SENTARA HEALTH PLANS

MEDICAL PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

<u>Directions:</u> The prescribing physician must sign and clearly print name (preprinted stamps not valid) on this request. All other information may be filled in by office staff; <u>fax to 1-844-668-1550</u>. No additional phone calls will be necessary if all information (including phone and fax #s) on this form is correct. <u>If information provided is not complete, correct, or legible, authorization can be delayed</u>.

<u>For Medicare Members:</u> Medicare Coverage for outpatient (Part B) drugs is outlined in the Medicare Benefit Policy Manual (Pub. 100-2), Chapter 15, §50 Drugs and Biologicals. In addition, National Coverage Determination (NCD) and Local Coverage Determinations (LCDs) may exist and compliance with these policies is required where applicable. They can be found at: https://www.cms.gov/medicare-coverage-database/overview-and-quick-search.aspx. Additional indications may be covered at the discretion of the health plan.

Non-Preferred Parenteral Iron Products

<u>Drug Requested</u>: (select ONE of drugs below) (Medical)

PREFERRED								
		ı	No prior authoriza	tion required				
	Feraheme ®	□ Fe	Ferrlecit [®] (sodium ☐ INFeD [®] (iron		on	□ Venofer [®] (iron		
	(ferumoxytol) (For	ferric gluconate		dextran) J1750		sucrose) J1756		
	ESRD on Dialysis)	complex) J2916						
	Q0139							
	NON-PREFERRED							
		Pr	ior authorization requ	ired as noted below				
□ Feraheme [®] (ferumoxytol)		ytol)	☐ Injectafer® (ferric ☐		□ Mo	noferric® (ferric		
(Non-ESRD) Q0138		carboxymalto	ltose) J1439 deri		somaltose) J1437			
			1					
MI	EMBER & PRESCR	IBER I	INFORMATIO	N: Authorization	may be de	layed if incomplete.		
Men	nber Name:							
Member Sentara #:		Date of Birth:						
Prescriber Name:								
Prescriber Signature:		Date:						
Offic	ce Contact Name:							
Phone Number:			Fax Number:					
NPI #:								

(Continued on next page)

DRUG INFORMATION: Authorization may be delayed if incomplete.					
Drug	g Form/Strength:				
Dosin	ing Schedule:]	Length of Therapy:			
Diagnosis:		ICD Code, if applicable:			
Weig	ght (if applicable):	Date weight obtained:			
	Standard Review. In checking this box, the timeframe doe or the member's ability to regain maximum function and w				
supp	LINICAL CRITERIA: Check below all that apply. A port each line checked, all documentation, including lab revided or request may be denied.				
Len	ngth of Authorization: 2 months				
Dia	agnosis – Select <u>ONE</u> of the following diagnoses	below:			
	Diagnosis: Iron-deficiency anemia				
	Provider has submitted the following labs collected wit Serum ferritin (iron) <u>AND</u> total iron binding capaci Transferrin saturation (TSAT%) *Note: TSAT% =	ty (TIBC)			
	☐ Lab documentation show member's TSAT < 20%				
	 □ Provider has submitted documentation to confirm mem preferred parenteral iron preparations □ Feraheme[®] (ferumoxide) for ESRD on Dialysis □ Ferrlecit[®] (sodium ferric gluconate complex) □ INFeD[®] (iron dextran) □ Venofer[®] (iron sucrose) 	ber has tried and failed <u>ONE</u> of the following			
	Diagnosis: Moderate-to-severe restless leg sy	ndrome (RLS)			
	Member is 18 years of age and older				
_	☐ Serum ferritin (iron) <u>AND</u> total iron binding capace ☐ Transferrin saturation (TSAT%) *Note: TSAT% =	eity (TIBC) = (Serum iron/TIBC) x 100%			
	- Lao documentation shows member 5 15A1 \20/0 alter	and of an oral non supplement			

PA Non-Preferred Parenteral Iron Products (Medical)(CORE) (Continued from previous page

	Member has tried and had an unsatisfactory response, intolerance or contraindication to oral iron administration				
	rovider has submitted documentation to confirm member has tried and failed <u>ONE</u> of the following referred parenteral iron preparations				
	☐ Feraheme [®] (ferumoxytol) for ESRD on Dialysis				
	Ferrlecit® (sodium ferric gluconate complex)				
	□ INFeD® (iron dextran)				
	□ Venofer® (iron sucrose)				
□ I	Diagnosis: Management of cancer and chemotherapy-induced anemia				
	Provider has submitted the following labs collected within the last 30 days:				
	☐ Serum ferritin (iron) AND total iron binding capacity (TIBC)				
	□ Transferrin saturation (TSAT%) *Note: TSAT% = (Serum iron/TIBC) x 100%				
	Provider has submitted documentation to confirm member has tried and failed ONE of the following preferred parenteral iron preparations				
	☐ Feraheme® (ferumoxytol) for ESRD on Dialysis				
	☐ Ferrlecit® (sodium ferric gluconate complex)				
	□ INFeD® (iron dextran)				
	□ Venofer® (iron sucrose)				
	Member has functional iron deficiency and must meet ONE of the following:				
	\square Member has a TSAT < 50% with the goal of avoiding allogenic transfusion				
	☐ Member has a TSAT < 50% and requested medication will be used in combination with erythropoises-stimulating agents (ESAs)				

(Continued on next page)

PA Non-Preferred Parenteral Iron Products (Medical)(CORE) (Continued from previous page

Med	ication being provided by: Please check applicable box below.
□ L	ocation/site of drug administration:
N	PI or DEA # of administering location:
	<u>OR</u>
\Box S_1	pecialty Pharmacy
standar urgent i	gent reviews: Practitioner should call Sentara Health Pre-Authorization Department if they believe a d review would subject the member to adverse health consequences. Sentara Health's definition of is a lack of treatment that could seriously jeopardize the life or health of the member or the member's to regain maximum function.
**	Use of samples to initiate therapy does not meet step edit/ preauthorization criteria.**
	vious therapies will be verified through pharmacy paid claims or submitted chart notes.*