

Hemophilia Product Prior Authorization Form

Please complete this form in its entirety and provide relevant progress notes and/or bleeding diaries and **fax to 1-888-656-0841**. All lab results must be faxed in.

This request form pertains to the following products:

Feiba	Helixate FS	Alphanate	Hemlibra	Wilate
Feiba NF	Kogenate FS	Humate-P	BeneFIX	Idelvion
NovoSeven	Novoeight	AlphaNine SD	lxinity	Vonvendi
RT Hemofil M	Recombinate	Mononine	Rixubis	Afstyla
Koate-DVI	Xyntha	Bebulin	Alprolix	
Monoclate-P	Adynovate	Kovaltry	Coagadex	
Nuwiq	Eloctate	Profilnine	Corifact	
Advate	Obizur	Rebinyn	Tretten	

I. Demographic Information

Patient Information					
First Name	Last Name	Patient Gender			
Patient DOB	Patient Phone # Alternative Phone #				
Patient Address:					
City	State	Zip code			
Provider Information					
Prescriber Name	Contact Name	Contact Phone #			
NPI	Fax#				
Prescriber Address:					
City	State Zip code				

Rendering Provider (Dispensing Pharmacy) Information							
Pharmacy Name				NPI			NABP
Contact Name	Phone #				Fax #		
Insurance Information							
Policy Holder Name			ID# of Insurance Card				
Name of Insurance Company			Group #				
Primary Diagnosis							
 □ Congenital Hemophilia A (Congenital Factor VIII Deficiency) □ Acquired Hemophilia A (Aquired Factor VIII Deficiency) □ Hemophilia B (Congenital Factor IX Deficiency) □ von Willebrand Disease □ Congenital Factor XIII Deficiency □ Congenital Factor XIII A-subunit Deficiency □ Hereditary Factor X Deficiency □ Congenital Factor VII Deficiency □ Glanzmann's Thrombasthenia 							
ICD 10 Code							
Patient Inventory (Medication on Hand)							
Total Number of Doses on Hand	Total Units on Hand (IU)			Date Verified			
Clinical Information							
Name of Treating Facility							

Treatment status		Product Name				
☐ Treatment-naïve						
☐ Treatment-experien	nced					
Was the patient on a different factor product previously?						
☐ Yes						
□ No						
If yes, which product and reason for product switching:						
Member's Height Member's Weight			Severity of Disease			
			☐ Mild (6% to 25% factor level)			
			☐ Moderate (1% to 5% factor level)			
			☐ Severe (< 1% factor level)			
Dose (IU)	Number of Doses F	Requested	Total Dose Requested (IU)			
Dosing Instructions		Retrospec	Retrospective request?			
		☐ Yes				
		□ No				
Type of Use (Check all that applies)		Place of Administration:				
☐ Episodic		□ но	☐ Home infusion			
☐ Prophylaxis		☐ Outpatient Hemophilia Treatment Cente				
☐ Acute Bleeding Episode		(HTC)				
☐ Dental Procedure		☐ Outpatient Hospital				
Date of Procedure:		Provider's office				
☐ Surgical Prophylaxis		☐ Self-administration				
Date of Procedure:						
Number and Location of bleeds in the past 12 months:						
Does the patient have a diagnosis confirmed by blood coagulation testing?						
□ Yes						
□ No						

Please provide the following information regarding factor levels				
☐ Factor VIII for Hemophilia A				
☐ Factor IX for Hemophilia B				
☐ Factor X for Hereditary Factor X Deficiency				
☐ Factor XIII for Congenital Factor XIII or Factor XXIII A-subunit Deficiencies				
☐ VW Factor for von Willebrand Disease				
a. Baseline Factor Level				
b. Date of Factor Level				
c. Desired (Target) Factor Level				
Does the patient have inhibitors to factor products?				
☐ Yes ☐ No				
If so, are documentations of inhibitor tests attached? (e.g., Bethesda inhibitor assay)				
☐ Yes				
□ No				
Has the patient previously received Immune Tolerance Induction (ITI)?				
☐ Yes				
□ No				
If yes, date and duration of the trial and patient response:				
Did the patient experience at least two documented episodes of spontaneous bleeding into the joints?				
☐ Yes ☐ No				
For minimally treated patients (< 50 exposure days to factor products) previously receiving a different factor product, how often will inhibitor testing be performed?				

Was a pharmacokinetics (PK) test performed for this patient?					
□ Yes □ No					
If so, are PK testing resul	ts attached?				
☐ Yes ☐ No					
If patient has a diagnosis	of Glanzmaı	nn's Thrombast	henia, has the p	oatient tried	platelet transfusions?
□ Yes □ No					
If yes, date of the tria	I and patient	response:			
If the patient has a diagn	osis of von V	Villebrand Dise	ase (VWD), has	the patient	tried desmopressin?
☐ Yes ☐ No					
If no, is the patient contraindicated to desmopressin?					
□ Yes □ No					
If yes, what is the reason for contraindication:					
For acute bleeding episodes, please provide the following additional information:					
Location of Bleed	Type of Blee Minor Mode Major	rate	Start Date of Bleed:		End Date of Bleed:
Number of Doses Used		Dose (IU)		Total Amo	unt Used (IU)

To view current hemophilia policies and the Hemophilia Product Prior Authorization Form, please visit **FL.ExploreMyPlan.com/provider**.