Request for Patient Designated Plasma Protein and Related Products



INFORMATION TO BE PROVIDED BY REQUESTING HOSPITAL/PRESCRIBER

This form must be used for initial requests, renewals and changes. It is to only be used for products licensed in Canada. For unlicensed products, go to the Health Canada Special Access Program website. **Request** forms must be sent to **SAPPRPRequests@blood.ca** or to your **local Canadian Blood Services Distribution Site** at least 2 weeks before product is required (review may take longer if requesting access outside of listed criteria (i.e., exceptional access)). If approved, a contract number will be assigned which must be referenced on subsequent orders using the Order Form for Plasma Protein and Related Products Requiring Contracts or through the Online Ordering Portal.

Section I: Requesting Hospital Details and Patient Information (complete for all request types) Unless this is an emergency request, by completing and submitting this form, you agree that your patient has been provided the Privacy Notice for Patient Designated Plasma Protein and Related Products.						
Hospital Information Canadian Blood Services customer # if known: 500364						
Request Date (YYYY-MM-DD):						
Requesting Hospital Name:						
Ship to Hospital/Location:	Queen Elizabeth Hospital, Charlottetown, PE					
Hospital Contact 1*:	Dr. Jennifer Fesser					
_{Email:} jnfesser@ihis.org		Phone #: (902) 894-2535	Fax #: (902) 894-2415			
Hospital Contact 2*:	Ami MacQuarrie					
Email: anmacquarrie@ihis.org		Phone #: (902) 894-2329	Fax #: (902) 894-2415			
Ordering Prescriber:						
Email:		Phone #:	Fax #:			
*Contract Notification will go to th	e Hospital Contact(s) Email/F	ax#.				
Patient Information						
Last Name:		First Name:				
Date of Birth (YYYY-MM-DD):		Sex (M/F):				
Height (cm):		Weight (kg):				
Provincial/Territorial Health Card Number:						
Province/Territory of Residence:						
Section II: Request Type						
□ New Patient (pr	oceed to Section III)	Renewal (includes changes)	□ Further Information			
Canadian Blood Services Patient # Canadian Blood Services Contract #						
Section III: Product and Criteria						
Diagnosis:						
Panhematin (hemin)						
Amelioration of recurrent attacks of acute intermittent porphyria temporally related to the menstrual cycle in susceptible women, after initial carbohydrate therapy is known or suspected to be inadequate						
OR Grout viewer For urgent use						
Confidential	I	Page 1 of 3	F800135 (Revision 1)			

TEM-00003 Rev 2

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	Hemlibra (emicizumab)							
Prescribed by a hematologist with experience in the		Supporting Information ([#] required values)						
diagnosis and management of hemophilia A		FVIII inhibitor level (BU/mL) #						
AND one of the following:				□ %				
 Congenital hemophilia A with inhibitors to factor VIII (> 0.6 Bethesda Units/mL) confirmed on more than one occasion by an appropriate assay 			Intrinsic FVIII level [#]		🗆 IU/mL			
			Annual bleeding rate#					
	Severe congenital hemoph VIII level < 1%) without inh		Number of target joints					
	candidates for routine prop prevent bleeding or reduce bleeding episodes	ohylaxis to	Number of hospital/clini treatment of bleeds in th					
	Other** (provide rationale Information or include an a							
	Glassia (alpha-1 protein	ase inhibitor)						
	acia may be requested for	a adult nation to that	Supporting Information	on (# required val	ues)			
	issia may be requested fo et <u>ALL</u> of the following cr		Baseline serum A1-PI	evel [#]	□ µmol/L			
	Respirologist has confirm	ed the diagnosis of			□ mg/dL			
severe alpha-1 proteinase deficiency and clinical evi and indicated that patient			FEV1 (%)#					
		would benefit from						
	treatment with A1-PI prod A1-PI deficiency, defined		If baseline serum A1-PI level is unavailable, please clarify below:					
	<11 μ mol/L or < 57 mg/dL		□ Already on treatment with A1-PI product and no record of baseline level					
	treatment			□ Other (explain):				
	Clinical evidence of obstru	· · · · · ·						
	Nonsmoker for at least 6							
	Has not received a lung tr	ansplant						
Other Product**:								
**If patient does not meet listing criteria or product is identified as "Other", an exceptional access review will be required. Please note that additional information may be requested, and the timeline for review may increase.								
	Current Therapy or D N/	Ά						
Product Name Dose		Route of	Frequency of	Indication (e.g., prophylaxis, on				
			Administration	Administration	demand)			
New Requested Therapy or Same as Current Therapy								
Pro	duct Name	Dose	Route of Administration	Frequency of Administration	Indication (e.g., prophylaxis, on demand)			



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Other Supporting Information	(including rationale for chang	ge or initiation of therapy):

Section IV: Total Contract Quantities in Vials (refer to order form for product and available sizes)

Contracts will be created up to a maximum of **12 months**, A renewal request will be required every 12 months

Vial Size	Total Contract Quantity	Pick Up Quantity	Frequency of Pic (e.g., every 3 mo		Duration of Contract (max 12 months)	
Date of next product order (please comment if less than 1 week):		Comments (please inc	lude when next dose	e is due f	or STAT requests):	
Expiry date of approved contra (optional to fill out for records following CBS notification):	act					
Section V: Urgent Medical Review and SAP Information (CBS Use Only)						
The on-call medical officer can be contacted after hours to review urgent requests for patients that meet listing criteria . Exceptional access reviews cannot be completed by the on-call medical officer and should be sent to the PPRP Formulary team for regular review. Please forward the request form with all documentation of medical review to <u>SAPPRPRequests@blood.ca</u> .						
Decision of urgent medical officer review: Approve 30-day supply (specify amount below) Deny						
Comments:						
If medical review was obtained verbally, indicate results of review in comment section above. Include: as per (physician name), initial and date (e.g., as per Dr. Jane Doe, LA 2019-07-27)						
SAP Patient #:	SAP Contract #:	Completed/Enter	ered by:	Date:		