HYQVIA patients.
	If you choose to opt out of these services, please check this bo
	in you choose to ope out of these services, piease check this se
REQUIRED:	
Takeda Patient Support Enrollment	
By cigning helow, Lam electing to enroll in Takeda Patient Support Ser	vices ("Services") and direct all disclosures of my Information in connection with such
	rrance benefits and drug coverage, prior authorization support, financial assistance with
co-pays, patient assistance programs, alternate funding sources, other	er related programs, communication with me or my prescribing physician by mail, email,
or telephone about my medical condition, treatment, care management	nt, product information, and health insurance).
Signature of Patient (Required)/*Legal Re	presentative Signature
Signature of Patient (Required)/*Legal Re	presentative Signature
Signature of Patient (Required)/*Legal Re	presentative Signature
Signature of Patient (Required)/*Legal Re	presentative Signature
Signature of Patient (Required)/*Legal Report Text Communication Agreement Terms	
Text Communication Agreement Terms  By agreeing to these Takeda Patient Support ( "Program") text message terms and or below. You also consent to receive autodialed and/or prerecorded calls and/or text message.	5 & Conditions  conditions, you agree to receive text messages on your mobile device subject to the Terms & Conditions described nessages from or on behalf of the Program at the telephone number provided above. You understand that this consent vice. Such messages may be nonmarketing messages related to the Patient Support Program. There is no fee
Text Communication Agreement Terms  By agreeing to these Takeda Patient Support ( "Program") text message terms and of below. You also consent to receive autodialed and/or prerecorded calls and/or text m is not a condition of purchase or use of the Program or of any Takeda product or sern payable to Takeda to receive text messages; however, your carrier's message and do You represent that you are the account holder for the mobile telephone number (syour mobile telephone number. You may notify Takeda of a number change by conservice may include your phone number and/or email address, related carrier in	conditions, you agree to receive text messages on your mobile device subject to the Terms & Conditions described nessages from or on behalf of the Program at the telephone number provided above. You understand that this consent vice. Such messages may be nonmarketing messages related to the Patient Support Program. There is no fee atar rates may apply.  (s) that you provide to opt in to the Program. You are responsible for notifying Takeda immediately if you change alling 1-866-861-1750. Data obtained from you in connection with your registration for, and use of, this SMS formation, and elements of pharmacy claim information and will be used to administer this Program and to
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<sup>\*</sup>Required only if applicable.





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## Instructions for Completion of Form

- Complete sections 1-6 and FAX PAGES 1-4 to 1-866-861-1752 and attach a copy of the patient's insurance card (front and back)
- Do not submit to Takeda any documentation of labs, clinical history, or other documents supporting the prior authorization process



# Prescribing Physician Information



Patient Information



Insurance Information



Diagnosis/Medical Assessment



## HYQVIA Prescription, Training Request/Waiver, and Prescribing Physician Signature

- Please indicate the number of refills
- Check the appropriate box to specify whether you would like your patient to be trained by Takeda on self-administration or whether training has already occurred
- This is a prescription; a physician's signature and date are required

Infusion Parameters for Recombinant Human Hyaluronidase (Hy) and Immune Globulin Infusion 10% (Rate of Administration for Hy: 1-2 mL/min/site(s), and increase as tolerated				
Rate of Administration for Ig:	Patients <4	10 kg (<88 lb)	Patients ≥40 kg (≥88 lb)	
	First 2 Infusions	Subsequent 2 or 3 Infusions	First 2 Infusions	Subsequent 2 or 3 Infusions
Interval (Minutes)	Rate/site (mL/hour)	Rate/site (mL/hour)	Rate/site (mL/hour)	Rate/site (mL/hour)
5-15	5	10	10	10
5-15	10	20	30	30
5-15	20	40	60	120
5-15	40	80	120	240
Remainder of Infusion	80	160	240	300

