★actna* MEDICARE FORM Pegfilgrastim Precertification Request (Fylnetra, Fulphila*, Neulasta*, Neulasta Onpro*, Nyvepria*, Rolvedon™, Stimufend*, Udenyca*, Ziextenzo*) Page 1 of 4 (All fields must be completed and legible for precertification review.)				For Michigan MMP: FAX: 1-844-241-2495 PHONE: 1-855-676-5772 For other lines of business: Please use other form. Note: Fylnetra, Nyvepria, Rolvedon, Stimufend, Udenyca and Ziextenzo are non-preferred.		
Please indicate: Start of treatr			, ,			nd Neulasta/Neulasta preferred.
Precertification Requested By: _		last treatment			Fax:	
A. PATIENT INFORMATION			I none		1 ax.	
First Name:		Last Name:			DOB:	
		Last Maine.	Qit			710.
Address:			City:		State:	ZIP:
Home Phone:	Work Phone:		Cell Phone:		Email:	
Patient Current Weight: lbs_c	orkgs Pati	ient Height: inch	es or <u>cms</u> Al	llergies:		
B. INSURANCE INFORMATION		-		<i>,</i>		
Aetna Member ID #: Group #:			ner coverage? [] א Car			
Insured:		Insured:	Car			
	rovido ID #:		edicaid: 🗌 Yes 🗌 No		ida ID #i	
				o il yes, prov	nde ID #:	
C. PRESCRIBER INFORMATION First Name:		Last Name:		(Chock one):		D.O. 🗌 N.P. 🗌 P.A.
		Last Name.		(Check one).		
Address:		o	City:		State:	ZIP:
Phone: Fax:		St Lic #:	NPI #:	DEA #:		UPIN:
Provider Email:		Office Contact Name:			Phone:	
Specialty (Check one): Oncolog	gist 🗌 Hematologi	st 🗌 Other:				
Place of Administration: Self-administered Ph Home Infusion Center Center Name Phone: Outpatient Facility: Facility Name: Phone: Outpatient Facility: Facility Name: Phone: Outpatient Infusion Center: Center Addministration code(s) (CPT):			Address: City: Phone:	ffice rmacy	Retail Pha Home Care State:	rmacy e ZIP:
Phone:						
TIN:			NPI:			
NPI:						
E. PRODUCT INFORMATION Fylnetra (pegfilgrastim- pbbk) Fulphila (pegfilgrastim- jmdb)			irections for Use: irections for Use:			
□ Neulasta/Neulasta Onpro (pegfilg			irections for Use:			
☐ Nyvepria (pegfilgrastim-apgf)			irections for Use:			
🗌 Rolvedon (eflapegrastim-xnst)			irections for Use:			
Stimufend (pegfilgrastim-fpgk)	Dose:	D	irections for Use:			
Udenyca (pegfilgrastim-cbqv)			irections for Use:			
Ziextenzo (pegfilgrastim-bmez)		· · · · · ·	irections for Use:			
F. DIAGNOSIS INFORMATION - P	lease indicate prima	•		licable.		
Primary Indication:)] Other:		:C: 1:	
G. CLINICAL INFORMATION - R	-	ormation must be con	npleted in its <u>entirety</u> f	or all precert	mcation requ	ests.
For All requests (clinical documentation required): Please indicate the patient's absolute neutrophil count:mm³ Date obtained:/ / □ Yes □ No Does the patient have a nadir count that requires an immediate need for Fylnetra, Fulphila, Neulasta/Neulasta Onpro, Nyvepria, Rolvedon, Stimufend, Udenyca, or Ziextenzo? □ Yes □ No Will Fylnetra, Fulphila, Neulasta/Neulasta Onpro, Nyvepria, Rolvedon, Stimufend, Udenyca, or Ziextenzo be used with another colony stimulating factor? □ Yes □ No Is Fylnetra, Fulphila, Neulasta/Neulasta Onpro, Nyvepria, Rolvedon, Stimufend, Udenyca, or Ziextenzo part of a stem cell mobilization protocol?						



