★ action ** 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.) Please indicate: Start of treatment: Start date // / Continuation of therapy: Date of last treatment // /				For New Jersey HMO D-SNP: FAX: 1-833-322-0034 PHONE: 1-844-362-0934 For other lines of business: Please use other form. Note: Fylnetra, Nyvepria, Rolvedon, Stimufend, Udenyca and Ziextenzo are non-preferred. Fulphila and Neulasta/Neulasta Onpro are preferred.			
Precertification Requested By: _				Phone	:	Fa	K:
A. PATIENT INFORMATION							
First Name:		Last Name:				DOB:	
Address:			С	ity:		State:	ZIP:
Home Phone:	Work Phone:		С	ell Phone:		Email:	
Patient Current Weight: lbs_c	or kas Pat	ient Height <sup>.</sup>			Allergies:		
B. INSURANCE INFORMATION		ione noight	moneo		7 alorgios.		
Aetna Member ID #:		Does patient hav	e other	coverage?	🗆 Yes 🗖 No		
Group #:		If yes, provide ID					
Insured:		Insured:					
Medicare: Yes No If yes, p	vrovide ID #:		Modi		☐ No If yes, pro	wide ID #:	
C. PRESCRIBER INFORMATION			Wear				
First Name:		Last Name:			(Chack ana)		D.O. 🗌 N.P. 🗌 P.A.
		Last Maille.		0.1	(Check one)		
Address:		1		City:		State:	ZIP:
Phone: Fax:		St Lic #:		NPI #:	DEA #:		UPIN:
Provider Email:		Office Contact Na	ame:			Phone:	
Specialty (Check one): Oncolog	jist 🗌 Hematolog	ist 🗌 Other:					
Place of Administration:         Self-administered       Ph         Home Infusion Center         Center Name         Phone:       Phone:         Outpatient Facility: Facility Name:         Phone:       Phone:         Outpatient Infusion Center: Center         Administration code(s) (CPT):       Address:         City:       Phone:         TIN:       NPI:         E. PRODUCT INFORMATION         Fylnetra (pegfilgrastim- pbbk)	er Name: State: Fax: PIN:	ZIP:		Physician Physician Specialty Other: Name: Address: City: Phone: TIN: NPI:		Retail Pr     Home Ca     State:     Fax:     PIN:	narmacy are ZIP:
☐ Fulphila (pegfilgrastim-jmdb)			_				
🗌 Neulasta/Neulasta Onpro (pegfilg			Dire	ections for Use:			
🗌 Nyvepria (pegfilgrastim-apgf)	Dose:		Dire	rections for Use:			
Rolvedon (eflapegrastim-xnst)				Directions for Use:			
Stimufend (pegfilgrastim-fpgk)							
Udenyca (pegfilgrastim-cbqv)							
Ziextenzo (pegfilgrastim-bmez)			_	ctions for Use:			
F. DIAGNOSIS INFORMATION - P	lease indicate prima	ary ICD code and s	specify	any other where	e applicable.		
Primary Indication:				Other:			
G. CLINICAL INFORMATION - F For All requests (clinical documenta Please indicate the patient's absolute □ Yes □ No Does the patient have Stimufend, Udenyca, □ Yes □ No Will Fylnetra, Fulphila stimulating factor?	ation required): neutrophil count: e a nadir count that re or Ziextenzo?	mm <sup>3</sup> Date obtai equires an immedia	ned: te need	/ / I for Fylnetra, Ful	_ phila, Neulasta/Ne	eulasta Onpro	Nyvepria, Rolvedon,

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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Rolvedon<sup>™</sup>, Stimufend<sup>®</sup>, Udenyca<sup>®</sup>, Ziextenzo<sup>®</sup>)

