AvMed

PHARMACY PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

<u>Directions</u>: <u>The prescribing physician must sign and clearly print name (preprinted stamps not valid)</u> on this request. All other information may be filled in by office staff; <u>fax to 1-305-671-0200</u>. No additional phone calls will be necessary if all information (<u>including phone and fax #s</u>) on this form is correct. <u>If the information provided is not complete, correct, or legible, the authorization process can be delayed.</u>

Drug Requested: Pharmacy Benefit Oncology Medications

MEN	MBER & PRESCRIBEI	R INFORMATION: Authorization may be delayed if incomplete.	
Memb	er Name:		
		Date of Birth:	
Prescr	iber Name:		
		Date:	
	Number:		
		uthorization may be delayed if incomplete.	
Drug l	Form/Strength:		
		Length of Therapy:	
	osis:		
	t:		
each l		eck below all that apply. All criteria must be met for approval. To support on, including lab results, diagnostics, and/or chart notes, must be provided	
Initi	al Authorization: 12 mo		
	The requesting provider is an	n oncologist	
	AND		
	Use of the requested oncolog ensure diagnosis is docume	gy therapy is documented in literature and found in ONE of following (please ented above) : dance with a specific indication	
	Accepted off-label indication	on found in the most recent edition of any of the following:	
	☐ American Hospital Form	ulary Service Drug Information	
	□ National Comprehensive	Cancer Network's Drugs & Biologics Compendium	
	☐ Elsevier Gold Standard's	Clinical Pharmacology	
		(Continued on next page)	

OR

For medical necessity (Please provide clinical rationale and submit any chart notes/literature you feel would be pertinent in support of medical necessity. Note: experimental/investigational use as defined by the chemotherapy administration policy precludes medical necessity.)		
AND		
If a biomarker/genetic component is required for the drug's site of action please ensure the following:		
□ Submit/attach all genetic mutation, receptor, biomarker, laboratory documentation using an FDA-approved test including both the results and which test was utilized		
NOTE: Experimental/investigational use as defined by the chemotherapy administration policy precludes medical necessity		
AND		
Member has tried and failed current treatment-guideline and FDA label-recommended first-line agents [or		

AND

☐ Please list all previous chemotherapy regimens and dates (please attach chart notes)

Chemotherapy Regimen	Dates/Cycles Completed
1.	
2.	
3.	
4.	

has a documented intolerance, FDA-labeled contraindication, or hypersensitivity to first line therapies]

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	AND
	Requested dose must meet ONE of the following:
	☐ The quantity (dose) requested is in accordance with FDA approved labeling, and if applicable or necessary, age and weight conditions are met
	What is the quantity requested per DAY?
	OR
	The quantity (dose) requested is higher than the maximum dose recommendation found in FDA approved labeling (i.e., the package insert), and the prescriber has submitted clinical literature and medical documentation in support of the safety and efficacy of the higher dose (such as evidence from practice guidelines or clinical trials from peer-reviewed medical literature)
	** Please note: Chart documentation of the above is required to be submitted along with this request **
	AND
	If requesting Kisqali® (ribociclib) for advanced or metastatic hormone receptor-positive, HER2-negative breast cancer, member must have trial and failure of <u>ONE</u> of the following (verified by chart notes or pharmacy paid claims):
	☐ Ibrance® (palbociclib)
	□ Verzenio® (abemaciclib)
To su	uthorization: 12 months. Check below all that apply. All criteria must be met for approval. apport each line checked, all documentation, including lab results, diagnostics, and/or chart notes, be provided or request may be denied.
	Member is currently receiving the requested agent (please submit medical chart notes and documentation of therapy history)
	AND
	Member requires continuation of therapy and is NOT experiencing disease progression
	AND
	Ongoing treatment is consistent with FDA-labeling or compendia support
	AND

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☐ Member is **NOT** experiencing an FDA-labeled limitation of use or toxicity

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	The qu	uantity (dose) requested is in accordance with FDA approved labeling
	•	IF there is an adjustment in quantity (dose) requested, higher than the maximum dose recommendation in FDA-approved labeling (i.e., the package insert), the prescriber must submit clinical literature and medical documentation in support the safety and efficacy of the higher dose (such as evidence from practice guidelines or clinical trials from peer-reviewed medical literature).
		** Please note: Chart documentation of the above is required to be submitted along with this request **
I	Medic	eation being provided by Specialty Pharmacy – Proprium Rx

Not all drugs may be covered under every Plan

If a drug is non-formulary on a Plan, documentation of medical necessity will be required.

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.