

# **Pegfilgrastim Precertification Request**

(Fylnetra, Fulphila®, Neulasta®, Neulasta Onpro®, Nyvepria®, Rolvedon<sup>™</sup>, Stimufend<sup>®</sup>, Udenyca<sup>®</sup>, Ziextenzo<sup>®</sup>)

Page 1 of 4

(All fields must be completed and legible for precertification review.)

Please indicate: ☐ Start of treatment: Start date \_\_\_\_/

For Ohio MMP:

**FAX:** 1-855-734-9389 PHONE: 1-855-364-0974

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.

	Date of last treatment			•	•
Precertification Requested By:		Phone:		Fax	<b>«</b> :
A. PATIENT INFORMATION					
First Name:	Last Name:			DOB:	
Address:	•	City:		State:	ZIP:
Home Phone: Work	Phone:	Cell Phone:		Email:	
Patient Current Weight: lbs or k			Allergies:		
B. INSURANCE INFORMATION	go r daoint rioight.	mones ofome	morgios.		
Aetna Member ID #:	Does patient hav	e other coverage?	Yes No		
Group #:		If yes, provide ID#: Carrier Name:			
Insured:	Insured:				
Medicare: ☐ Yes ☐ No If yes, provide ID #	:	Medicaid: Yes 1	No If yes, pro	vide ID #:	
C. PRESCRIBER INFORMATION					
First Name:	Last Name:		(Check one):	M.D.	D.O.  N.P.  P.A.
Address:	<u>.</u>	City:		State:	ZIP:
Phone: Fax:	St Lic #:	NPI #:	DEA #:		UPIN:
Provider Email:	Office Contact Na	ame.	I	Phone:	
Specialty (Check one): Oncologist He				1	
D. DISPENSING PROVIDER/ADMINISTRATI					
Center Name	ZIP:	Address:  City: Phone: TIN:		_ State: Fax: _ PIN: _	ZIP:
<b>TIN:</b> PIN:		NPI:			
NPI:					
E. PRODUCT INFORMATION  ☐ Fylnetra (pegfilgrastim- pbbk)	Dose:	Directions for Use:			
☐ Fulphila (pegfilgrastim- jmdb)	Dose:				
☐ Neulasta/Neulasta Onpro (pegfilgrastim)	Dose:				
☐ Nyvepria (pegfilgrastim-apgf)	Dose:	Directions for Use:			
☐ Rolvedon (eflapegrastim-xnst)	Dose:				
Stimufend (pegfilgrastim-fpgk)	Dose:				
Udenyca (pegfilgrastim-cbqv)	Dose:				
☐ Ziextenzo (pegfilgrastim-bmez)	Dose:				
F. DIAGNOSIS INFORMATION - Please indic	ate primary ICD code and s		рисаріе.		
Primary Indication:	inical information movet be	Other:	for all rayonari	lifi a a ti a mana	w.c.c.tc
G. CLINICAL INFORMATION - Required cl  For All requests (clinical documentation required Please indicate the patient's absolute neutrophil of the patient have a nadir constitution of the patient have a	ed): count:mm³ Date obtai unt that requires an immedia o? Neulasta Onpro, Nyvepria, R	ned: //// te need for Fylnetra, Fulphil olvedon, Stimufend, Udenyo	a, Neulasta/Ne ca, or Ziextenzo	ulasta Onpro,	Nyvepria, Rolvedon, n another colony



# **Pegfilgrastim Precertification Request**

(Fylnetra, Fulphila®, Neulasta®, Neulasta Onpro®, Nyvepria®, Rolvedon™, Stimufend®, Udenyca®, Ziextenzo®)

Page 2 of 4

(All fields must be completed and legible for precertification review.)

For Ohio MMP:

**FAX:** 1-855-734-9389 **PHONE:** 1-855-364-0974

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.

