Product(s)	Administration**	Storage/Stability				
Humate-P	Humate P given over 3-20 min (Max 4mL/min)	Prior to reconstitution: 2-8°C or room temp (≤25C), do not freeze.	Reconstitution: Bring to room temperature and reconstitute with provided diluent. Swirl, do not shake or filter.			
FEIBA NH	FEIBA max rate of 2 units/kg/min	`	After reconstitution: Room temp (≤25°C), use within 3hr			
BeneFix	Benefix given at a rate of 2-4mL/min over several minutes	Prior to reconstitution: 2- 8°C, do not freeze. May be stored at room temp (≤25C) for up to 6 months				
Kogenate	Over 1 to 15 minutes IVP unless ordered as Continuous Infusion	Prior to reconstitution: 2-8°C, do not freeze, Helixate FS may be stored at room temp ≤ 25C, up to 12 months.	After reconstitution: Room temp (≤25°C), use within 3hr Continuous infusion of Helixate FS or Recombinate			
		200, up to 12 months.	10 IU/mL dilution in NS. Dose based on replacement requirements. Infuse within 12hrs			
NovoSeven RT	given as IVP over 2-5 minutes (can also be ordered as Continuous Infusion)	Prior to reconstitution: <25°C	After reconstitution: Room temp (≤25°C) of refrigerated IVP: Use within 3hr; do not freeze; do not store in syringes CI: Do not dilute. Infuse within 12 hours (stability questionable beyond 12-24 hours)			

*Refer to JDH/UConn IV Guidelines for further information. Do not use an inline filter. Flush with Normal Saline, Store all products in original packaging

Humate-P Dosing for VWD Hemorrhage DOSED IN VWF:RCo ONLY								
VWD Type	Severity	Dosage (IU VWF:RCo/kg)						
Type 1 – Mild	Minor	Use DDAVP IV 0.3mcg/kg by slow infusion						
(VWF:RCo >30%)	Minor w/o DDAVP or Major	Load 40-60 IU/kg then 40-50IU/kg q8-12hr x3 days ¹ , then 40-50IU/kg qday ²						
Type 1 – Moderate/	Minor	40-50 IU/kg (x1-2 doses)						
Severe (VWF:RCo <30%)	Major	Load 50-75 IU/kg then 40-60 IU/kg q8-12hr x3 days ¹ , then 40-60 IU/kg qday ²						
Type 2 and 3 (all)	Minor	40-50 IU/kg (x1-2 doses)						
	Major	Load 60-80 IU/kg then 40-60 IU/kg q8-12hr x3 days ¹ , then 40-60 IU/kg qday ²						

BILLING INFO

- EVERY bolus and IV factor dispensed including Kcentra <u>must</u> be entered in the Cubixx log clearly and neatly and initialed clearly by the dispensing Pharmacist.
- Entering pharmacists must physically enter in the number of labels needed to ensure doses are sent, using ... labels for inpatients.
- Billing personnel will verify all factor billing with the log, and MAK or the MAR.
- At present, Factor Products are done in Dose Edge
- Bolus doses in excess of 3 vials should be drawn up and labeled by Pharmacy once need of dose is verified.

¹To maintain VWF:RCo trough level >50% ²For up to 7 days

References: LexiComp, NovoSeven RT (NovoNordisk) package insert http://www.novo-pi.com/novosevenrt.pdf

Recombinate (Baxter) package insert http://factorviia.com/pi.pdf

BeneFix (Wyeth) package insert http://www.pfizerpro.com/resources/minisites/hemophilia/docs/BeneFIX-Pl.pdf FEIBA (Baxter) package insert http://www.feiba.com/us/forms/feiba_nf_pl.pdf

Humate-P (CSLBehring) package insert http://www.humate-p.com/Professional/Prescribing-Information.aspx

Thigpen and Limdi, Reversal of Oral Anticoagulation. *Pharmacotherapy*, vol 33, issue 11, pp. 1199-1213, *November*, 2013

KCentra package insert, http://www.kcentra.com/Kcentra-resources.aspx



UCONN JOHN DEMPSEY HOSPITAL

Department of Pharmacy, November 2018 Revised by Ruth Kalish, RPh UConn Health
Department of Pharmacy

Procoagulant (Factor) Agent



Helpful Hints

For Factors VIII and IX, IU/kg dose is a general guideline, as dosage should be based on desired/expected goal factor concentration.

