Patient Enrollment Form | Rare Blood Disorders

Phone: 1-844-668-6732 Fax: 1-866-488-6576

Monday - Friday 8:00 AM to 8:00 PM ET



* Indi	cates a required field New start	☐ Reautho	orization 🗆] Restartir	ng treat	tment 🗆 1	Transitioning fr	om:			Page 1 of 3
SERVICES REQUESTED	Select a product: NovoSeven® RT Novoeight® Tretten® Rebinyn® Esperoct® Trial Program³ Patient has commercial prescription coverage, such as an HMO or PPO Patient has been newly prescribed a Novo Nordisk factor product for an indicated condition Patient is Novo Nordisk factor product-naïve JumpStart™ request. Shipping schedule to be confirmed with patient by NovoCare®. NovoCare® Savings Offer (if eligible) For complete copay terms and conditions, visit NovoFactorSavings.com Patients who have been prescribed one of the above products for an FDA-approved indication and who have commercial insurance may be eligible to receive a limited supply of free product from Trial and/or JumpStart™. Patient is not eligible if he/she participates in or seeks reimbursement or submits a claim for reimbursement to any federal or state health care program with prescription drug coverage, such as Medicaid, Medicare, Medigap, VA, DOD, TRICARE, or any similar federal or state health care program. Trial and JumpStart™ products are provided at no cost to the patient or the HCP, is not contingent on any product purchase, and the patient and HCP must not: (1) bill any third party for the free product, or (2) resell the free product. No purchase necessary.										
	Patient name:*								DOB (MM/DD/Y	YYY):*	
ш	Gender *:*										
PATIENT/INSURANCE INFORMATION	Home address (No P.O. box):						City:		State:	Zip:*	
₹ <u>o</u>	Shipping address (If different from Home Addres	is):					City:		State:	Zip:*	
ISL	Email:				Primary	/ phone:		Shi	p drug to: 🗖 Patien	t's home 🗖 Pr	escribing HCP
E S	Primary guardian/caregiver (required if patient	under 18 years old	d): *					Relationship to	patient:		
	Primary pharmacy insurance: (Please attach a	copy of the insu	rance card if avai	lable)				<u> </u>	Phone:		
	Rx # ID:	Rx Group #:		,		Rx PCN #:			Rx BIN #:		
4	Secondary pharmacy insurance: (Please attack		nauranaa aard if a	(aldalia)		TOX 1 CIV II.					
		1	risurarice card ii a	ivaliable)		D., DCN #			Phone:		
	Rx # ID:	Rx Group #:				Rx PCN #:			Rx BIN #:		
	Novo Nordisk and its partners recognize that p Please indicate the gender on file with the par			e or female. F	łowever,	many insurance	companies still rec	uire that one of	these two fields be u	sed for each of	their members.
D66 - Congenital hemophilia A (without inhibitors) D66 - Congenital hemophilia A (with inhibitors) D66 - Congenital hemophilia B (with inhibitors) D67 - Congenital hemophilia B (without inhibitors) D69.1 - Qualitative platelet defect (Glanzmann's Thrombasthenia) Other diagnosis: D68.2 - Other congenital factor deficiency (FVII) Patient Weight (kg): Patient Weight (kg):											
MEDICAL ASSESSMENT	IV Access: PIV/butterfly Implanted Port PICC Central Line Additional Information:										
4	Order Information: Complete prescription information below or submit a prescription with the strengths and assay limits. Quantity limits apply.										
		nformation belo	1	· · ·		e strengths and	assay limits. Qua	ntity limits appl	у.	_	l
NO N	Product name:		Dose:	Directions	:					Qty:	Refills:
E											
8								1			
PRESCRIPT	Highlights of prescribing information on following pages. For full prescribing information, see product sp						package insert.	Do you intend	I to buy and bill? Pharmacy Fax:	Yes 🗆 No	
4	Preferred pharmacy:					armacy Phone:	acy Phone:				
	Pharmacy address:				City	y:		State:		Zip:	
	Prescriber name:*							License	#:*		
	Practice name and office contact:						Preferred method of contact: ☐ Phone ☐ Fax ☐ Email				
	DEA #: Tax ID #:					NPI #.*					
	Phone:*	Fax:*				Email:*					
~8	Address:*				Cit	V:*		State:*		Zip:*	
PRESCRIBER AUTHORIZATION	Prescriber release:* By signing below, I hereby certify that: (a) I am a licensed practitioner, in good standing under applicable state law; (b) in my medical judgment, I have determined that the product(s) being prescribed which are detailed on the accompanying prescribing information (and listed only with those on label indications, appropriate treatment type, approved patient population, labeled dosing, and frequency) is to treat a diagnosis(es) consistent with indications dosing, and appropriate uses as described in the product's prescribing information; (c) the information I have provided on this enrollment form is, to the best of my knowledge, true, complete, and accurate in all respects; and (d) I have obtained the necessary authorization from the patient, or where appropriate the patient's parent, caregiver, and/or legal representative to use, disclose, share, and/or release the above-referenced information along with other protected health information (as defined in the Health Insurance Portability and Accountability Act of 1996 ("HIPAA")) for the sole purpose of providing patient assistance. Further, I appoint NovoCare®, on my behalf, to convey this prescription to the dispensing pharmacy. I will immediately notify Novo Nordisk Inc., its employees, or partners, including AssistRx, Inc. (collectively, "NovoCare®") if the above-named patient, or where appropriate the patient's parent, caregiver, and/or legal representative, revokes their consent to share their PHI with NovoCare®. I give you permission to contact me with any questions related to NovoCare®.										
	Prescriber signature (no signature stamps):*								Da	te:*	