#### Initial Treatment Interval and Ramp-Up Schedule for PI For patients previously on another IgG treatment, the first dose should be given approximately 1 week after the last infusion of their previous treatment. Ramp-up schedule if switching from SCIG PI: Ramp-up schedule if switching from IVIG Treatment Interval Dose Interval Dose 1st week | Total grams x 0.25 | Total grams x 0.33 Switch from IVIG 1st dose Total grams x 0.25 2nd week | Total grams x 0.50 | Total grams x 0.67 2nd dose Total grams x 0.50 No Infusion 3rd dose Total grams x 0.75 3rd Infusior 4th week Total grams x 0.75 Total grams No Infusion No Infusion Total grams 7th week 4th dose Total grams

#### Initial Treatment Interval and Ramp-Up Schedule for CIDP . Doses less than or equal to 0.4 g/kg can be administered without ramp-up Patients must be on stable doses of IVIG for 12 weeks before switching to HYQVIA Ramp-up schedule if switching from IVIG Dose Interval Dose Week Switch from IVIG No Infusion 1st dose Total grams x 0.25 2nd dose Total grams x 0.25 3rd dose Total grams x 0.50 4th dose Total grams x 0.75 5th dose Total grams Total grams=total monthly equivalent dose in grams

## 6

## Patient HIPAA Authorization and Takeda Patient Support Enrollment

The patient signature is required to allow personal health information to be shared by third parties to Takeda to facilitate access to HYQVIA (insurance benefits, self-administration training, transfer Rx to specialty pharmacy provider, etc.).

## Checking the Takeda Patient Support Enrollment box allows patients to receive product support services from Takeda, if eligible

• Benefits investigation

Total grams=total monthly equivalent dose in grams

- Infusion training (if applicable)
- Co-pay support (when applicable) and information about third-party financial assistance programs, as necessary
- Enrollment in Takeda Patient Support—Patient Support Manager assignment and product support services

## What happens next?

- Once the completed form has been submitted to Takeda Patient Support, a dedicated Patient Support Manager will be assigned to your eligible patient
- The Patient Support Manager will contact the patient directly to inform him or her of the services available through Takeda Patient Support and to begin the insurance verification process
- The Patient Support Manager will work with the insurance company to determine insurance benefits
- The Patient Support Manager will assess the patient's eligibility for co-pay support (when applicable) and provide information about third-party financial assistance programs, as necessary
- If requested, the Patient Support Manager will set up Takeda-provided self-administration training services

Please see Important Safety Information on page 6 and click for <u>Full Prescribing Information</u> including Boxed Warning regarding Thrombosis.

HyQvia [Immune Globulin Infusion 10% (Human) with Recombinant Human Hyaluronidase]

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### **INDICATIONS**

HYQVIA is indicated for the treatment of primary immunodeficiency (PI) in adults and pediatric patients two years of age and older and for chronic inflammatory demyelinating polyneuropathy (CIDP) as maintenance therapy to prevent relapse of neuromuscular disability and impairment in adults. HYQVIA is for subcutaneous use only.

### IMPORTANT SAFETY INFORMATION

#### WARNING: THROMBOSIS

- Thrombosis may occur with immune globulin (IG) products, including HYQVIA. Risk factors may include advanced age, prolonged immobilization, hypercoagulable conditions, history of venous or arterial thrombosis, use of estrogens, indwelling vascular catheters, hyperviscosity, and cardiovascular risk factors.
   Thrombosis may occur in the absence of known risk factors.
- For patients at risk of thrombosis, administer HYQVIA at the minimum dose and infusion rate practicable. Ensure adequate hydration in patients before administration.
- Monitor for signs and symptoms of thrombosis and assess blood viscosity in patients at risk of hyperviscosity.

#### **Contraindications**

- History of anaphylactic or severe systemic hypersensitivity reactions to human IG
- IgA-deficient patients with antibodies to IgA and a history of hypersensitivity to human IG
- Known systemic hypersensitivity to hyaluronidase including Recombinant Human Hyaluronidase of HYQVIA
- Known systemic hypersensitivity to human albumin (in the hyaluronidase solution)

## **Warnings and Precautions**

**Hypersensitivity:** Severe hypersensitivity reactions may occur, even in patients who have tolerated previous treatment with human IG. If a hypersensitivity reaction occurs, discontinue infusion immediately and institute appropriate treatment. IgA-deficient patients with antibodies to IgA are at greater risk of developing potentially severe hypersensitivity reactions, including anaphylaxis.

