



MEDICARE FORM

Erythropoiesis Stimulating Agents, HIF Inhibitors Medication Precertification Request

Page 1 of 3

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

For Medicare Advantage Part B: Phone: 1-866-503-0857 (TTY:711) FAX: 1-844-268-7263

For other lines of business: Please use other form

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

Please indicate: Start of treatment: Start date Continuation of therapy: Date of last treatment

Precertification Requested By: Phone: Fax:

A. PATIENT INFORMATION

Form section A containing fields for Patient Information: First Name, Last Name, DOB, Address, City, State, ZIP, Home Phone, Work Phone, Cell Phone, Email, Current Weight, Height, Allergies.

B. INSURANCE INFORMATION

Form section B containing fields for Insurance Information: Aetna Member ID #, Group #, Insured, Does patient have other coverage?, Carrier Name.

C. PRESCRIBER INFORMATION

Form section C containing fields for Prescriber Information: First Name, Last Name, Address, City, State, ZIP, Phone, Fax, St Lic #, NPI #, DEA #, UPIN, Provider Email, Office Contact Name, Phone.

D. DISPENSING PROVIDER/ADMINISTRATION INFORMATION

Form section D containing fields for Dispensing Provider/Pharmacy: Place of Administration, Dispensing Provider/Pharmacy, Address, City, State, ZIP, Phone, Fax, TIN, PIN, NPI.

E. PRODUCT INFORMATION

Form section E containing fields for Product Information: Request is for (Aranesp, Epogen, Jesduvrog, Mircerca, Procrit, Retacrit), Dose/Frequency, HCPCS Code.

F. DIAGNOSIS INFORMATION - Please indicate primary ICD code and specify any other where applicable.

Form section F containing fields for Diagnosis Information: 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.

Form section G containing fields for Clinical Information: For All Requests, For Initial Requests, Note: Epogen, Jesduvrog and Retacrit are non-preferred. The preferred products are Aranesp and Procrit.



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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 entirety for all precertification requests.

Is this request for Epogen (epoetin alfa)? Was treatment with Aranesp (darbeoetin alfa), Procrit (epoetin alfa), or Retacrit (epoetin alfa-epbx) ineffective? Please select: not tolerated, contraindicated. Please indicate the length of time on therapy: / / - / /

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. Are any of the above symptoms affecting the patient's ability to perform activities of daily living?

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 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 / / Please indicate the result: ng/mL
Serum Ferritin Levels - Date of test / / Please indicate the result: ng/mL
Serum Transferrin Saturation (TSAT) - Date of test / / Please indicate the result: %

Please choose from one of the indications below:

Anemia of Prematurity:

Please indicate the patient's birth weight in grams: Please indicate the patient's gestational age in weeks:

Antineoplastic / Myelosuppressive Chemotherapy Induced Anemia (solid tumors, multiple myeloma, lymphoma, lymphocytic leukemia):

- Is the intent of the treatment to decrease the need for transfusions in persons who will receive chemotherapy?
Is the patient actively receiving chemotherapy? Date of most recent chemotherapy treatment / /
Is the intent of the treatment to be curative?
Is the planned chemotherapy treatment regimen to continue for a minimum of 2 months?

Continuation of treatment:

Has there been a decrease in the need for transfusions in patients who are receiving chemotherapy?

Chronic Kidney Disease (CKD / ESRD) Induced Anemia:

- Is the patient currently receiving dialysis? Please indicate the patient's creatinine clearance: mL/min Date of test / / Please indicate the patient's glomerular filtration: mL/min/1.73m2 Date of test / /
Based on the decline rate of Hgb levels is there a likelihood of red blood cell transfusion?
Will this request be used to reduce the risk of alloimmunization and/or other RBC transfusion-related risks?
If yes, please indicate how long patient has been receiving dialysis: months Date started: / /
Does patient have pretreatment hemoglobin (Hgb) less than or equal to 11 g/dL? g/dL Date of test / /

Hepatitis C with Chemotherapy Induced Anemia:

- Is the patient receiving interferon or pegylated interferon plus ribavirin?
Is the patient's Hgb less than 10 g/dL despite a reduction in the dose of ribavirin?

Human Immunodeficiency Virus (HIV) Disease Induced Anemia:

- Endogenous EPO level: mIU/mL Date of test / /
Is the patient currently receiving zidovudine?
Is the current zidovudine dose less than or equal to 4200 mg/week?

Myelodysplastic Syndrome Induced Anemia:

- Endogenous serum erythropoietin (EPO) levels are less than or equal to 500 IU/L. Endogenous EPO level: mIU/mL Date of test / /
Does the bone marrow have less than 15% blasts?
Has the patient required a blood transfusion of 2 or fewer units of blood per month?

For Continuation of Therapy:

Have the transfusion requirements been reduced by less than 50% after 6 months of therapy?

Myelofibrosis-associated Anemia:

- Endogenous EPO level: mIU/mL Date of test / /
Is the member transfusion dependent?

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G. CLINICAL INFORMATION (Continued) – Required clinical information must be completed in its entirety for all 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 ____/____/____ 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: ____/____/____
- Yes No Has the requested product been effective for treating the patient's diagnosis or condition?

H. ACKNOWLEDGEMENT

Request Completed By (Signature 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.