Patient First Name	Patient Last Name	Patient Phone	Patient DOB
C. CLINICAL INFORMATION (com	figured. Dequired clinical information my	at he completed in its entirety for all	propertification requests
For All requests (clinical documen	tinued) - Required clinical information mu	st be completed in its <u>entirety</u> for all	precertification requests.
☐ Yes ☐ No Will Fylnetra, Fulphi	a, Neulasta/Neulasta Onpro, Nyvepria, R	olvedon, Stimufend, Udenyca, or Zie	xtenzo be given with weekly
	a, Neulasta/Neulasta Onpro, Nyvepria, Ra as another colony stimulating factor?	olvedon, Stimufend, Udenyca, or Zie	xtenzo be used in the same
	tly receiving concomitant chemotherapy a	nd radiation therapy?	
For Initiation requests:			
☐ Yes ☐ No Has the patient had (pegfilgrastim-fpgk),☐ Yes ☐ No Has the patient had☐ Fulphila (pegfilgra	Udenyca (pegfilgrastim-cbqv), or Ziexten a trial and failure, intolerance, or contrain	obbk), Nyvepria (pegfilgrastim-apgf), zo (pegfilgrastim-bmez) within the last dication to any of the following? (sele to (pegfilgrastim)	Rolvedon (eflapegrastim-xnst), Stimufend st 365 days? ect all that apply)
☐ Fulphila (pegfilgra	stim-jmdb) Neulasta/Neulasta Onpre	o (pegfilgrastim)	
☐ Yes ☐ No Is this the initia ☐ Yes ☐ No Is this the first Please provide the chemothera ☐ Advanced HIV infection	ys of chemotherapy been completed? I induction of chemotherapy? cost-remission course of chemotherapy? py regimen and date started: Regimen: essive anti-retroviral medication the patie		
	eutropenic?		
☐ Yes ☐ No Is the medicati		of neutropenia and neutropenia-rela	ation
Congenital, cyclic or idiopathic			
☐ Yes ☐ No Is the patient c☐ Yes ☐ No Is Fylnetra (per Rolvedon (efla	gfilgrastim-pbbk), Fulphila (pegfilgrastim-jr pegrastim-xnst), Stimufend (pegfilgrastim- hronic administration to reduce the incide	ndb), Neulasta/Neulasta Onpro (peg fpgk),Udenyca (pegfilgrastim-cbqv),	filgrastim), Nyvepria (pegfilgrastim-apgf), or Ziextenzo (pegfilgrastim-bmez)being
☐ Chronic Myeloid Leukemia	,		
☐ Drug- induced agranulocytosis	nia secondary to use of any of the followinosutinib)   Gleevec (imatinib)   Iclust		o) 🔲 Tasigna (nilotinib)
	de the medication(s) that caused the agra	nulocytosis:	
☐ Glycogen storage disease (GSI☐ Yes ☐ No Does the patie	D) type 1	,	
☐ Hairy Cell Leukemia			
	nt have clinical evidence of neutropenic fe	ver following chemotherapy?	
☐ Increase dose intensity chemo ☐ Yes ☐ No Is the patient b disease contro	eing treated in a setting in which clinical re	esearch demonstrates that dose-inte	nsive therapy produces improvement in
> Please indic	ate the type of cancer the patient is being		
Please ente	the exact chemotherapy regimen patient	is currently being treated with:	

Continued on next page



# **Pegfilgrastim Precertification Request**

(Fylnetra, Fulphila®, Neulasta®, Neulasta Onpro®, Nyvepria®, Rolvedon™, Stimufend®, Udenyca®, Ziextenzo®)

Page 3 of 4

(All fields must be completed and legible for precertification review.)

For Ohio MMP:

**FAX:** 1-855-734-9389 **PHONE:** 1-855-364-0974

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.