**Thrombosis:** Has been reported to occur following treatment with IG products, including HYQVIA and in the absence of known risk factors. In patients at risk, administer at the minimum dose and infusion rate practicable. Ensure adequate hydration before administration. Monitor for signs and symptoms of thrombosis and assess blood viscosity in patients at risk for hyperviscosity.

Immunogenicity of Recombinant Human Hyaluronidase (rHuPH20): Non-neutralizing antibodies to the Recombinant Human Hyaluronidase component can develop. The clinical significance of these antibodies or whether they interfere with fertilization in humans is unknown.

**Aseptic Meningitis Syndrome:** Has been reported with use of IG, including HYQVIA and may occur more frequently in females. The syndrome usually begins within several hours to two days following IG treatment

Conduct a thorough neurological exam on patients exhibiting signs and symptoms, to rule out other causes of meningitis. Discontinuing IG treatment has resulted in remission within several days without sequelae.

**Hemolysis:** HYQVIA contains blood group antibodies which may cause a positive direct antiglobulin reaction and hemolysis. Monitor patients for signs and symptoms of hemolysis and delayed hemolytic anemia and, if present, perform appropriate confirmatory lab testing.

Renal Dysfunction/Failure: Acute renal dysfunction/failure, acute tubular necrosis, proximal tubular nephropathy, osmotic nephrosis, and death may occur with intravenous (IV) use of IG products, especially those containing sucrose. Ensure patients are not volume depleted prior to infusion. In patients at risk due to pre-existing renal insufficiency or predisposition to acute renal failure, assess renal function before initiation and throughout treatment, and consider lower, more frequent dosing. If renal function deteriorates, consider discontinuation.

**Spread of Localized Infection:** Do not infuse HYQVIA into or around an infected area due to potential risk of spreading a localized infection

**Transfusion-Related Acute Lung Injury:** Non-cardiogenic pulmonary edema may occur with IV administered IG. Monitor patients for pulmonary adverse reactions. If suspected, perform appropriate tests for presence of anti-neutrophil and anti-HLA antibodies in both product and patient serum. May be managed using oxygen therapy with adequate ventilatory support.

**Transmittable Infectious Agents:** Because HYQVIA is made from human plasma, it may carry a risk of transmitting infectious agents (e.g. viruses, other pathogens). No cases of transmission of viral diseases or variant Creutzfeldt-Jakob disease (vCJD) have been associated with HYQVIA.

Interference with Lab Tests: False positive serological test results and certain assay readings, with the potential for misleading interpretation, may occur as the result of passively transferred antibodies.

#### **Adverse Reactions**

The most common adverse reactions observed in >5% of patients in the clinical trials were:

<u>Primary Immunodeficiency (PI)</u>: local adverse reactions including pain, erythema, edema, and pruritus, and systemic adverse reactions including, headache, antibody formation against Recombinant Human Hyaluronidase (rHuPH20), fatigue, nausea, pyrexia, and vomiting.

<u>Chronic Inflammatory Demyelinating Polyneuropathy (CIDP)</u>: local reactions, headache, pyrexia, nausea, fatigue, erythema, pruritus, increased lipase, abdominal pain, back pain, and pain in extremity.

#### **Drug Interactions**

Passive transfer of antibodies may transiently interfere with the immune responses to live attenuated virus vaccines (e.g., measles, mumps, rubella, and varicella).

### **Use In Specific Populations**

**Pregnancy:** Limited human data are available on the use of HYQVIA during pregnancy. The effects of antibodies to the Recombinant Human Hyaluronidase on the human embryo or fetal development are unknown. It is not known whether HYQVIA can cause fetal harm when administered to a pregnant woman or if it can affect reproductive capacity. HYQVIA should be given to a pregnant woman only if clearly needed.

Please click for Full Prescribing Information.

