

PRIOR AUTHORIZATION REQUEST FORM **HEMOPHILIA AND BLOOD PRODUCTS**

For authorization, please answer each question and fax this form PLUS chart notes back to Real Rx at 385-425-4052.

Failure to submit clinical documentation to support this request will result in a dismissal of the request.

If you have prior authorization questions, please call for assistance 385-425-5094.						
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Disclaimer: Prior authorization request for	ms are subject to change in acco	rdance v	with Fede	ral and State notice requirements.		
Date:	Member Name:		ID#:			
DOB:	Gender:		Phys	sician:		
Office Phone:	Office Fax:		Offic	ice Contact:		
Height/Weight:			HCP	CS Code:		
Member must try formulary preferred dru	as before a request for a non-pr	eferred (drua mav	be considered. If treatment with		
preferred products has not been successfu		-				
reason for failure. Reasons for failure mu		-		-		
HEMOPHILIA A AGENTS						
\square Advate $^{ ext{@}}$ (antihemophilic factor (recomb	inant), □ Alphanate® (antihemo	philic fac	ctor (hum	an), □ Desmopressin (DDAVP),		
☐ Helixate FS® (antihemophilic factor (rec			-			
\square Humate-P $^{ ext{@}}$ (antihemophilic factor (hum		-				
factor (recombinant), 🗆 Kovaltry® (antiher			-			
\square Novoeight $^{ ext{@}}$ (antihemophilic factor (reco	-		-			
(antihemophilic factor (recombinant), ☐ F		_				
•	·	•				
VIIa (recombinant)-jncw) □ Wilate® (antihemophilic factor (human), □ Xyntha® (antihemophilic factor (recombinant) Long-Acting Products: □ Adynovate® (antihemophilic factor (recombinant), □ Afstyla® (antihemophilic factor (recombinant),						
☐ Eloctate™(antihemophilic factor (recom	• • • • • • • • • • • • • • • • • • • •	,	y.a (a	(,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,		
HEMOPHILIA B AGENTS						
☐ Alphanine SD® (coagulation Factor IX), ☐ Benefix® (coagulation factor IX (recombinant), ☐ Mononine® (coagulation Factor IX),						
□ Ixinity® (coagulation factor IX (recombinant), □ Profilnine® (factor IX complex), □ Rixubis® (coagulation factor IX						
(recombinant)), Sevenfact® (coagulation factor VIIa (recombinant)-jncw)						
Long-Acting Products: □ Alprolix™ (coagulation factor IX (recombinant))						
VON WILLEBRAND DISEASE (VWD) AGENTS						
\square Alphanate $^{ ext{@}}$ (antihemophilic factor (hum	ian), 🗆 Stimate® (Desmopressin	(DDAVP), 🗆 Hum	ate-P® (antihemophilic factor		
(human), 🗆 Wilate (coagulation factor VIII complex (human)						
Dosing/Frequency:						
If the request is for reauthorization, proceed to reauthorization section.						
Questions Yes No Comments/Notes						
1. Is this request for an expedited rev						

If the request is for reauthorization, proceed to reauthorization section.			
Questions	Yes	No	Comments/Notes
1. Is this request for an expedited review?			
By checking the "Yes" box to request an expedited review (24			
hours), you are certifying that applying the standard review			