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB
G. CLINICAL INFORMATION (continued) –	Boguirod clinical information must be comp	lated in its antiraty for all properti	fication requests
G. CLINICAL INFORMATION (continued) – For All requests (clinical documentation req		neled in its <u>entirely</u> for all precerti	lication requests.
Yes No Will Fylnetra, Fulphila, Neulas	/	mufend, Udenyca, or Ziextenzo I	be given with weekly
chemotherapy regimens?		mufend, Udenyca, or Ziextenzo I	be used in the same
chemotherapy cycle as another Yes No Is the patient currently receiving		n therapy?	
For Initiation requests:			
☐ Yes ☐ No Has the patient had a trial and	apy with Fylnetra (pegfilgrastim-pbbk), Nyve (pegfilgrastim-cbqv), or Ziextenzo (pegfilgr failure, intolerance, or contraindication to a )	epria (pegfilgrastim-apgf), Rolved astim-bmez) within the last 365 d any of the following? (select all the stim) the following preferred products	on (eflapegrastim-xnst), Stimufend ays? at apply)
		5411)	
<ul> <li>☐ Acute lymphoblastic leukemia (ALL)</li> <li>☐ Yes ☐ No Has the first days of cher</li> <li>☐ Yes ☐ No Is this the initial induction</li> </ul>	of chemotherapy?		
Yes No Is this the first post-remis	sion course of chemotherapy? n and date started: Regimen:		Date started: / /
Advanced HIV infection			
Please indicate the myelosuppressive and Please I No Is the patient neutropenion of t	ti-retroviral medication the patient is receivi c?	ng:	
Bone Marrow Transplantation			
	documented diagnosis of non-myeloid mali		stieves executions?
Yes No is the medication being r	equested to reduce the duration of neutrop mveloablative chemotherapy?	enia and neutropenia-related infe	clious complications?
	eatment will be followed by:  Autologous	bone marrow transplantation	
	☐ Allogeneic ☐ None	bone marrow transplantation	
Congenital, cyclic or idiopathic neutrope		tana anin 🗖 avalin anytana anin	
Please identify which documented type of neut			
Yes No Is Fylnetra (pegfilgrastim Rolvedon (eflapegrastim	-pbbk), Fulphila (pegfilgrastim-jmdb), Neula -xnst), Stimufend (pegfilgrastim-fpgk),Uden ministration to reduce the incidence and du	yca (pegfilgrastim-cbqv), or Ziext	enzo (pegfilgrastim-bmez)being
Chronic Myeloid Leukemia			
Yes No Does the patient have real	sistant neutropenia? dary to use of any of the following medicati	one?	
	Gleevec (imatinib) Clusig (ponatin		asigna (nilotinib)
Drug- induced agranulocytosis		, _ , , , _ , _	<b>č</b>
☐ Yes ☐ No Is the agranulocytosis ca			
Glycogen storage disease (GSD) type 1	dication(s) that caused the agranulocytosis		
Yes No Does the patient have a l	ow neutrophil count?		
Hairy Cell Leukemia			
	nical evidence of neutropenic fever followin	g chemotherapy?	
□ Increase dose intensity chemotherapy re □ Yes □ No Is the patient being treated disease control?	ed in a setting in which clinical research der	nonstrates that dose-intensive th	erapy produces improvement in
Please indicate the typ	be of cancer the patient is being treated for:		
Please enter the exact	chemotherapy regimen patient is currently	being treated with:	

