SENTARA HEALTH PLANS

MEDICAL PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

<u>Directions:</u> 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; <u>fax to 1-844-668-1550</u>. No additional phone calls will be necessary if all information (including phone and fax #s) on this form is correct. <u>If information provided is not complete, correct, or legible, authorization can be delayed</u>.

<u>For Medicare Members:</u> Medicare Coverage for outpatient (Part B) drugs is outlined in the Medicare Benefit Policy Manual (Pub. 100-2), Chapter 15, §50 Drugs and Biologicals. In addition, National Coverage Determination (NCD) and Local Coverage Determinations (LCDs) may exist and compliance with these policies is required where applicable. They can be found at: https://www.cms.gov/medicare-coverage-database/overview-and-quick-search.aspx. Additional indications may be covered at the discretion of the health plan.

Pemetrexed Injections

The Sentara Health Plans Oncology Program is administered by OncoHealth

- ❖ For any oncology indications, the most efficient way to submit a prior authorization request is through the OncoHealth OneUM Provider Portal at https://oneum.oncohealth.us.Fax to 1-800-264-6128.
 OncoHealth can also be contacted by Phone: 1-888-916-2616.
- ❖ Commercial customers <u>NOT</u> enrolled in the OncoHealth program, please fax requests to Sentara Health plans at fax number 1-800-750-9692.

Drug Requested: (Please select drug below)

	PRE	FERRED [Aut	horization Requ	uired]	
	J9294 Pemetrexed	□ J9296 Pemetrexed		□ J9297 Pemetrexed	
	Injection (Hospira mfg.)	Injection (Accord mfg.)	Injection (Sandoz mfg.)	
	□ J9322 Pemetrexed Injection (Bluepoint mfg.)		□ J9323 Pemetrexed Ditromethamine		
		_	Authorization R	-	
	J9305 Alimta®	□ J9292 Axtl	le TM	☐ J9304 Pemfexy® (Pemetrexed Injection)	
	(Pemetrexed Injection)	(Pemetrex	ed Injection)	(Pemetrexed Injection)	
	J9314 Pemetrexed Injection	on (Teva	□ J9324 Pemrydi RTU® (Pemetrexed		
	mfg.)		Ready-to-Use)		
M	EMBER & PRESCRIBER	INFORMATI	ON: Authorization	on may be delayed if incomplete.	
Mei	mber Name:				
Member Sentara #: Date of Birth:					
Pre	scriber Name:			_	
Prescriber Signature:			Date:		
Off	ice Contact Name:				
Phone Number:			Fax Nui	mber:	
NDI	ſ # .				

			thorization may be delayed if incomplete.
Dosii	ng So	chedule:	Length of Therapy:
Diag	nosis	:	ICD Code, if applicable:
Weig	ht (i	f applicable):	Date weight obtained:
			is box, the timeframe does not jeopardize the life or health of the member maximum function and would not subject the member to severe pain.
supp	ort e		ck below all that apply. All criteria must be met for approval. To nentation, including lab results, diagnostics, and/or chart notes, must be
Init	tial 1	Authorization: 6 mont	hs
	Th	e requesting provider is an	oncologist
		AND	
	htt the	tps://www.sentarahealthpl e detailed policy description	ocumented above; please access the following webpage, lans.com/en/providers/clinical-reference/medical-policies/medical, for on (Chemotherapy and Supportive Care Medical 316)) ance with a specific indication
	Ac	ccepted off-label indication	n found in the most recent edition of any of the following:
		American Hospital Formu	lary Service Drug Information (Supportive)
		-	Cancer Network's Drugs & Biologics Compendium (use must be ecommendations carrying a Category 1 or 2A level of evidence)
			Clinical Pharmacology (Supportive)
		-	rugDex® (Class I, IIa, or IIb)
		Wolters Kluwer Lexi-Drug	gs® (Level A)
		OR	
		you feel would be pertine	ease provide clinical rationale and submit any chart notes/literature ent in support of medical necessity. Note: experimental/investigationa emotherapy 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 has a documented intolerance, FDA-labeled contraindication, or hypersensitivity to first line therapies]

AND

□ Provider must submit documentation of a contraindication, failure, or intolerance to any of the preferred agents prior to approval of a non-preferred product (NOTE: Step therapy applies to all overlapping compendia supported indications/regimens)

AND

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

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

AND

	Re	quested 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
		requesting the brand formulation of any therapy with generic availability, provider must submit cumentation to confirm treatment failure, contraindication or intolerance to the generic product
suppo	ort e	orization: 6 months. Check below all that apply. All criteria must be met for approval. To ach line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be or request may be denied.
	Me	ember is currently receiving the requested medication and must meet BOTH of the following: All initial authorization criteria continues to be met
		Provider must submit documentation of contraindication, failure, or intolerance to any of the preferred agents prior to continued approval of the non-preferred products (NOTE: please see initial authorization section; step therapy applies to all overlapping compendia supported indications/regimens)
		AND
	Me	ember requires continuation of therapy and is NOT experiencing disease progression
		AND
	On	going treatment is consistent with FDA-labeling or compendia support
		AND
	Me	ember is NOT experiencing an FDA-labeled limitation of use or toxicity
		AND

PA Pemetrexed Injections (Medical)(CORE)

(Continued from previous page)

□ T	he quantity (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 **
Medic	ation being provided by: Please check applicable box below.
□ Loc	cation/site of drug administration:
NP	I or DEA # of administering location:
	<u>OR</u>
□ Spe	ecialty Pharmacy
standard urgent is	nt reviews: Practitioner should call Sentara Health Plans Pre-Authorization Department if they believe a review would subject the member to adverse health consequences. Sentara Health Plan's definition of a lack of treatment that could seriously jeopardize the life or health of the member or the member's regain maximum function.

**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. *