	time frame (72 hours) may place the member's life, health, or ability to regain maximum function in serious jeopardy.			
_	, , ,			Black and the day of the state
۷.	Does the member have a diagnosis of moderate or severe			Please provide documentation
	hemophilia A, hemophilia B, or laboratory confirmed diagnosis			
_	of type 2B or type 3 Von Willebrand's disease?			Black of the day of the day
3.	For members with mild hemophilia A OR type 1, 2A, 2M, or 2N			Please provide documentation
	Von Willebrand Disease, has the member tried and failed, or			
	has a contraindication/intolerance, or a clinical reason for not			
_	using desmopressin (DDAVP)?			
4.	Is the request made by, or in consultation with, a			
	hematologist?			
5.	Will the request be made for one of the following?			Please provide documentation
	 Treatment and control of bleeding episodes 			
	Perioperative management of bleeding			
	Prevention of bleeding episodes			
	ADYNOVATE®, AFSTYLA®, ELOCTATE	™ OR A	ALPROLIX	
1.	Has the member tried and failed or had an			Please provide documentation
	intolerance/contraindication to a shorter acting recombinant			
	factor OR has the physician provided rationale for use of longer			
	acting recombinant factor?			
	NOVO-SEVEN RT®	_		
1.	Does the member have one of the following FDA-approved			Please provide documentation
	indications?			
	Hemophilia A or B with inhibitors			
	Acquired hemophilia			
	Congenital factor VII deficiency			
	Glanzmann thrombasthenia			
	HEMLIBRA®			
	Does the member have diagnosis of Hemophilia A?			
2.	Is the request for routine prophylaxis or reducing frequency of			
	bleeding episodes?			
3.	Will it be used in combination with Immune Tolerance			
	Induction (ITI)?			
4.	Does the member have at least 2 documented episodes of			Please provide documentation
	spontaneous bleeding into joints?			
5.	For members with Hemophilia A with inhibitors, are the high			Please provide documentation
	titer factor VII inhibitors ≥5 Bethesda units?			
_	ther factor vir initiations is bethesad anits.			
6.	For members with Hemophilia A without inhibitors, does the			Please provide documentation
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6.	For members with Hemophilia A without inhibitors, does the			Please provide documentation
6.	For members with Hemophilia A without inhibitors, does the member have one of the following:			Please provide documentation
6.	For members with Hemophilia A without inhibitors, does the member have one of the following: • Diagnosis of severe Hemophilia A AND documentation of			Please provide documentation
6.	For members with Hemophilia A without inhibitors, does the member have one of the following: • Diagnosis of severe Hemophilia A AND documentation of endogenous factor VIII levels <1%, OR			Please provide documentation
6.	 For members with Hemophilia A without inhibitors, does the member have one of the following: Diagnosis of severe Hemophilia A AND documentation of endogenous factor VIII levels <1%, OR Diagnosis of moderate Hemophilia A AND documentation 			Please provide documentation
6.	 For members with Hemophilia A without inhibitors, does the member have one of the following: Diagnosis of severe Hemophilia A AND documentation of endogenous factor VIII levels <1%, OR Diagnosis of moderate Hemophilia A AND documentation of endogenous factor levels of 1% to 5%, OR 			Please provide documentation
	 For members with Hemophilia A without inhibitors, does the member have one of the following: Diagnosis of severe Hemophilia A AND documentation of endogenous factor VIII levels <1%, OR Diagnosis of moderate Hemophilia A AND documentation of endogenous factor levels of 1% to 5%, OR Diagnosis of mild hemophilia A AND documentation of 			Please provide documentation Please provide documentation
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7.	 For members with Hemophilia A without inhibitors, does the member have one of the following: Diagnosis of severe Hemophilia A AND documentation of endogenous factor VIII levels <1%, OR Diagnosis of moderate Hemophilia A AND documentation of endogenous factor levels of 1% to 5%, OR Diagnosis of mild hemophilia A AND documentation of endogenous factor levels of ≥ 5% Has the member tried and failed, or have documented contraindications, to two prophylactic factor VIII replacement products? 			Please provide documentation
7.	 For members with Hemophilia A without inhibitors, does the member have one of the following: Diagnosis of severe Hemophilia A AND documentation of endogenous factor VIII levels <1%, OR Diagnosis of moderate Hemophilia A AND documentation of endogenous factor levels of 1% to 5%, OR Diagnosis of mild hemophilia A AND documentation of endogenous factor levels of ≥ 5% Has the member tried and failed, or have documented contraindications, to two prophylactic factor VIII replacement products? Is the member is currently on Hemlibra AND diagnosed with 			Please provide documentation

SEVENFACT®				
9.	Does the member have one of the following FDA-approved indications?			Please provide documentation
	Hemophilia A or B with inhibitors			
	REAUTHORIZATIO	N		
	Is the request for reauthorization of therapy?			
2.	Has the member provided the current number of on-hand doses since previous authorization?			Please provide documentation
3.	For patients using Hemlibra®, has therapy shown to be effective with evidence of a positive clinical response?			Please provide documentation
4.	Does documentation demonstrate medical necessity which may include, but is not limited to, documentation of bleeding episodes?			Please provide documentation
Ac	lditional information:			
	ysician Signature:			

** Failure to submit clinical documentation to support this request will result in a dismissal of the request.**

Policy PHARM-110

Origination Date: 11/12/2020 Reviewed/Revised Date: 10/26/2022 Next Review Date: 10/26/2023 Current Effective Date: 11/01/2022

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