## BlueCross BlueShield of Alabama

## PEEHIP ONLY Hemophilia Product Prior Authorization Form

Please complete this form in its entirety and provide relevant progress notes and/or bleeding diaries and **fax to 1-866-606-6021**. 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	Ixinity	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		

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

Treatment status		Product Name		
☐ Treatment-naïve				
☐ Treatment-experienced				
Was the patient on a differ	Was the patient on a different factor product previously?			
□ Yes				
□ No				
If yes, which product a	nd reason for produc	ct 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)	
			D Severe (*176 factor fever)	
Dose (IU)	Number of Doses F	Requested	Total Dose Requested (IU)	
Dosing Instructions	,	Retrospective request?		
		☐ Yes		
		□ No		
Type of Use (Check all that applies)		Place of Administration:		
☐ Episodic		☐ Home infusion		
☐ Prophylaxis		☐ Outpatient Hemophilia Treatment Center		
☐ Acute Bleeding Episode		(HTC		
☐ Dental Procedure		☐ Outpatient Hospital☐ Provider's office		
Date of Procedure:  ☐ Surgical Prophylaxis				
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				
<u> </u>				

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?
<u> </u>

Was a pharmacokinetics (PK) test performed for this patient?					
☐ Yes	□ Yes				
□ No	□ No				
If so, are PK testing resul	If so, are PK testing results attached?				
☐ Yes					
□ No					
If patient has a diagnosis	If patient has a diagnosis of Glanzmann's Thrombasthenia, has the patient tried platelet transfusions?				
☐ Yes					
□ No	□ No				
If yes, date of the tria	l and patient	response:			
If the patient has a diagr	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:					
Tor acate biccamig cpisc	acs, picasc	provide the for	iowing addition	iai iiiioiiiiat	1011.
Location of Bleed	ed Type of Bleed Si		Start Date of Bleed:		End Date of Bleed:
☐ Moderate					
	☐ Major	•			
Number of Doses Used		Dose (IU)		Total Amo	unt Used (IU)
				1	

To view current hemophilia policies and the Hemophilia Product Prior Authorization Form, please visit **AlabamaBlue.com/providers/policies**.