

## **MEDICARE FORM**

## Feraheme® (ferumoxytol) and Injectafer® (ferric carboxymaltose) Medication Precertification Request

Page 1 of 2

(All fields must be completed and legible for precertification review.)

For Illinois MMP: FAX: 1-855-320-8445 PHONE: 1-866-600-2139

For other lines of business:

Please use other form.

Note: Feraheme, Injectafer, and Monoferric are non-preferred. The preferred products are Ferrlecit (sodium ferric gluconate), Infed,

and Venofer.

	Start of treatment: Start date _			•			
<del></del>	Continuation of therapy, Date o	of last treatment	1 1				
Precertification Requ	ested By:		Phone:		Fax:		
A. PATIENT INFORMA	ATION						
First Name:		Last Name:			DOB:		
Address:			City:		State:	ZIP:	
Home Phone:	Work Phone:		Cell Phone:		Email:		
Patient Current Weight:	kgs	Patient Height:	inches orcm	s Allergie	es:		
B. INSURANCE INFOR	RMATION						
Aetna Member ID #: Does patient have other			her coverage?	er coverage?			
			yes, provide ID#: Carrier Name:				
Insured:		Insured:					
Medicare: ☐ Yes ☐ No If yes, provide ID #: Medicaid: ☐ Yes ☐ No If yes, provide ID #:							
C. PRESCRIBER INFORMATION							
First Name:		Last Name:		(Check One	e):	D.O. 🗌 N.P. 🗌 P.A.	
Address:		-	City:		State:	ZIP:	
Phone:	Fax:	St Lic #:	NPI #:	DEA #:	1	UPIN:	
Provider Email:		Office Contact Name	);		Phone:	1	
Specialty (Check one):	☐ Hematologist ☐ Interna	I Medicine	r:		l .		
D. DISPENSING PROV	VIDER/ADMINISTRATION INFO	DRMATION					
Place of Administration			Dispensing Provid	er/Pharmac	v: Patient Sele	ected choice	
☐ Self-administered	☐ Physician's Office		☐ Physician's Offic	-	_ ☐ Retail Pharn		
Outpatient Infusion C	-		_ ☐ Specialty Pharm		_ ☐ Other	,	
Center Name:					<u> </u>		
☐ Home Infusion Cente			Name:				
Agency Name:			Address:				
1	s) (CPT):		_ City:				
Address:	State:	7ID:	_ Phone:				
	State Fax:		· · · · ·				
	PIN:		NPI:				
NPI:			_				
E. PRODUCT INFORM	IATION						
Request is for: Feraheme Injectafer Dose: Frequency:							
F. DIAGNOSIS INFORMATION - Please indicate primary ICD code and specify any other where applicable.							
Primary ICD Code:		_ Secondary ICD Co	ode :	Other I	CD Code:	_	
G. CLINICAL INFORM	ATION - Required clinical inform	nation must be complet	ed in its <u>entirety</u> for all pre	ecertification	requests.		
For All Requests (clini	ical documentation required for	or all requests):			•		
Note: Feraheme, Injectafer, and Monoferric are non-preferred. The preferred products are Ferrlecit (sodium ferric gluconate), Infed, and Venofer.							
☐ Yes ☐ No Has the patient had prior therapy with Feraheme (ferumoxytol injection) within the last 365 days?							
☐ Yes ☐ No Has the patient had prior therapy with Injectafer (ferric carboxymaltose injection) within the last 365 days?							
Yes No Has the patient had prior therapy with Monoferric (ferric derisomaltose injection) within the last 365 days?							
Yes No Has the patient had a trial and failure, intolerance, or contraindication to any of the following? (select all that apply)							
☐ Ferrlecit (sodium ferric gluconate) ☐ Infed (iron dextran) ☐ Venofer (iron sucrose)							
Please explain if there are any other medical reason(s) that the patient cannot use any of the following preferred products when indicated for the patient's diagnosis (select all that apply).							
,	appiy). ecit (sodium ferric gluconate)	Infed (iron devtran)	Venofer (iron sucrose)				
	Con (300ium femo giuconate)	imou (iioii uextiali)	volidiei (iidii sudiuse)				



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB				
G. CLINICAL INFORMATION (conf	 <i>tinued)</i>	nation must be completed in its <u>entiret</u>	v for all precertification requests.				
Please indicate the patient's serum ferritin level:							
Please indicate the patient's transferrin saturation (TSAT) level:							
☐ Yes ☐ No Was the serum ferritin and/or transferrin saturation level drawn within the last 30 days?							
☐ Yes ☐ No Is this a request for continuation of therapy?							
Yes No Does the patient have a contraindication, intolerance or ineffective response to Ferrlecit, Infed, or Venofer?							
For chronic kidney disease indications only:							
Yes No Does the patient have iron deficiency anemia associated with chronic kidney disease?							
Yes No Is the patient non-dialysis dependent (NDD) or undergoing peritoneal dialysis?							
Please explain: The patient is non-dialysis dependent (NDD) The patient is undergoing peritoneal dialysis							
For all other non- chronic kidney disease indications:							
☐ The patient is unable to tolerate oral iron compounds ☐ The patient is losing iron (blood) at a rate that is too rapid for oral intake to compensate for the loss							
The patient is losing from (blood) at a rate that is too rapid for oral intake to compensate for the loss  The patient is losing from (blood) at a rate that is too rapid for oral intake to compensate for the loss  The patient is losing from (blood) at a rate that is too rapid for oral intake to compensate for the loss  The patient is losing from (blood) at a rate that is too rapid for oral intake to compensate for the loss							
aggravated by oral iron therapy							
☐ The patient is unable to maintain iron balance on treatment with hemodialysis							
☐ The patient is donating large amounts of blood for autologous programs							
The patient has failed to heed instructions for oral iron supplementation or are incapable of accepting or following them							
The patient has heart failure and iron deficiency with or without anemia							
☐ The patient has iron deficiency and chemotherapy-induced anemia☐ The patient has iron deficiency anemia due to heavy uterine bleeding							
☐ The patient has iron deficiency following gastric bypass surgery and/or subtotal gastric resection and who exhibited decreased absorption							
of oral iron							
H. ACKNOWLEDGEMENT							
Request Completed By (Signatur	e Required):		Date: //				
Any person who knowingly files a request for authorization of coverage of a medical procedure or service with the intent to injure, defraud or deceive							
	any insurance company by providing materially false information or conceals material information for the purpose of misleading, commits a fraudulent insurance act, which is a crime and subjects such person to criminal and civil penalties.						

The plan may request additional information or clarification, if needed, to evaluate requests.