

MEDICARE FORM Erythropoiesis Stimulating Agents, HIF Inhibitors Medication Precertification Request

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

For Medicare Advantage Part B: For other lines of business: Please use commercial form.

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

Would you like to use electronic prior authorization? Consider using **Availity**, our electronic prior authorization portal. Learn more about **Availity** from the links in the table below.

For phone or fax requests, refer to the table below for routing information. To determine which box to use, refer to the patient's Aetna ID card. State specific special needs and Medicare-Medicaid Plans may be designated on the front of the ID card or in the website URL on the back of the card. If you don't see your specific plan listed, call the number on the back of the member's ID card to confirm routing information.

For Aetna Medicare Advantage and Allina Health Aetna Medicare members send request to:

Phone: 1-866-503-0857 (TTY: 711)

Fax: 1-844-268-7263

Availity: https://www.aetna.com/health-care-professionals/resource-center/availity.html

For Aetna Medicare Advantage Virginia Dual Eligible Special Needs Plans (HMO D-SNP)

send request to:

Phone: <u>1-855-463-0933</u> Fax: 1-833-280-5224

Availity: https://www.aetnabetterhealth.com/virginia-hmosnp/providers/portal

For Aetna Assure Premier Plus Medicare Advantage New Jersey Dual Eligible Special Needs Plans

(HMO D-SNP) send request to:

Phone: <u>1-844-362-0934</u> Fax: 1-833-322-0034

Availity: https://www.aetnabetterhealth.com/new-jersey-hmosnp/providers/portal.html

For Aetna Better Health of Illinois Premier Medicare Medicaid Plan (MMP) send request to:

Phone: <u>1-866-600-2139</u> FAX: <u>1-855-320-8445</u>

Availity: https://www.aetnabetterhealth.com/illinois/providers/portal

For Aetna Better Health of Ohio Premier Medicare Medicaid Plan (MMP) send request to:

Phone: <u>1-855-364-0974</u> Fax: <u>1-855-734-9389</u>

Availity: https://www.aetnabetterhealth.com/ohio/providers/portal

For Aetna Better Health of Michigan Premier Medicare Medicaid Plan (MMP) send request to:

Phone: <u>1-855-676-5772</u> Fax: <u>1-844-241-2495</u>

Availity: https://www.aetnabetterhealth.com/michigan/providers/portal.html



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Dose/Frequency: (Failure to provide dose & frequency may delay request) F. DIAGNOSIS INFORMATION - Please indicate primary ICD code and specify any other where applicable. Primary ICD Code: Secondary ICD Code: Other ICD Code: G. CLINICAL INFORMATION - Required clinical information must be completed in its entirety for all precertification 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?							
Last Name:							
Last Name: City: State: ZIP:							
City: State: ZIP:							
Mork Phone:							
Dispensing Provider/Pharmacy:							
Does patient have other coverage? Yes No Group #: Insured: Insure							
Actina Member ID #:							
If yes, provide ID#;							
Insured: C. PRESCRIBER INFORMATION First Name: Last Name: Check One: M.D. D.O. N.	· ·						
C. PRESCRIBER INFORMATION First Name:							
First Name:							
Phone: Fax: St Lic #: NPI #: DEA #: UPIN: Provider Email: Office Contact Name: Phone: D. DISPENSING PROVIDER/ADMINISTRATION INFORMATION Place of Administration: Genter Physician's Office Home Genter Physician's Office Phone: Genter Name:	.P. 🗌 P.A.						
Provider Email:							
D. DISPENSING PROVIDER/ADMINISTRATION INFORMATION Place of Administration: Gelf-administered Physician's Office Home Outpatient Dialysis Center Physician's Office Gutpatient Infusion Center Outpatient Dialysis Center Physician's Office Retail Pharmacy Specialty Pharmacy Specialty Pharmacy Other: Name: Administration code(s) (CPT): Address: Address: City: State: ZIP: Phone: Fax: TIN: PIN: TIN: TIN: PIN: TIN: TIN: PIN: TIN: TI							
Place of Administration: Gelf-administered Physician's Office Home Outpatient Dialysis Center Physician's Office Gutpatient Infusion Center Phone: Retail Pharmacy Specialty Pharmacy Mail Order Other: Name: Administration code(s) (CPT): Address: ZIP: Address: ZIP: Phone: Fax: ZIP: Phone: Phon							
Self-administered Physician's Office Home Outpatient Dialysis Center Physician's Office Outpatient Influsion Center Center Name: Retail Pharmacy Specialty Phar							
City:							
Phone: Fax: TIN: PIN: PIN: NPI: TIN: NPI: PIN: NPI: NPI							
NPI:							
E. PRODUCT INFORMATION Request is for: Aranesp (darbepoetin alfa) Epogen (epoetin alfa) Jesduvroq (daprodustat) Vafseo (vadadustat) Mircera (methoxy polyethylene glycol/epoetin beta) Procrit (epoetin alfa) Retacrit (epoetin alfa-e Dose/Frequency: HCPCS Code: (Failure to provide dose & frequency may delay request) F. DIAGNOSIS INFORMATION - Please indicate primary ICD code and specify any other where applicable. Primary ICD Code: Secondary ICD Code: Other ICD Code: G. CLINICAL INFORMATION - Required clinical information must be completed in its entirety for all precertification 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?							
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☐ 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: Epogen, Jesduvroq, Retacrit and Vafseo are non-preferred. The preferred products are Aranesp and Procrit. Preferred products may vary based on indication. Yes No Has the patient had prior therapy with the requested product within the last 365 days? No Has the patient had a trial and failure of any of the following? (If yes, select all that apply) Aranesp (darbepoetin alfa) Procrit (epoetin alfa) When was the member's trial and failure of the preferred drug? Please describe the nature of the failure of the preferred drug							
No Has the patient had an adverse reaction to any of the following? (If yes, select all that apply) ☐ Aranesp (darbepoetin alfa) ☐ Procrit (epoetin alfa) ───────────────────────────────────							