For Factor VIII, 1 IU/kg can be expected to raise the factor levels by 2 percentage points. For Factor IX, 1 IU/kg can be expected to raise factor levels by 0.75 percentage points

Humate P is the only product available for patients with von Willebrand Disease (VWD). Dose is based on VWF:RCo units only for VWD. VWF units varies in ratio to Factor VIII units with an average of 2.4:1.

* Per Pharmacy & Therapeutics committee, Pharmacy can adjust dose within +/- 10% based on current dosage vial sizes in pharmacy to prevent waste. Rounding up to the nearest vial size is preferred.

ASD Healthcare has provided a Cubixx refrigerator to store the consignment product in. If there is a mechanical issue with the refrigerator contact card is taped to the refrigerator.

The pharmacy will be billed for the product when it has been removed from the Cubixx unit for more than 180 minutes

When the inventory falls below the designated par levels, the Cubixx unit will automatically generate an order. The turnaround time for system generated orders is 2 days.

Do not deface any packaging for all factor agents (e.g. write on or label the box, remove the filter, remove the package insert) or JDH Pharmacy will be required to pay for the product since they are on consignment. Store products in the original package to protect from light.

Pharmacists ONLY: If an order needs to be placed urgently off hours, call 1-800-746-6273. The operator will have an associate call back. Identify yourself as John Dempsey Hospital, acct. # 206267.