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RARE BLOOD DISORDERS | HIGHLIGHTS OF PRESCRIBING INFORMATION

Novoeight®: Hemophilia A (congenital FVIII deficiency)

Treatment type	Patient Population/Bleed type	Dose/Target factor level	Frequency
Prophylaxis	Adult (≥ 12 yrs)	20-50 IU/kg	3 times weekly
	Adult (≥ 12 yrs)	20-40 IU/kg	Every other day
	Pediatric	25-60 IU/kg	3 times weekly
	Pediatric	25-50 IU/kg	Every other day
On-demand	Minor bleed	20-40 IU/dL	Every 12-24 hours
	Moderate bleed	30-60 IU/dL	Every 12-24 hours
	Major bleed	60-100 IU/dL	Every 8-24 hours
Perioperative	Minor surgery	30-60 IU/dL	Every 24 hours
	Major surgery	80-100 IU/dL	Every 8-24 hours

Esperoct®: Hemophilia A (congenital FVIII deficiency)

Treatment type	Patient Population/Bleed type	Dose	Frequency			
Prophylaxis	Adult (≥ 12 yrs)	50 IU/kg	Every 4 days*			
	Pediatric (< 12 years)	65 IU/kg	2 times weekly*			
On-demand	Adult: Minor bleed	40 IU/kg	1 dose should be sufficient			
	Adult: Moderate bleed	40 IU/kg	An additional dose may be administered after 24 hours			
	Adult: Major bleed	50 IU/kg	Additional dose(s) may be administered approximately every 24 hours			
	Pediatric: any bleed	65 IU/kg	Minor: 1 dose should be sufficient			
			Moderate: An additional dose may be administered after 24 hours			
			Major: Additional dose(s) may be administered approximately every 24 hours			
Perioperative	Adult: Minor or Major	50 IU/kg	Minor: Every 24 hours			
			Major: Every 24 hours for the first week, then approximately every 48 hours until wound healing has occurred			
	Pediatric: Minor or Major	65 IU/kg	Minor: Every 24 hours			
			Major: Every 24 hours for the first week, then approximately every 48 hours until wound healing has occurred			

^{*}Frequency can be adjusted based on bleeding episodes

Rebinyn®: Hemophilia B (congenital FIX deficiency); All doses/frequencies are for both adult and pediatric populations

Treatment type	Patient Population/Bleed type	Dose	Frequency			
Prophylaxis	ophylaxis N/A 40 IU/kg Onc		Once weekly*			
On-demand	Minor/Moderate bleed	40 IU/kg	1 dose should be sufficient, but additional doses of 40 IU/kg can be given			
	Major bleed	80 IU/kg	Additional doses of 40 IU/kg can be given			
Perioperative	Minor surgery	40 IU/kg	1 pre-op dose should be sufficient, additional doses of 40 IU/kg can be given			
	Major surgery	80 IU/kg	Pre-op dose; additional doses of 40 IU/kg can be given every 1-3 days within 1st week			
*Frequency can be ad	*Frequency can be adjusted based on bleeding episodes and physical activity					

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Tretten®: Congenital FXIII A-subunit deficiency; adult and pediatric populations

т	reatment type	Patient Population/Bleed type	Dose	Frequency
P	rophylaxis	N/A	35 IU/kg	Monthly*

^{*}Consider dose adjustment if adequate coverage is not achieved

NovoSeven® RT

Indication	Treatment type	Patient Population/Bleed type	Dose	Frequency			
Hemophilia A with inhibitors or hemophilia B with inhibitors	On-demand	Adult/pediatric: all other bleeds	90 mcg/kg	Every 2 hours until hemostasis is achieved, or until the treatment has been judged to be inadequate			
		Adult/pediatric: severe bleeds	90 mcg/kg	Every 2 hours until hemostasis and then post hemostatic every 3-6 hours			
	Perioperative	Adult/pediatric: minor surgery	90 mcg/kg	Immediately before surgery, every 2 hours during surgery, every 2 hours after surgery for 48 hours and then every 2-6 hours until healing occurs			
		Adult/pediatric: major surgery	90 mcg/kg	Immediately before surgery, every 2 hours during surgery, every 2 hours after surgery for 5 days and then every 4 hours or by continuous infusion (50 mcg/kg/hr) until healing occurs			
Congenital FVII deficiency	On-demand	Adult/pediatric	15-30 mcg/kg	Every 4-6 hours until hemostasis achieved			
	Perioperative	Adult/pediatric	15-30 mcg/kg	Immediately before surgery, every 4-6 hours during surgery and until healing occurs			
Glanzmann's thrombasthenia (with refractoriness to platelet transfusions)	On-demand	Adult/pediatric	90 mcg/kg	Every 2-6 hours until hemostasis achieved			
	Perioperative	Adult/pediatric	90 mcg/kg	Immediately before surgery, every 2 hours during surgery and every 2-6 hours post surgery			
Acquired hemophilia	On-demand	Adult	70-90 mcg/kg	Every 2-3 hours until hemostasis achieved			
	Perioperative	Adult	70-90 mcg/kg	Immediately before surgery, every 2-3 hours during surgery and until hemostasis achieved			