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB
G. CLINICAL INFORMATION (continued) – I	Required clinical information must be comp	leted in its entirety for all precertif	ication requests.
What is the expected percentage of febrile neu	tropenia incidence from the chemotherapy ] 10-19% (Intermediate risk)  □ 20% or gi	regimen? reater  (high risk)	
	to be at high risk for chemotherapy-induced of the following reasons that categorizes th		implications?
Active infections	Age greater than or equal to 65 years	Bone marrow compromise	
	/ement by tumor producing cytopenias		
Poor performance s	status 🔲 Previous chemotherapy 🗌 Pre	evious radiation therapy [] Previ	ous episodes of FN
	orbidities: Cardiovascular disease	HIV infection 🔲 Liver dysfunctio	
Intermittent use in patients with myelody			
☐ Yes ☐ No Does the patient have sy			
Yes No Has the patient been test		_	
	sult of the test and date obtained:	Da	te obtained: / /
☐ Yes ☐ No Does the patient present ☐ Yes ☐ No Has a serum erythropoiet			
	sult of the test and date obtained:	Da	te obtained: / /
	that the patient is being treated with curativ orednisone) or more aggressive regimens?	ve chemotherapy (e.g. (R- CHOP	) rituximab, cyclophosphamide,
$\rightarrow$ Please indicate the particular	tient's chemotherapy regimen:		
Primary prophylaxis of neutropenia			
	documented diagnosis of non-myeloid malig	gnancy?	
Yes No Is the patient receiving m			
	e of cancer the patient is being treated for: chemotherapy regimen patient is currently		
What is the expected percentage of febrile neu			
	] 10-19% (Intermediate risk)	-	
	to be at high risk for chemotherapy-induced		mplications?
	of the following reasons that categorizes th		
	☐ Age greater than or equal to 65 years [		
	vement by tumor producing cytopenias		
	status 🔲 Previous chemotherapy 🗌 Pre	evious radiation therapy D Previ	ous episodes of FN
☐ Recent surgery ☐ Other serious co-m	orbidities: Cardiovascular disease	HIV infection 🔲 Liver dysfunctio	
☐ Radiation therapy alone			
— 13	adiation therapy expected due to neutrope	nia?	
Secondary prophylaxis of neutropenia			
	documented diagnosis of non-myeloid malig	gnancy?	
Yes No Did the patient experienc	e a febrile neutropenic complication from a	prior cycle of chemotherapy?	
	utropenic complication the patient experien	ced from the prior cycle of chemo	otherapy:
Neutropenic complicat			
	or cycle of chemotherapy that the patient re		
	e a dose-limiting neutropenic event (a nadii ior cycle of similar chemotherapy?	i or day or treatment count impac	ung me planned dose of
	<i>i</i> th the same dose and schedule planned for	or current cycle?	
	imary prophylaxis against febrile neutropen		

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Patient First Name	ame Patient Last Name Patient Phone		Patient DOB	Patient DOB		
G. CLINICAL INFOR	MATION (continued) –	Required clinical information must	be completed in its entirety for all pre-	ecertification request	S.	
☐ Therapeutic use	in a high-risk, febrile ne	utropenic patient				
	•	gnostic factors pertains to the pati	ent:			
	Age greater than 68					
	<b>.</b> .	at the time of the development of				
		ide date of hospitalization: /	/			
	Invasive fungal infe		tion a command.	Dete	,	1
	Provide type	e of fungal infection and date infec	tion occurred:	Date:	/	1
		ide date of pneumonia infection:	1 1			
	Prior episodes of fe		, <u>, </u>			
	Prolonged neutrope	•				
	$\longrightarrow$ Yes $\square$ M	No Is the prolonged neutropenia	expected to last greater than 10 days	?		
	Profound neutroper	nia				
	Sepsis syndrome					
		ain:				
Treatment for rac						
		used the injury: grays (Gy)				
For Continuation rec		, , <u> </u>				
Neul (pegi (Sam	asta/Neulasta Onpro (peg filgrastim-cbqv), or Ziexter	yfilgrastim),Nyvepria (pegfilgrastim nzo (pegfilgrastim-bmez)? a, Neulasta/Neulasta Onpro, Nyve	oles of Fylnetra (pegfilgrastim-pbbk), I-apgf), Rolvedon (eflapegrastim-xnst pria, Rolvedon, Stimufend, Udenyca,	), Stimufend (pegfilg	rastim-fpgk), l	-
(pegt		ilgrastim-apgf), Rolvedon (eflapeg	bk) Fulphila (pegfilgrastim-jmdb), Net rastim-xnst), Stimufend (pegfilgrastim			qv), or
H. ACKNOWLEDG	EMENT					
Request Complete	d By (Signature Requi	red):		Date:	1	1
			a medical procedure or service with	h the intent to injur	e defraud or	deceive
			Is material information for the purp			

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.