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB
G. CLINICAL INFORMATION (continued) –	Required alinical information must be com	nloted in its optiraty for all p	reporting requests
For All requests (clinical documentation req	•	pieted in its <u>entirety</u> for all pi	recentification requests.
Yes No Will Fylnetra, Fulphila, Neulas chemotherapy regimens?	,	timufend, Udenyca, or Ziext	enzo be given with weekly
Yes No Will Fylnetra, Fulphila, Neulas chemotherapy cycle as another		timufend, Udenyca, or Ziext	enzo be used in the same
Yes No Is the patient currently receivir		n therapy?	
For Initiation requests:			
Yes No Has the patient had a trial and	apy with Fylnetra (pegfilgrastim-pbbk), Nyv (pegfilgrastim-cbqv), or Ziextenzo (pegfilg	epria (pegfilgrastim-apgf), R rastim-bmez) within the last any of the following? (select	Rolvedon (eflapegrastim-xnst), Stimufend 365 days?
Please explain if there are any other medical re	eason(s) that the patient cannot use any o	f the following preferred pro	ducts (select all that apply)
🗌 Fulphila (pegfilgrastim-jmdb)	astim)	
☐ Acute lymphoblastic leukemia (ALL) ☐ Yes No Has the first days of cher ☐ Yes No Is this the initial induction ☐ Yes No Is this the first post-remis	of chemotherapy? sion course of chemotherapy?		
Please provide the chemotherapy regime	n and date started: Regimen:		Date started: / /
Please indicate the myelosuppressive and	ti-retroviral medication the patient is receiv	ing:	
Yes No Is the patient neutropenio	?	•	
Yes No Is the medication being r			ed infectious complications?
□ Yes □ No Is the patient undergoing	myeloablative chemotherapy? eatment will be followed by: Autologou	s bone marrow transplantat	ion
		bone marrow transplantation	
Congenital, cyclic or idiopathic neutrope Please identify which documented type of neut ☐ Yes ☐ No Is the patient currently sy	ropenia that patient has: 🗌 congenital ne mptomatic?		
requested for chronic ad oropharyngeal ulcers)?	-pbbk), Fulphila (pegfilgrastim-jmdb), Neu -xnst), Stimufend (pegfilgrastim-fpgk),Ude ministration to reduce the incidence and d	nyca (pegfilgrastim-cbqv), o	r Ziextenzo (pegfilgrastim-bmez)being
Chronic Myeloid Leukemia	sistant neutropenia?		
Yes No Is the neutropenia secon	dary to use of any of the following medica		
□ Drug- induced agranulocytosis	Gleevec (imatinib) Clusig (ponati	nib) 🗌 Sprycel (dasatinib)	🗌 Tasigna (nilotinib)
Yes No Is the agranulocytosis ca	used by chemotherapy?		
Please provide the me	dication(s) that caused the agranulocytos	s:	
☐ Glycogen storage disease (GSD) type 1 ☐ Yes ☐ No Does the patient have a l	ow neutrophil count?		
Hairy Cell Leukemia			
Yes No Does the patient have cli	nical evidence of neutropenic fever followi	ng chemotherapy?	
□ Increase dose intensity chemotherapy re			
disease control?	ed in a setting in which clinical research de		sive therapy produces improvement in
	be of cancer the patient is being treated fo		
Please enter the exact	t chemotherapy regimen patient is current	y being treated with:	