Department of Pharmacy

Product	Half- Life (h)	How Supplied	Indications: FDA-Approv (<i>Unapproved</i>)	/ed	Type of Bleed	Dose		Interval	Duration	Goal Factor Concentration
BeneFIX *	44.00	250, 500,	Hemophilia B		Minor	Up to 40 IU/kg		12-24hr	Until resolved	20-30%
Factor IX (recombinant)	11-36 it)	1000, and 2000 IU vials w/ 5ml diluent	Number of factor IX IU re patient weight (kg) x desi		Moderate	Up to	65 IU/kg	12-24hr	up to 7 days	25-50%
Pfizer 1-800-505-4426	Final conc: 50, 100, 200,	level increase (as % norr x 1.3 (as units/kg or units	mal or IU/dL)	Major	·	130 IU/kg	12-24hr	7-10 days or until resolved	50-100%	
		400IU/mL	, ·	,	Surgery	Up to 130 IU/kg pre- op, then less		12-24hr PRN to Keep ≤50%	7-14 day or until resolved	Pre-op: 100% Post-op ≤ 50%
FEIBA NF* Activated Pro-	ated Pro- 4-7	approx. 500IU or 1000 IU/	Hemophilia A/B with inl	h factor VIII s >5 Bethes- of life threat-	Joint	50-10 units/	0 Feiba kg	12hr DIC or throm-	Consult special- ist if ineffective	NA
thrombin Com- plex Concen- trate (plasma-	Dura- tion =	20ml and approx. 2500 IU/50ml vials	(acquired hemophilia with or factor IX inhibitor titers da units (BU) Treatment of ening bleeds associated v		Mucous mem- brane	50-100 Feiba units/kg		bosis may occur if 200unit max dose exceeded	or >3 doses in 24hr 50 unit/kg dose	
derived) Baxter	8-12h	10/00mm viaio			Soft tissue					
Healthcare (800) 422-9837			dabigatran)		Severe (CNS)		eiba units/kg of 200 units/kg	12hr	- can be q6h	
NovoSeven RT	1.7-	1mg, 2mg, 5mg, 8mg	Hemophilia A or B with		Hemophilia A or B	90 m	cg/kg	q2hr initially. Cont. Post-op	Until hemosta- sis achieved	NA
Factor VIIa (recombinant)	3.1	vials as 1mg/mL	factor VIII or factor IX, acquired hemophilia, or congenital factor VII deficiency (warfarin-related intracerebral hemorrhage; treatment of refractory bleeding after cardiac surgery in nonhemophiliac patients) CI = 1 mg/ml FVIIa diluent		Acquired hemoph.	70-90 mcg/kg		until healed Q2h up to 5 days, Consult		Treat until bleeding
Novo Nordisk (800) 727-6500					Congenital factor VII deficiency	IV Pu 15-30	sh: Imcg/kg	IVP q4-6hr until hemostasis,	doses or inade- quate response	stopped, sur- gery healed or judged ineffective
						Continued 12	nuous: undilut- -50mcg/kg/hr	then CI based on levels		
Helixate FS *	11-15	250, 500,	Hemophilia A (bleeding/s		Minor	10-20	IU/kg	12-24hr	Until resolved	20-40%
Factor VIII (recombinant)		1000, 2000 units/ 2.5ml	prophylaxis of joint bleeding and to reduce risk of joint damage in children with hemophilia A with no preexisting joint damage Wt (kg) x desired % increase/2 = dose Continuous infusion = 10 units/ml NS JDH does not carry Recombinate*, but dosing guidelines are similar		Moderate/minor surg	15-30 IU/kg		12-24hr	3 days or until- resolved	30-60%
CSL Behring (800) 504-5434		Final conc: 100IU/mL, 200IU/mL,			Major	40-50 IU/kg f/b 20-25IU/kg		8-12hr	3-7 days or until bleed resolved	80-100%
		400IU/mL, 800IU/ml			Major Surgery	50 IU/kg load Cl dose based on levels		6-24hr	10-14 days or until healed	Pre-op: 100% Monitor levels
Humate-P * Factor VIII 8-28 (plasma-derived) (HA)		250/600IU/5ml 500/1200IU/10	Hemophilia A (classical hemophilia) (this table); Not preferred agent.		Minor	15 IU FVIII/kg f/b 7.5 IU FVIII/kg		12-24hr	Until resolved	30%
CSL Behring (800) 504-5434	SL Behring	ml 1000/2400IU/15 ml	See Reverse for Severe VWD, including mild or moderate disease where use of desmopressin is known		Moderate	25 IU FVIII/kg f/b 15 IU FVIII/kg		8-24hr	3-7 days or until resolved	30-50%
See Guidelines for Von Wil- lebrand's Ds on reverse	(VWD)	VIII/VWF:RCo Ratio Approx 2.4:1	or suspected to be inade Not indicated for the prop spontaneous bleeding ep	quate <u>Note</u> : ohylaxis of	Surgery or Major	40-50 IU FVIII/kg f/b 20-25 IU FVIII/kg		8-24hr	7-14days or until resolved	Pre-op + initial Post-op: 80- 100% x7days f/b 30-50% x7d
KCentra* Indication		ndication			Dose (in Factor IX con- tent) Administration		Administration	Caution	Storage	Monitoring
Prothrombin complex concentrate, heat treated, (Factors II, VII, IX, X)		duced coagulation factor deficiency in patient with acute major bleed or urgent		4 - 6 > 6	25 units/kg max 2500 t 35 units/kg max 3500 t 50 units/kg max 5000 t Do not exceed max do	units units	0.12ml/kg/min or approx. 3 units/kg/min. NTE 8.4ml/min	Do not let blood enter syringe to avoid clot for- mation. Monitor for clotting	Refrigerate. Use within 4 hrs of reconstitution if refrigerated	INR declines within minutes, duration 6-8hrs May affect aPTT