

# Filgrastim Precertification Request (Granix®, Leukine, Neupogen®, Nivestym®, Releuko®, Zarxio)

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(filgrastim-sndz) be used within 7 days of Neulasta (pegfilgrastim)?

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

For Illinois MMP: FAX: 1-855-320-8445 **PHONE**: 1-866-600-2139

For other lines of business: Please use other form.

Note: Granix, Leukine, Neupogen, Nivestym, and

Releuko are non-preferred.

Please indicate:   Stort of tree	tmont: Start data			,	Zarxio	is preferred.
Please indicate: Start of trea		last treatment /	1			
Precertification Requested By:				:	Fa	ax:
A. PATIENT INFORMATION						
First Name:		Last Name:			DOB:	
Address:		City:			State:	ZIP:
Home Phone:	Work Phone:		Cell Phone:		Email:	
Patient Current Weight: lbs		nt Height: inches	or cms	Allergies:	I	
B. INSURANCE INFORMATION	g-	<u></u>				
Aetna Member ID #:		Does patient have other	r coverage?	☐ Yes ☐ No		
Group #:		If yes, provide ID#:Carrier Name:				
Insured:		Insured:				
C. PRESCRIBER INFORMATION		Lead Maria		(0)		
First Name:		Last Name:		(Check one):		D.O. N.P. P.A.
Address:		City:		T	State:	ZIP:
Phone: Fax	:	St Lic #:	NPI #:	DEA #:		UPIN:
Provider Email:	Offi	ce Contact Name:		Phone:		
☐ Outpatient Infusion Center Center Name: ☐ Home Infusion Center Agency Name: ☐ Administration code(s) (CPT): Address:	ysician's Office [ Phone: Phone:	] Home	☐ Physician ☐ Specialty ☐ Other: _ Name: _ Address: _	Pharmacy	Retail P Mail Ord	•
E. PRODUCT INFORMATION		Discretions for the	-			
	se:					
<ul><li>☐ Leukine (sargramostim)</li><li>☐ Nivestym (filgrastim-aafi)</li><li>Do</li></ul>	se:					
	se:					
☐ Releuko (filgrastim-ayow) Do						
	se:					
F. DIAGNOSIS INFORMATION -	Please indicate prima		<u> </u>	applicable.		
Primary Indication:	•		Other:	''		
G. CLINICAL INFORMATION - R	eguired clinical inform			r all precertification	n requests	
For All requests (clinical documen	•	<u>'</u>	<del></del>	•	'	
Please indicate the patient's absolut  Yes No Does the patient ha Nivestym (filgrastim  Yes No Is the requested do:  Yes No H	e neutrophil count: ve a nadir count that re -aafi), Releuko (filgras se less than 180 mcg ( as the patient tried Zar ] Yes	mm³ Date obtained: _equires an immediate need tim-ayow), or Zarxio (filgra 0.3 mL)? xio (filgrastim-sndz)? e patient have a contraind s ☐ No Is the patient co	I for Granix (tbo-fi stim-sndz)? ication to Zarxio (1 mpleting an existii	filgrastim-sndz)? ng chemotherapy ro		
☐ Yes ☐ No Will Granix (tbo-filgr	ractim) Louking (acres		to remain unchang	,	ouko (filara:	etim avow) or Zarvia
(filgrastim-sndz) be	used with another colo Granix (tbo-filgrastim) n-sndz) part of a stem /ill Granix (tbo-filgrastin	ny stimulating factor?	Neupogen (filgras Nivestym (filgrasti	tim), Nivestym (filg	rastim-aafi)	, Releuko (filgrastim-ayow),
☐ Yes ☐ No Will Granix (tbo-filgr	astim), Leukine (sargraused in the same chem	amostim), Neupogen (filgra notherapy cycle as anothe	astim), Nivestym ( r colony stimulatir		euko (filgra:	stim-ayow) or Zarxio
☐ Yes ☐ No. Will Granix (tho-filer	, ,			ilarastim-aafi) Rele	uko (filaras	tim-avow) or Zarvio



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Please use other form.

Note: Granix, Leukine, Neupogen, Nivestym, and Releuko are non-preferred. Zarxio is preferred.