Patient First Name		Patient Last Name	Patient Phone	Patient DOB
G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its entirety for all precertification requests.				
		ropenia incidence from the chemothera		
		10-19% (Intermediate risk) 20% o		
		to be at high risk for chemotherapy-indu		emplications?
		of the following reasons that categorizes		
		Age greater than or equal to 65 years ement by tumor producing cytopenias		utropenia
		tatus Previous chemotherapy		
	Recent surgery	tatae : Tevieue enemetrierapy ::	Tovious radiation therapy	ious opisouse or i ii
_	_ ,	orbidities: Cardiovascular disease		
☐ Intermittent use in p	ationts with myolody			-
	es the patient have syn			
		ed for 5q gene deletion?		
$\vdash$	Please indicate the res	ult of the test and date obtained:	Da	te obtained: / /
		with other cytogenetic abnormalities?	_	
		in test been completed?		
	Please indicate the res	ult of the test and date obtained:	Da	te obtained://
Lymphoma			tive alcomothermany (a.m. (D. CHOF	) \ mit
		that the patient is being treated with currednisone) or more aggressive regimen		) rituximab, cyclopnospnamide,
		ient's chemotherapy regimen:		
	•			
☐ Primary prophylaxis	•	ocumented diagnosis of non-myeloid m	alignancy?	
		/elosuppressive chemotherapy?	angriancy:	
		e of cancer the patient is being treated t	or:	
		chemotherapy regimen patient is currer		
	•	ropenia incidence from the chemothera	•	
		10-19% (Intermediate risk) 20% o		
		to be at high risk for chemotherapy-indu		omplications?
		of the following reasons that categorizes		
☐ Active infections ☐ Age greater than or equal to 65 years ☐ Bone marrow compromise ☐ Bone marrow involvement by tumor producing cytopenias ☐ Open wounds ☐ Persistent neutropenia ☐ Poor nutritional status				
☐ Poor performance status ☐ Previous chemotherapy ☐ Previous radiation therapy ☐ Previous episodes of FN				
	☐ Recent surgery			
		orbidities:   Cardiovascular disease	☐ HIV infection ☐ Liver dysfunction	on 🔲 Renal dysfunction
		Other- Please explain:		
☐ Radiation therapy al				
∐ Yes ∐ No Are	e prolonged delays in ra	adiation therapy expected due to neutro	penia?	
☐ Secondary prophyla				
		ocumented diagnosis of non-myeloid m		
		e a febrile neutropenic complication fron		
Please indicate the neutropenic complication the patient experienced from the prior cycle of chemotherapy:				
Neutropenic complication: Please indicate the prior cycle of chemotherapy that the patient received with the neutropenic complication:				
	•	•	•	
Yes ☐ No Did the patient experience a dose-limiting neutropenic event (a nadir or day of treatment count impacting the planned dose of chemotherapy) from a prior cycle of similar chemotherapy?				5 - F
☐ Yes ☐ No Was the patient treated with the same dose and schedule planned for current cycle?				
☐ Yes ☐ No Did the patient receive primary prophylaxis against febrile neutropenia?				

Continued on next page



### **Pegfilgrastim Precertification Request**

(Fylnetra, Fulphila®, Neulasta®, Neulasta Onpro®, Nyvepria®, Rolvedon™, Stimufend®, Udenyca®, Ziextenzo®)

Page 4 of 4

(All fields must be completed and legible for precertification review.)

For Ohio MMP:

**FAX:** 1-855-734-9389 **PHONE:** 1-855-364-0974

For other lines of business:

Please use other form.

Onpro are preferred.

Note: Fylnetra, Nyvepria, Rolvedon, Stimufend, Udenyca and Ziextenzo are non-preferred. Fulphila and Neulasta/Neulasta

Patient First Na	ame	Patient Last Name	Patient Phone	Patient DOB
G. CLINICAL II	NFORMATION (continued) -	l Required clinical information must be cor	mpleted in its <u>entirety</u> for all pr	recertification requests.
☐ Therapeution	c use in a high-risk, febrile ne dicate which of the following pro Age greater than 69 Being hospitalized Please provice type Provide type Provide type Prior episodes of fee Profound neutrope Sepsis syndrome Other	utropenic patient gnostic factors pertains to the patient: 5 years at the time of the development of fever de date of hospitalization:/_/ ction of fungal infection and date infection oc de date of pneumonia infection:/ brile neutropenia enia No Is the prolonged neutropenia expect	curred:// ed to last greater than 10 day	Date://
☐ Treatment f	Please explation injury	ain:		
		used the injury: grays (Gy)		
For Continuation	<u> </u>			
∐ Yes ∐ No	Neulasta/Neulasta Onpro (peg (pegfilgrastim-cbqv), or Ziexte	nzo (pegfilgrastim-bmez)? ı, Neulasta/Neulasta Onpro, Nyvepria, R	Rolvedon (eflapegrastim-xns	Fulphila (pegfilgrastim-jmdb), t), Stimufend (pegfilgrastim-fpgk), Udenyca , or Ziextenzo) does not guarantee coverage
Yes No				eulasta/Neulasta Onpro n-fpgk), Udenyca (pegfilgrastim-cbqv), or
H. ACKNOWL	.EDGEMENT			
Request Com	pleted By (Signature Requi	red):		Date:/ /
any insurance	company by providing materi		erial information for the pur	th the intent to injure, defraud or deceive cose of misleading, commits a fraudulent

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