ANTIHEMOPHILIA AGENTS PRIOR AUTHORIZATION FORM





(form effective 1/8/2024)

Fax to PerformRxSM at **1-888-981-5202**, or to speak to a representative call **1-866-610-2774**.

PRIOR AUTHORIZATION REQUEST INFORMATION						
☐ New request ☐ I	Renewal request	Total # pages:				
Name of office contact: Contact's phone number:						
PATIENT INFOR	MATION					
Patient name:			F	Patient ID #:	DOB:	
Street address:						
Apt #:	City/state/zip:			Phone:		
PRESCRIBER INFORMATION						
Prescriber name:				Specialty:		NPI:
Street address:						
Suite #:	City/state/zip:					
Phone:	Fax:					
CLINICAL INFOR	RMATION					
Product requested: ☐ He	emlibra	☐ Factor (name):		J-code:	Weight:	lbs/kg
Strength/vial size:				# of vials:	NDC#:	
Strength/vial size:				# of vials:	NDC#:	
Administration date: (to) Dispense date:						
DX code (required):			1	Diagnosis (submit documentation):		
Directions:			1	Total quantity requested:	Duration:	
PHARMACY INFORMATION (Prescriber to identify the pharmacy that is to dispense the medication): Deliver to: □ Patient's Home □ Physician's Office □ Patient's Preferred Pharmacy Name: NPI#: Pharmacy Phone #: □ Pharmacy Fax #: □ Leaken will does that the policy agrees with the pharmacy shoes for delivery of this medication.						
□ I acknowledge that the patient agrees with the pharmacy chosen for delivery of this medication.						
INITIAL REQUESTS (Complete the section(s) below applicable to the patient and this request and SUBMIT DOCUMENTATION for each item.) 1. For HEMLIBRA (emicizumab), one of the following:						
2. For a BYPASSING AGENT (e.g., FEIBA NF, NovoSeven): For routine prophylaxis: Has hemophilia A with inhibitors AND (check all that apply): Failed to achieve clinical goals with Hemlibra Has a medical reason why Hemlibra cannot be used Has been using the requested bypassing agent for routine prophylaxis within the past 90 days For use other than routine prophylaxis (e.g., episodic/on-demand treatment, intermittent/per				☐ Has hemophilia B with inhibitors ☐ Has acquired hemophilia ☐ Has congenital factor VII deficiency ☐ Has Glanzmann's thrombasthenia		
☐ Has hemophilia A with inhibitors						
3. For a non-preferred FACTOR VIII, FACTOR IX, or VWF: \[\begin{align*} \text{Has been using the requested product within the past 90 days AND has a medical reason to continue using the requested product \[\begin{align*} \text{Failed to achieve clinical goals with or has a contraindication or an intolerance to the preferred FVIII, FIX, or FVIII/VWF products with the same half-life (standard v. extended half-life), if applicable. Refer to https://papdl.com/preferred drug-list for a list of preferred and non-preferred drugs in this class. \[\begin{align*} Has a diagnosis for which no preferred Antihemophilia Agents are appropriate. Refer to https://papdl.com/preferred-drug-list for a list of preferred and non-preferred drugs in this class.						
RENEWAL REQUESTS						
4. Experienced a positive	ve clinical response sinc	e starting the requested medication	on: 🗆 Yes 🔝	□ No		
PLEASE FAX COMPLETED FORM WITH REQUIRED CLINICAL DOCUMENTATION						
Prescriber signature:					Date:	

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