

MEDICARE FORM

Erythropoiesis Stimulating Agents Injectable Medication Precertification Request

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

For New Jersey HMO D-SNP: FAX: 1-833-322-0034 PHONE: 1-844-362-0934

For other lines of business: Please use other form

Note: Epogen and Retacrit are non-preferred. The preferred products are Aranesp and Procrit.

Please indicate: [_// st treatment/				·	
Precertification Req	uested By:			Phone:		Fax:		
A. PATIENT INFOR	MATION							
First Name:			Last Name:			DOB:		
Address:				City:		State:	ZIP:	
Home Phone:		Work Phone:		Cell Phone:		Email:		
Current Weight:	lbs orkgs	Height:	inches orcm	s Allergies:				
B. INSURANCE INF	ORMATION							
			Does patient have other coverage? ☐ Yes ☐ No					
Aetna Member ID #: Group #:			If yes, provide ID#: Carrier Name:					
Insured:			Insured:	_				
C. PRESCRIBER IN	FORMATION							
First Name:			Last Name:	С	heck One:	☐ M.D. ☐ D.O. ☐ N.P. ☐ P.A.		
Address:				City:		State:	ZIP:	
Phone:	Fax:		St Lic #:	NPI #:	DEA #:		UPIN:	
Provider Email:			e Contact Name:		Phone:			
D. DISPENSING PR	OVIDER/ADMINIST	RATION INFOR	MATION					
□	center c: checker phone: checker phone: checker pe(s) (CPT): checker pe(s) (CPT): checker pe(s) (CPT): checker pe(s) (CPT): checker procedure proc	State: ZII Fax: PIN: etin alfa)	P: ogen (epoetin alfa) [acrit (epoetin alfa-ep	☐ Outpatient Dialysis Co ☐ Retail Pharmacy ☐ Mail Order Name:	Sta	pecialty Pharma ther: te: Z Fax: PIN:	IP:beta)	
Primary ICD Code:		Secondar	ry ICD Code:	Othe	er ICD Code	I		
G. CLINICAL INFO	RMATION - Require	d clinical informat	tion must be completed	d in its <u>entirety</u> for all pre	certification	requests.		
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?								



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB				
G. CLINICAL INFORMATION (Continued)	- Paguired clinical information must be	completed in its entirety for all n	propertification requests				
	·	completed in its <u>entirety</u> for all p	necertification requests.				
Yes No Is this request for Epogen (epo	oeun ana) <i>?</i> It with Aranesp (darbepoetin alfa), Procrit (d	enoetin alfa) or Retacrit (enoetin	alfa-enbx) ineffective?				
	No Was treatment with Aranesp (darbepoe						
	not tolerated, or contraindicated?	, , , , , , , , , , , , , , , , , , , ,	, ,				
	ct: not tolerated contraindicated						
	time on therapy: / _ /						
Yes No Does the patient experience s							
> Please indicate which of the	following symptoms the patient experience		eakness Ihtheadedness				
☐ Yes ☐ No. Are any of the	above symptoms affecting the patient's at	_ ` `	,				
☐ Yes ☐ No Does the patient exhibit anging		,	9				
> Please indicate which of the	following symptoms of anemia the patient ϵ	exhibits: angina syncope	☐ tachycardia				
Which of the following laboratory test(s) has the							
Check all that apply and supply date and result	ts:						
☐ Iron Stores from Bone Marrow Iron	- Date of test / Please indicate the	e indicate the result:ng/mL	-				
Serum Transferrin Saturation (TSA	T) - Date of test / / Please indicate the	ase indicate the result: %					
Please choose from one of the indications b							
☐ Anemia of Prematurity:							
Please indicate the patient's birth weigh	ght in grams:						
Please indicate the patient's gestation							
☐ Antineoplastic / Myelosuppressive Chem	notherapy Induced Anemia (solid tumors atment to decrease the need for transfusio						
Yes No Is the patient actively		its in persons who will receive chi	emotrierapy :				
	chemotherapy treatment / /						
☐ Yes ☐ No Is the intent of the tre							
Yes No Is the planned chemo	therapy treatment regimen to continue for	a minimum of 2 months?					
Continuation of treatment:			2				
	rease in the need for transfusions in patier	its who are receiving chemothera	py?				
☐ Chronic Kidney Disease (CKD / ESRD) In ☐ Yes ☐ No Is the patient currentl							
Please indicate the p	atient's creatinine clearance:mL/min	Date of test ///	<u></u>				
	atient's glomerular filtration:mL/min/						
	/A Based on the decline rate of Hgb level						
	his request be used to reduce the risk of al tinuation request for a member currently on		; transfusion-related risks?				
The No is this a con	at apply to the patient: ☐ acute myocardial	infarction (AMI)	ypotension □ angina				
onesk un un		tion of greater than 6000ft	ypotonoion				
	☐ anemia with Hgb	less than 11g/dL has significantly	interfered with activities of daily living				
☐ Hepatitis C with Chemotherapy Induced							
	ng interferon or pegylated interferon plus rik						
	ess than10 g/dL despite a reduction in the	Jose of fibaviliti?					
Human Immunodeficiency Virus (HIV) Di Endogenous EPO level:mIU/m							
Yes No Is the patient currently							
<u> </u>	line dose less than or equal to 4200 mg/we	eek?					
☐ Myelodysplastic Syndrome Induced Ane	mia:						
☐ Endogenous serum erythropoietin (EPO) levels are less than or equal to 500 IU/L.							
Endogenous EPO level:mIU/mL Date of test/							
☐ Yes ☐ No Does the bone marrow have less than 15% blasts? ☐ Yes ☐ No Has the patient required a blood transfusion of 2 or fewer units of blood per month?							
For Continuation of Therapy:							
	requirements been reduced by less than 5	0% after 6 months of therapy?					
☐ Myelofibrosis-associated Anemia:							
Endogenous EPO level:mIU/m Yes No Is the member transfu	L Date of test/ usion dependent?						



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G. CLINICAL INFORMATION (Continued)	Required clinical information must be	completed in its entirety for all p	precertification requests.				
☐ Miscellaneous Induced Anemias: Check all that apply and supply requested information: ☐ The underlying chronic disease has been identified. →> Please identify the underlying chronic disease: ☐ The patient cannot or will not receive whole blood or components as replacement for traumatic/surgical blood loss. ☐ 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 / <							
☐ 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: Yes ☐ No Has the patient's hemoglobin (Hgb) risen by at least 1 g/dL while on erythropoietin stimulating treatment?							
If yes, please indicate the pre-treatment hemoglobin level:g/dL Date obtained:/							
H. ACKNOWLEDGEMENT							
Request Completed By (Signature Requi	red):		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.