

Jakafi (Ruxolitinib) **Prior Authorization Request Form**



Caterpillar Prescription Drug Benefit

Instructions: Please fill out all applicable sections completely and legibly. Attach any additional documentation that is important for the review (e.g., chart notes or lab data, to support the authorization request). Information contained in this form is Protected Health Information under HIPAA.

| MEMBER INFORMATION | | | | |
|---|------------------------|----------------------------|------------------|--|
| LAST NAME: | | FIRST NAME: | | |
| PHONE NUMBER: | | DATE OF BIRTH: | | |
| STREET ADDRESS: | | • | | |
| CITY: | | STATE: ZIP C | ODE: | |
| PATIENT INSURANCE ID NU | MBER: | • | | |
| | GHT (IN/CM): WEIG | | | |
| FOLLOWING LINK: HTTPS://MAGELLANRX.COM/MEMBER/EXTERNAL/COMMERCIAL/COMMON/DOC/EN-US/PHI DISCLOSURE AUTHORIZATION.PDF | | | | |
| PATIENT'S AUTHORIZED REPRESENTATIVE (IF APPLICABLE):AUTHORIZED REPRESENTATIVE'S PHONE NUMBER: | | | | |
| AUTHORIZED REPRESENTATI | VE 3 PHONE NOWIBER. | | | |
| PRESCRIBER INFORMATION | | | | |
| LAST NAME: | | FIRST NAME: | | |
| PRESCRIBER SPECIALTY: | | EMAIL ADDRESS: | | |
| NPI NUMBER: | | DEA NUMBER: | | |
| PHONE NUMBER: | _ | FAX NUMBER: | | |
| STREET ADDRESS: | | • | | |
| CITY: | | STATE: ZIP CODE: | | |
| REQUESTOR (if different than prescriber): | | OFFICE CONTACT PERSON: | | |
| | | | | |
| MEDICATION OR MEDICAL | DISPENSINGINFORMATION | | | |
| MEDICATION NAME: | | | | |
| DOSE/STRENGTH: | FREQUENCY: | LENGTH OF THERAPY/REFILLS: | QUANTITY: | |
| NEW THERAPY DURATION OF THERAPY (SPI | RENEWAL ECIFIC DATES): | IF RENEWAL: DATE THE | ERAPY INITIATED: | |

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| MEMBER'S LAST NAME: MEMBER'S FIRST NAME: | | | | |
|---|---|--------------------------------------|--|--|
| 1. HAS THE PATIENT TRIED ANY OTHER | R MEDICATIONS FOR THIS CONDITION? | YES (if yes, complete below) NO | | |
| MEDICATION/THERAPY (SPECIFY DRUG NAME AND DOSAGE): | DURATION OF THERAPY (SPECIFY DATES): | RESPONSE/REASON FOR FAILURE/ALLERGY: | | |
| 2. LIST DIAGNOSES: | | ICD-10: | | |
| □ Myelofibrosis □ Polycythemia vera □ Acute Graft-versus-host-disease (GVHD) □ Chronic Graft vs. Host Disease (cGVHD) □ Other diagnosis:ICD | -10 | | | |
| 3. REQUIRED CLINICAL INFORMATION PRIOR AUTHORIZATION. | I: PLEASE PROVIDE ALL RELEVANT CLINIC | CALINFORMATION TO SUPPORT A | | |
| For myelofibrosis, answer the following: Does the patient have a diagnosis of intermediate or high risk myelofibrosis (including primary myelofibrosis, postpolycythemia vera myelofibrosis or post-essential thrombocythemia myelofibrosis)? Yes No | | | | |
| Has the diagnosis of myelofibrosis been verified with a bone marrow biopsy? ☐ Yes ☐ No Please include a copy of the bone marrow biopsy report | | | | |
| Does the patient have abnormal blood cells, confirmed by a peripheral blood smear report? Yes No Please include a copy of the peripheral blood smear report | | | | |
| For polycythemia vera, answer the following: Does the patient have documentation of polycythemia vera? □ Yes □ No | | | | |
| Has the patient had an inadequate response or intolerance to hydroxyurea? ☐ Yes ☐ No | | | | |
| For acute graft-versus-host-disease(G | VHD), answer the following: | | | |
| Has patient had a history of hematologic malignancy? ☐ Yes ☐ No Please provide documentation. | | | | |
| Has patient undergone ONLY one allogeneic hematopoetic stem cell transplant? \Box Yes \Box No <i>Please provide documentation</i> . | | | | |
| Does patient have GVHD overlap ("acute on chronic") syndrome (as defined by NIH guidelines)? \square Yes \square No Please provide documentation. | | | | |
| Does the patient's acute GVHD involve the liver and/or upper GI tract and/or lower GI tract and/or >50% body surface area of the skin? No Please provide documentation. | | | | |
| Has the patient's acute GVHD progressed after 3 or more days of methylprednisolone > 2mg/kg/day(or equivalent)? \Box Yes \Box No Please provide documentation. | | | | |





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| Prescriber Signature or Electronic I.D. Verification: CONFIDENTIALITY NOTICE: The desuments assemble this transmission contain confidential and the second | |
|---|--|
| ATTESTATION: I attest the information provided is true and accurate to the bethe Health Plan, insurer, Medical Group or its designees may perform a routine information necessary to verify the accuracy of the information reported on the information repo | e audit and request the medical his form. |
| Please note: Not all drugs/diagnosis are covered on all plans. This request may information is received. | y be denied unless all required |
| Are there any other comments, diagnoses, symptoms, medications tried or f physician feels is important to this review? | failed, and/or any other information the |
| Renewal for Graft versus Host Disease: Does the patient have active disease? Yes No Is the patient requiring a taper dose schedule in order to wean off Jakafi(rux) | olitinib)? □ Yes □ No |
| Does the patient have a confirmed diagnosis of glucocorticoid-refractory or g defined per 2014 National Institutes of Health (NIH) consensus criteria as a la after administration of minimum prednisone 1 mg/kg/dayfor≥1 week (or exwithout improvement despite continued treatment with prednisone at >0.5 for ≥4 weeks (or equivalent) OR Increase to prednisone dose to >0.25 mg/kg to taper the dose (or equivalent) □ Yes □ No | ack of response or disease progression quivalent) OR disease persistence mg/kg/dayor 1 mg/kg/every other day |
| Has the patient had a trial with systemic corticosteroids for the treatment of | fcGVHD? □ Yes □ No |
| Has the patient undergone an allogeneic stem cell transplantation? \Box Yes | □ No |
| For Chronic Graft vs. Host Disease (cGVHD), answer the following: | |
| Has the patient been unable to tolerate tapering from corticosteroids? \Box Ye documentation. | es □ No <i>Please provide</i> |
| Has the patient's acute GVHD developed in another organ after receiving me equivalent) for skin GVDH or skin/upper GI GVDH? Yes No Please provide | |
| Did the patient's acute GVHD show no improvement after 7 days of methylp equivalent)? Yes No Please provide documentation. | oreanisolone > 2mg/kg/day(or |

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FAX THIS FORM TO: 800-424-7640

MAIL REQUESTS TO: Magellan Rx Management Prior Authorization Program

Attn:CP-4201 P.O.Box 64811 St. Paul, MN 55164-0811

Phone: 877-228-7909

