

## **MEDICARE FORM**

## **Erythropoiesis Stimulating Agents Injectable** Medication Precertification Request

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

Virginia (HMO D-SNP) FAX: 1-833-280-5224 PHONE: 1-855-463-0933

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:	☐ Start of treatment: Start date _ ☐ Continuation of therapy: Date of		· /				
Precertification R	equested By:		Phone:		Fax:		
A. PATIENT INFO	ORMATION						
First Name:		Last Name:			DOB:		
Address:			City:		State:	ZIP:	
Home Phone:	Work Phone	:	Cell Phone:		Email:		
Current Weight:	lbs orkgs Height:	inches orc	ms Allergies:				
B. INSURANCE	INFORMATION						
Aetna Member ID	) #:	Does patient have o	Does patient have other coverage?				
		If yes, provide ID#: Carrier Name:					
Insured:		Insured:					
C. PRESCRIBER		Leat Name:		Chaoli Onei			
First Name:		Last Name:	0:1	Check One.		D.O. N.P. P.A.	
Address:			City:		State:	ZIP:	
Phone:	Fax:	St Lic #:	NPI #:	DEA #:		UPIN:	
Provider Email:	O PROVIDER/ADMINISTRATION INF	ffice Contact Name:		Phone:			
Outpatient Infus Center Na Home Infusion Agency N Administration Address: City: Phone: TIN: NPI: E. PRODUCT INI Request is for:	ame:	_ ZIP: Epogen (epoetin alfa) Retacrit (epoetin alfa-e may delay request)	Retail Pharma     Mail Order     Name: Address: City: Phone: TIN: NPI: MPI: Mircera (methorepbx)	Sta	Specialty Pha Dther: ate: _ Fax: _ PIN: glycol/epoc	armacy ZIP: etin beta)	
	NFORMATION - Please indicate prim						
,	e: Secor FORMATION - Required clinical infor	ndary ICD Code:					
For All Requests: Yes No W Yes No Is Yes No Is For Initial Reques Note: Epogen and Yes No Ha Yes No Ha	(Clinical documentation required to ill Aranesp (darbepoetin alfa), Procrit (d r Retacrit (epoetin alfa-epbx) be used of the patient currently taking iron supple Hemoglobin (Hgb) result?mg/dl ts: I Retacrit are non-preferred. The preference as the patient had prior therapy with the as the patient had a trial and failure, int Aranesp (darbepoetin alfa) _ Pro- nere are any other medical reason(s) the	for all requests) epoetin alfa), Epogen (eponcomitantly? ments? L Date of test // ferred products are Ara e requested product within olerance, or contraindicat port (epoetin alfa) nat the patient cannot use	oetin alfa), Mircera (r / nesp and Procrit. Pr n the last 365 days? tion to any of the follo	nethoxy polyethyler referred products of wing? (select all the	ne glycol/epo may vary ba at apply)	sed on indication.	



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB					
G. CLINICAL INFORMATION (Continued)	- Required clinical information must be o	completed in its <u>entirety</u> for all p	recertification requests.					
□ Yes       No       Is this request for Epogen (epoetin alfa)?         □ Yes       No       Was treatment with Aranesp (darbepoetin alfa), Procrit (epoetin alfa), or Retacrit (epoetin alfa-epbx) ineffective?         □ Yes       No       Was treatment with Aranesp (darbepoetin alfa), Procrit (epoetin alfa), or Retacrit (epoetin alfa-epbx) ineffective?         □ Yes       No       Was treatment with Aranesp (darbepoetin alfa), Procrit (epoetin alfa), or Retacrit (epoetin alfa-epbx) not tolerated, or contraindicated?         □ Please select:       □ not tolerated       □ contraindicated								
		/ /						
Please indicate the length of time on therapy:/ // / 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								
Yes No Does the patient exhibit angin	following symptoms of anemia the patient e		-					
Which of the following laboratory test(s) has the patient had within the past 12 months?         Check all that apply and supply date and results:         Iron Stores from Bone Marrow Iron - Date of test       /         Serum Ferritin Levels - Date of test       /         Serum Transferrin Saturation (TSAT) - Date of test       /								
Please choose from one of the indications b								
<ul> <li>Anemia of Prematurity:         <ul> <li>Please indicate the patient's birth weight in grams:</li> <li>Please indicate the patient's gestational age in weeks:</li> </ul> </li> <li>Antineoplastic / Myelosuppressive Chemotherapy Induced Anemia (solid tumors, multiple myeloma, lymphoma, lymphocytic leukemia):         <ul> <li>Yes</li> <li>No</li> <li>Is the intent of the treatment to decrease the need for transfusions in persons who will receive chemotherapy?</li> <li>Yes</li> <li>No</li> <li>Is the patient actively receiving chemotherapy?</li> </ul> </li> </ul>								
Yes No Is the intent of the tre	chemotherapy treatment /// eatment to be curative? otherapy treatment regimen to continue for							
	crease in the need for transfusions in patier	its who are receiving chemothera	py?					
□ Chronic Kidney Disease (CKD / ESRD) Ir	aduced Anemia:         y receiving dialysis?         atient's creatinine clearance:mL/min         atient's glomerular filtration:mL/min/         I/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        living at an elevat        anemia with Hgb	Date of test // 1.73m <sup>2</sup> Date of test // s is there a likelihood of red blood lloimmunization and/or other RBC n dialysis? infarction (AMI)	/ d cell transfusion? c transfusion-related risks?					
	Anemia: ng interferon or pegylated interferon plus rik ess than10 g/dL despite a reduction in the e							
□ Human Immunodeficiency Virus (HIV) Di Endogenous EPO level:mIU/m □ Yes □ No Is the patient current □ Yes □ No Is the current zidovuc	L Date of test / /	ek?						
↓       Endogenous EPO level:         ↓       Yes       No       Does the bone marro         ↓       Yes       No       Has the patient requi         For Continuation of Therapy:       ↓       Yes       No         ↓       Yes       No       Have the transfusion         ↓       Yes       No       Have the transfusion	(EPO) levels are less than or equal to 500 _mIU/mL Date of test/ / w have less than 15% blasts? red a blood transfusion of 2 or fewer units of requirements been reduced by less than 50	_ of blood per month?						
Endogenous EPO level:mIU/m								



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G. CLINICAL INFORMATION (Continued)	- Required clinical information must be	e completed in its <u>entirety</u> for	all precertification requests.					
<ul> <li>The patient cannot or will not rece</li> <li>The patient is scheduled to undergo</li> </ul>	is been identified. $\longrightarrow$ Please identify t ive whole blood or components as replace go high-risk surgery. $\longrightarrow$ Is there an in	ement for traumatic/surgical blo creased risk of or intolerance to	od loss. b blood transfusions?  Yes No					
Date of surgery/ Type of surgery:								
Continuation of Treatment:	noglobin (Hgb) risen by at least 1 g/dL wh ationale for continuation of treatment requ	ile on erythropoietin stimulating	g treatment?					
If yes, please indicate	the pre-treatment hemoglobin level:	g/dL Date obtained: /	1					
H. ACKNOWLEDGEMENT								
Request Completed By (Signature Requi	Date: / /							
Any person who knowingly files a request for any insurance company by providing materi								

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.