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB		
G. CLINICAL INFORMATION (Contin	l nued) – Required clinical information m	ust be completed in its entirety	for all precertification requests.		
G. CLINICAL INFORMATION (Continued) – Required clinical information must be completed in its entirety for all precertification requests. Please explain if there are any contraindications or 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) Aranesp (darbepoetin alfa) Procrit (epoetin alfa)					
	· · · /				
☐ 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) ☐ not tolerated, or contraindicated? ☐ Please select: ☐ not tolerated ☐ contraindicated					
	gth 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 Please indicate which	vof the above symptoms affecting the pati angina, syncope, or tachycardia from ane of the following symptoms of anemia the p	emia? patient exhibits: ☐ angina ☐ sy			
	has the patient had within the past 12 mo	nths?			
	resuits: w Iron - Date of test/_/ of test/ Please indi		ng/mL		
	(TSAT) - Date of test //		%		
Please choose from one of the indicat	ions below:				
☐ Anemia of Prematurity: Please indicate the patient's bir Please indicate the patient's ge	th weight in grams: stational age in weeks:				
	chemotherapy treatment regimen to continual and decrease in the need for transfusions in the need for the ne	nsfusions in persons who will red / nue for a minimum of 2 months?	ceive chemotherapy?		
☐ Yes ☐ No☐ Yes, please indicate	urrently receiving dialysis? a the patient's creatinine clearance: the patient's glomerular filtration: N/A Based on the decline rate of How Will this request be used to reduce the rhow long patient has been receiving dialy patient have pretreatment hemoglobin (How the patient have pretreatment hemoglobin (How the patient have pretreatment hemoglobin)	gb levels is there a likelihood of re isk of alloimmunization and/or oth rsis: months Date started	ed blood cell transfusion? ner RBC transfusion-related risks? ://		
-	luced Anemia: eceiving interferon or pegylated interferon Hgb less than10 g/dL despite a reductior				
☐ Yes ☐ No Is the patient c	mIU/mL Date of test / /	_			
☐ Myelodysplastic Syndrome Induce ☐ Endogenous serum erythrop ☐ Endogenous EPO level ☐ Yes ☐ No Does the bone ☐ Yes ☐ No Has the patient For Continuation of Therapy:		to 500 IU/L. / r units of blood per month?	apy?		
☐ Myelofibrosis-associated Anemia: Endogenous EPO level: ☐ Yes ☐ No Is the member	mIU/mL Date of test / / transfusion dependent?	-			



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB
G. CLINICAL INFORMATION (Contin	nued) – Required clinical information must l	pe completed in its entirety f	or all precertification requests.
☐ The patient cannot or will no ☐ The patient is scheduled to u ☐ Date of surgery ☐ Continuation of Treatment: ☐ Yes ☐ No Has the patient ☐ If no, please sup ☐ If yes, please in	requested information: ase has been identified. Please identified. Please identified. Please identified. Indergo high-risk surgery. Is there an ingered in the property of surgery: Property at least 1 g/dL with poply rationale for continuation of treatment required in the pre-treatment hemoglobin level: Please identified in the property of the property in the path of the property in the path of the property is a property of the property of the property in the path of the property in the path of the property	cement for traumatic/surgical ncreased risk of or intolerance while on erythropoietin stimulatuest:g/dL Date obtained:	blood loss. e to blood transfusions? Yes No ting treatment?
H. ACKNOWLEDGEMENT			
Request Completed By (Signature F	Required):		Date: //
any insurance company by providing	uest for authorization of coverage of a me materially false information or conceals ma bjects such person to criminal and civil per	terial information for the pu	with the intent to injure, defraud or deceive rpose of misleading, commits a fraudulent

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