

MEDICARE FORM

Erythropoiesis Stimulating Agents Injectable Medication Precertification Request

Page 1 of 3

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

For Michigan MMP: FAX: 1-844-241-2495 PHONE: 1-855-676-5772

For other lines of business:

Please use other form

Note: Procrit and Epogen are nonpreferred. The preferred products are Aranesp, Mircera and Retacrit.

Please indicate: St		date <u>/ /</u> Date of last treatment	1 1				
		Date of last treatment			_		
Precertification Reques	=		Phone:		Fax:		
A. PATIENT INFORMA	IION	1			000		
First Name:		Last Name:	lo:		DOB:	1715	
Address:	I		City:		State:	ZIP:	
Home Phone:	ı	Phone:	Cell Phone:		Email:		
		Height: inches	or cms	Allergies:			
B. INSURANCE INFOR	MATION						
Aetna Member ID #:		Does patient have	Does patient have other coverage? ☐ Yes ☐ No				
Group #:			If yes, provide ID#: Carrier Name:				
Insured:		Insured:					
C. PRESCRIBER INFO	RMATION						
First Name:		Last Name:	1	Check One: L		O. N.P. P.A.	
Address:			City:	T==	State:	ZIP:	
Phone:	Fax:	St Lic #:	NPI #:	DEA #:		UPIN:	
Office Contact Name:				Phone:			
D. DISPENSING PROV	DER/ADMINISTRATION	ON INFORMATION	Dispensing Provide				
Place of Administration: Self-administered Physician's Office Home Outpatient Infusion Center Phone: Center Name: Home Infusion Center Phone: Agency Name: Administration code(s) (CPT): Address:			☐ Outpatient Dialy ☐ Retail Pharmacy ☐ Mail Order Name: Address: Phone: TIN:	rsis Center	Specialty Pharm Other:Fax:	acy	
E. PRODUCT INFORMA	ATION						
Request is for: Aranesp (darbepoetin alfa) Epogen (epoetin alfa) Mircera (methoxy polyethylene glycol/epoetin beta) Procrit (epoetin alfa) Retacrit (epoetin alfa-epbx) Dose/Frequency: HCPCS Code: Failure to provide dose & frequency may delay request)							
		te primary ICD code and spe					
		Secondary ICD Code:al information must be compl					
For All Requests: (Clinical documentation required for all requests) Yes No Will Aranesp (darbepoetin alfa), Procrit (epoetin alfa), Epogen (epoetin alfa), Mircera (methoxy polyethylene glycol/epoetin beta), or Retacrit (epoetin alfa-epbx) be used concomitantly? Yes No Is the patient currently taking iron supplements? Hemoglobin (Hgb) result?mg/dL Date of test/_/ For Initial Requests: Note: Procrit and Epogen are non-preferred. The preferred products are Aranesp, Mircera and Retacrit. Preferred products may vary based on indication. Yes No Has the patient had prior therapy with the requested product within the last 365 days? Yes No Has the patient had a trial, intolerance, or contraindication to any of the following? (select all that apply) Aranesp (darbepoetin alfa) Mircera (methoxy polyethylene glycol-epoetin beta) Retacrit (epoetin alfa-epbx) 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)							
☐ Ara	nesp (darbepoetin alfa)	☐ Mircera (methoxy polyethy	lene glycol-epoetin beta) Retacrit (epo	petin alfa-epbx)		

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB				
G. CLINICAL INFORMATION (Continued)	■ Required clinical information must be €	l completed in its entirety for all p	recertification requests.				
G. CLINICAL INFORMATION (Continued) – Required clinical information must be completed in its entirety for all precertification requests. Yes							
-	time on therapy:/	1 1					
Yes No Does the patient experience shortness of breath, weakness, fatigue, or lightheadedness from anemia? Please indicate which of the following symptoms the patient experiences: shortness of breath weakness fatigue lightheadedness lightheadedness No Are any of the above symptoms affecting the patient's ability to perform activities of daily living? Yes No Does the patient exhibit angina, syncope, or tachycardia from anemia? Please indicate which of the following symptoms of anemia the patient exhibits: angina syncope tachycardia							
Which of the following laboratory test(s) has the Check all that apply and supply date and result	e patient had within the past 12 months? ts: - Date of test / Please t / Please indicate the T) - Date of test / Ple	se indicate the result:ng/ml e result:ng/mL					
☐ Anemia of Prematurity: Please indicate the patient's birth weight							
Yes ☐ No Is the patient actively Date of most recent of ☐ Yes ☐ No Is the intent of the tre	notherapy Induced Anemia (solid tumors eatment to decrease the need for transfusion receiving chemotherapy? Chemotherapy treatment//	ns in persons who will receive che					
Continuation of treatment: ☐ Yes ☐ No. Has there been a dec	crease in the need for transfusions in patien	nts who are receiving chemothera	nv?				
☐ Chronic Kidney Disease (CKD / ESRD) In Yes ☐ No Is the patient currently Please indicate the particular properties of the partic	nduced Anemia: y receiving dialysis? atient's creatinine clearance:mL/min atient's glomerular filtration:mL/min l/A Based on the decline rate of Hgb level this request be used to reduce the risk of al tinuation request for a member currently on at apply to the patient: acute myocardial	n Date of test // /1.73m² Date of test // s is there a likelihood of red blood lloimmunization and/or other RBC in dialysis? infarction (AMI) orthostatic hitton of greater than 6000ft	/ d cell transfusion? c transfusion-related risks?				
	Anemia: ng interferon or pegylated interferon plus rik ess than10 g/dL despite a reduction in the o						
☐ Human Immunodeficiency Virus (HIV) Dis Endogenous EPO level:mIU/m ☐ Yes ☐ No Is the patient current! ☐ Yes ☐ No Is the current zidovud	L Date of test ///	eek?					
☐ Yes ☐ No Does the bone marrow ☐ Yes ☐ No Has the patient requirements ☐ Yes ☐ No Has the patient requirements ☐ Yes ☐ No Have the transfusion ☐ Myelofibrosis-associated Anemia:	(EPO) levels are less than or equal to 500 _mIU/mL Date of test/ / whave less than 15% blasts? red a blood transfusion of 2 or fewer units or requirements been reduced by less than 50 graphs.	 of blood per month?					
Endogenous EPO level:mIU/m Yes \[\] No Is the member transfu	L Date of test / / usion dependent?						



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB				
G. CLINICAL INFORMATION (Continued)	- Required clinical information must be	completed in its <u>entirety</u> for all p	recertification requests.				
Miscellaneous Induced Anemias: Check all that apply and supply reque		o underlying chronic discoso:					
☐ The underlying chronic disease has been identified. —> Please identify the underlying chronic disease:							
The patient is scheduled to undergo high-risk surgery. —> Is there an increased risk of or intolerance to blood transfusions? 🗌 Yes 🔲 No							
Date of surgery/ Type of surgery:							
Continuation of Treatment:							
☐ Yes ☐ No Has the patient's hemoglobin (Hgb) risen by at least 1 g/dL while on erythropoietin stimulating treatment? If no, please supply rationale for continuation of treatment request:							
If yes, please indicate	the pre-treatment hemoglobin level:	g/dL Date obtained:/					
H. ACKNOWLEDGEMENT							
Request Completed By (Signature Require	red):		Date:/				
Any person who knowingly files a request fo any insurance company by providing materia insurance act, which is a crime and subjects	ally false information or conceals materia	il information for the purpose of					

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