SENTARA COMMUNITY PLAN (MEDICAID)

PHARMACY PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

Directions: The prescribing physician must sign and clearly print name (preprinted stamps not valid) on this request. All other information may be filled in by office staff; fax to 1-800-750-9692. No additional phone calls will be necessary if all information (including phone and fax #s) on this form is correct. If the information provided is not complete, correct, or legible, the authorization process can be delayed.

Drug Requested: Graft-Versus-Host Disease (GVHD) Drugs (select drug below) □ **Imbruvica**® (ibrutinib) □ **Jakafi**[®] (ruxolitinib) **❖ FOR AN ONCOLOGY INDICATION, PLEASE REFER TO THE FOLLOWING** PRIOR AUTHORIZATION FORM: PAOralOncology.pdf (sitecorecontenthub.cloud) MEMBER & PRESCRIBER INFORMATION: Authorization may be delayed if incomplete. Member Name: _____ Member Sentara #: Date of Birth: Prescriber Name: Prescriber Signature: Date: Office Contact Name: _____ Phone Number: Fax Number: DEA OR NPI #: _____ **DRUG INFORMATION:** Authorization may be delayed if incomplete. Drug Form/Strength: Dosing Schedule: _____ Length of Therapy: _____ ICD Code, if applicable: _____ Diagnosis: Weight (within last 30 days: **Maximum Allowable Daily Dosage:**

- Jakafi[®] (ruxolitinib):
 - ☐ Acute & Chronic GVHD: 20 mg per day
- Imbruvica® (ibrutinib):
 - ☐ Chronic GVHD: 420 mg per day

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suppo	NICAL CRITERIA: Check below all that apply. All criteria must be met for approval. To ort each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be ided or request may be denied.				
	Provider is an oncologist/hematologist				
	Complete subsequent criteria for the applicable indication below:				
□ D	Diagnosis: Acute Graft-Versus-Host Disease (aGVHD) - Jakafi® (ruxolitinib) ONLY				
<u>Initi</u>	ial Authorization: 6 months				
	Medication will be used for disease related to allogeneic hematopoietic stem cell transplantation				
	Members is at least 12 years of age or older				
	Member has acute graft versus host disease (aGVHD) that is refractory to treatment with corticosteroids				
	Provider has submitted recent progress notes and/or clinical assessment recording the symptomology and staging/grading of acute GVHD organ involvement				
□ D	Diagnosis: Acute Graft-Versus-Host Disease (aGVHD) - Jakafi® (ruxolitinib) ONLY				
Rea	uthorization: 6 months				
	Member has experienced treatment response as evidenced by stabilization or improvement in disease (please submit recent progress notes and/or clinical assessment recording improvement in aGVHI organ involvement)				
	ONE of the following must be met:				
	☐ Member has been able to discontinue therapeutic doses of corticosteroids, <u>AND</u> additional therapy authorization will be utilized for tapering ruxolitinib. NOTE: Taper by one dose level approximately every 8 weeks (10 mg twice daily to 5 mg twice daily to 5 mg once daily)				
	☐ Member requires re-treatment because aGVHD signs/symptoms recurred during or after the tapering of ruxolitinib (please submit recent progress notes and/or clinical assessment recording changes/worsening of aGVHD organ involvement)				
	Diagnosis: Chronic Graft-Versus-Host Disease (cGVHD) - Jakafi® (ruolitinib) or mbruvica® (ibrutinib)				
<u>Initi</u>	ial Authorization: 6 months				
For Ja	akafi® Requests:				
	Medication will be used for disease related to allogeneic hematopoietic stem cell transplantation				
	Member is at least 12 years of age or older				
	Member has chronic graft versus host disease (cGVHD) that is refractory to treatment with corticosteroids				

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Member has failed one or two lines of systemic therapy (e.g., mycophenolate, methotrexate) for the treatment of chronic graft versus host disease (cGVHD) <u>AND</u> will be used in combination with systemic corticosteroids
 Provider has submitted recent progress notes and/or clinical assessment recording the symptomology and staging/severity of chronic GVHD (i.e. NIH Global Severity Score, NIH Organ-specific Score)

For Imbruvica® Requests:

- ☐ Medication will be used for disease related to allogeneic hematopoietic stem cell transplantation
- ☐ Member is at least 1 year of age or older
- ☐ Medication will be used as a single agent or in conjunction with systemic steroids
- ☐ Member has failed one or more previous lines of systemic therapy for the treatment of cGVHD (e.g., corticosteroids or immunosuppressant's such as cyclosporine)
- □ Provider has submitted recent progress notes and/or clinical assessment recording the symptomology and staging/severity of chronic GVHD (i.e. NIH Global Severity Score, or NIH Organ-specific Score)
- □ Diagnosis: Chronic Graft-Versus-Host Disease (cGVHD) Jakafi® (ruolitinib) or Imbruvica® (ibrutinib)

Reauthorization: 6 months

For Jakafi® Requests:

- ☐ Member has experienced treatment response as evidenced by stabilization or improvement in disease [please submit recent progress notes and/or clinical assessment recording the response in symptomology and staging/severity of chronic GVHD (i.e., NIH Global Severity Score, or NIH Organ-specific Score)]
- □ **ONE** of the following must be met:
 - ☐ Member has been able to discontinue therapeutic doses of corticosteroids <u>AND</u> additional therapy authorization will be utilized for tapering ruxolitinib. NOTE: Taper by one dose level approximately every 8 weeks (10 mg twice daily to 5 mg twice daily to 5 mg once daily)
 - ☐ Member requires re-treatment because cGVHD signs/symptoms recurred during or after the tapering of ruxolitinib [please submit recent progress notes and/or clinical assessment recording the response in symptomology and staging/severity of chronic GVHD (i.e., NIH Global Severity Score, or NIH Organ-specific Score)]

For Imbruvica® Requests:

☐ Member has experienced treatment response as evidenced by stabilization or improvement in disease [please submit recent progress notes and/or clinical assessment recording the response in symptomology and staging/severity of chronic GVHD (i.e. NIH Global Severity Score, or NIH Organ-specific Score)]

Medication being provided by Specialty Pharmacy - PropriumRx

Use of samples to initiate therapy does not meet step edit/ preauthorization criteria.

Previous therapies will be verified through pharmacy paid claims or submitted chart notes.