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G. CLINICAL INFORMATION (continued) –			recertification requests.		
What is the expected percentage of febrile neu $\Box 0.9\%$ (Low risk)] 10-19% (Intermediate risk)				
☐ Yes ☐ No Is the patient considered			ious complications?		
		orizes the patient to be at high risk:			
		years 🔲 Bone marrow compromis			
			ent neutropenia 🔲 Poor nutritional status		
	status 🔲 Previous chemotherapy	Previous radiation therapy	Previous episodes of FN		
Cther earlieus on m			function Donal dysfunction		
		se	•		
☐ Intermittent use in patients with myelody					
Yes No Does the patient have sy	mptomatic anemia?				
Yes No Has the patient been test					
☐ Yes ☐ No Does the patient present	sult of the test and date obtained:		_ Date obtained: / _/		
Yes No Has a serum erythropoiet	, .	55 !			
			_ Date obtained:/ /		
Yes No Is there clinical evidence			CHOP) rituximab, cyclophosphamide,		
	orednisone) or more aggressive reg				
Please indicate the part	tient's chemotherapy regimen:				
Primary prophylaxis of neutropenia					
Yes No Does the patient have a c		oid malignancy?			
Yes No Is the patient receiving m		atad far			
Please indicate the typ		currently being treated with:			
What is the expected percentage of febrile neu					
] 10-19% (Intermediate risk)				
Yes No Is the patient considered			ious complications?		
Please indicate which of the following reasons that categorizes the patient to be at high risk:					
		years 🔲 Bone marrow compromis			
			ent neutropenia		
□ Poor performance s	status U Previous chemotherapy	Previous radiation therapy	Previous episodes of FN		
	orbidities: 🔲 Cardiovascular dise	ase 🔲 HIV infection 🔲 Liver dys	stunction Renal dysfunction		
Radiation therapy alone					
Yes No Are prolonged delays in r	adiation therapy expected due to r	eutropenia?			
Secondary prophylaxis of neutropenia					
Yes No Does the patient have a c	documented diagnosis of non-myel	oid malignancy?			
Yes No Did the patient experienc					
		experienced from the prior cycle of	chemotherapy:		
Neutropenic complicat					
Please indicate the prive Please indicate the please indicate		patient received with the neutropen	•		
	ior cycle of similar chemotherapy?	t (a fracil of day of treatment count	inipacing the plained dose of		
☐ Yes ☐ No Was the patient treated w		lanned for current cycle?			
Yes No Did the patient receive pr					

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Patient First Name		Patient Last Name	Patient Phone	Patient DOB	Patient DOB	
G. CLINICAL INFORMA	TION (continued) –	Required clinical information mu	st be completed in its <u>entirety</u> for all pre	certification requests.		
☐ Therapeutic use in a	a high-risk, febrile ne	utropenic patient				
	•	gnostic factors pertains to the pa	atient:			
	Age greater than 6	,				
		at the time of the development of				
		ide date of hospitalization:				
	Invasive fungal infe		ection occurred:	Deter	1 1	,
	Provide type	or lungar intection and date inte			/	
		ide date of pneumonia infection:				
	Prior episodes of fe					
	Prolonged neutrope	enia				
	·	1 0 1	a expected to last greater than 10 days?	?		
	Profound neutrope	nia				
	☐ Sepsis syndrome ☐ Other					
		ain:				
Treatment for radiat		ann				
		used the injury: grays (G	y)			
For Continuation reque	<u>sts:</u>					
Neulasta (pegfilgr (Samplir	a/Neulasta Onpro (pec astim-cbqv), or Ziexte	ŋfilgrastim),Ŋyvepria (pegfilgrast nzo (pegfilgrastim-bmez)? a, Neulasta/Neulasta Onpro, Ny∖	nples of Fylnetra (pegfilgrastim-pbbk), F im-apgf), Rolvedon (eflapegrastim-xnst) /epria, Rolvedon, Stimufend, Udenyca, o	, Stimufend (pegfilgra	stim-fpgk), Uo	-
(pegfilgr		lgrastim-apgf), Rolvedon (eflape	obbk) Fulphila (pegfilgrastim-jmdb), Neu grastim-xnst), Stimufend (pegfilgrastim-			/), or
H. ACKNOWLEDGEM	ENT					
Request Completed B	y (Signature Requi	red):		Date:	/ /	
			f a medical procedure or service with	the intent to injure	defraud or d	leceive
			eals material information for the purpo			

L The plan may request additional information or clarification, if needed, to evaluate requests.

insurance act, which is a crime and subjects such person to criminal and civil penalties.