Patient First Name	Patient Last Name	Patient Phone	Patient DOB		
G. CLINICAL INFORMATION (continued) – F	Required clinical information must be compl	eted in its <u>entirety</u> for all precertif	ication requests.		
G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its entirety for all precertification requests.  For Initiation requests:  Note: Granix, Leukine, Neupogen, Nivestym, and Releuko are non-preferred. Zarxio is preferred.  Yes No Has the patient had prior therapy with Granix (tbo-filgrastim), Leukine (sargramostim), Neupogen (filgrastim), Nivestym (filgrastim-aafi), or Releuko (filgrastim-ayow) within the last 365 days?  Yes No Has the patient had a trial and failure, intolerance, or contraindication to Zarxio (filgrastim-sndz)?  Please explain if there are any other medical reason(s) that the patient cannot use Zarxio (filgrastim-sndz).					
Granix (tbo-filgrastim):  ☐ Yes ☐ No Does the patient have a solid to significant incidence of febrile.	umor or non-myeloid malignancy and will renewtropenia for primary or secondary proph		herapy associated with a clinically		
Leukine (sargramostim):  ☐ Acute myeloid leukemia ☐ Yes ☐ No Is the patient receiving incomplete indicate the regire ☐ Yes ☐ No Is the patient receiving co	duction chemotherapy? nen: nsolidation chemotherapy?	<b>,</b>			
→ Please indicate the regir  Adjunct to progenitor cell-transplantation Please indicate which type of transplant a	· ·				
<ul><li>☐ Advanced HIV infection</li><li>Please indicate the myelosuppressive anti</li><li>☐ Yes ☐ No Is the patient neutropenic</li></ul>	i-retroviral medication the patient is receivir ?	ng:			
☐ Yes ☐ No Is the medication being re☐ Yes ☐ No Is the patient undergoing	atment will be followed by: 🔲 Autologous b	enia and neutropenia-related infe	ctious complications?		
☐ Congenital, cyclic or idiopathic neutrope Please identify which documented type of ☐ Yes ☐ No Is the patient currently syn	neutropenia that patient has: $\square$ congenita	l neutropenia ☐ cyclic neutrope	nia ☐ idiopathic neutropenia		
	ication(s) that caused the agranulocytosis:				
<ul> <li>☐ Hematopoietic Subsyndrome of Acute Radiation Syndrome (H-ARS)</li> <li>☐ Yes ☐ No Is the medication being requested for the treatment of radiation-induced myelosuppression following a radiological/nuclear incident?</li> </ul>					
☐ Intermittent use in patients with myelody ☐ Yes ☐ No Does the patient have syr ☐ Yes ☐ No Has the patient been teste	rsplastic syndromes mptomatic anemia? ed for 5q gene deletion? It of the test and date obtained:	, ,,	ate obtained://		
☐ Yes ☐ No Has a serum erythropoiet  → Please indicate the resu ☐ Neuroblastoma	in test been completed? It of the test and date obtained:	Da	ate obtained://		
(Aldesleukin), (Proleukin), is	onsidered high-risk? ation be used in combination with ALL of the sotretinoin (13-cis-retinoic acid)? Vill the requested medication be used in col				

Continued on next page



## **Filgrastim Precertification Request** (Granix<sup>®</sup>, Leukine, Neupogen<sup>®</sup>, Nivestym<sup>®</sup>, Releuko®, Zarxio®)

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

For Illinois MMP: **FAX:** 1-855-320-8445 PHONE: 1-866-600-2139

For other lines of business:

Please use other form.

Note: Granix, Leukine, Neupogen, Nivestym, and Releuko are non-preferred.

Zarxio is 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. Primary prophylaxis of neutropenia ☐ Yes ☐ No Does the patient have a documented diagnosis of non-myeloid malignancy? ☐ Yes ☐ No Is the patient receiving myelosuppressive chemotherapy? Please indicate the type of cancer the patient is being treated for: Please enter the exact chemotherapy regimen patient is currently being treated with: What is the expected percentage of febrile neutropenia incidence from the chemotherapy regimen? □ 0-9% (Low risk) □ 10-19% (Intermediate risk) □ 20% or greater (high risk) ☐ Yes ☐ No Is the patient considered to be at high risk for chemotherapy-induced febrile neutropenia infectious complications? > Please indicate which of the following reasons that categorizes the patient to be at high risk: ☐ 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 ☐ Other serious co-morbidities: ☐ Cardiovascular disease ☐ HIV infection ☐ Liver dvsfunction ☐ Renal dvsfunction Other- Please explain: \_\_\_ ☐ Secondary prophylaxis of neutropenia Yes No Does the patient have a documented diagnosis of non-myeloid malignancy? ☐ Yes ☐ No Did the patient experience a febrile neutropenic complication from a prior cycle of chemotherapy? > 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? → ☐ 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? ☐ Therapeutic use in a high-risk, febrile neutropenic patient Please indicate which of the following prognostic factors pertains to the patient: ☐ Age greater than 65 years ☐ Being hospitalized at the time of the development of fever Please provide date of hospitalization: \_\_\_ / / ☐ Invasive fungal infection Provide type of fungal infection and date infection occurred: \_\_\_\_\_\_ Date: \_\_\_\_/ ☐ Pneumonia Please provide date of pneumonia infection: \_\_\_\_/ ☐ Prior episodes of febrile neutropenia ☐ Prolonged neutropenia → ☐ Yes ☐ No Is the prolonged neutropenia expected to last greater than 10 days? ☐ Profound neutropenia ☐ Sepsis syndrome ☐ Other → Please explain: Neupogen (filgrastim), Nivestym (filgrastim-aafi), Releuko (filgrastim-ayow), Zarxio (filgrastim-sndz): ☐ Acute lymphoblastic leukemia (ALL) ☐ Yes ☐ No Has the first days of chemotherapy been completed? ☐ Yes ☐ No Is this the initial induction of chemotherapy? ☐ Yes ☐ No Is this the first post-remission course of chemotherapy? Please provide the chemotherapy regimen and date started: Regimen: \_\_\_\_\_\_\_ Date started: \_\_\_ / \_\_\_\_ ☐ Acute myeloid leukemia ☐ Yes ☐ No Is the patient receiving induction chemotherapy? Please indicate the regimen:

☐ Yes ☐ No Is the patient receiving consolidation chemotherapy?

Continued on next page

Please indicate the regimen: \_\_\_\_

Yes No Is the patient receiving chemotherapy for relapsed or refractory disease?

Relapsed disease Refractory disease Please indicate the regimen: \_\_\_\_



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB		
G. CLINICAL INFORMATION (continued) –	Poguired clinical information must be comp	loted in its entirety for all proce	prification requests		
☐ Adjunct to progenitor cell-transplantation	<u> </u>		ertification requests.		
	and date received: Autologous Allog		/ /		
☐ Advanced HIV infection					
	ti-retroviral medication the patient is receivi	ng:			
Yes No Is the patient neutropenion	c?				
Bone Marrow Transplantation  ✓ Yes ✓ No. Does the nationt have a decimal between the partial transplantation.	decumented diagnosis of non myoloid mali	ananov?			
	<ul><li>☐ Yes</li><li>☐ No Does the patient have a documented diagnosis of non-myeloid malignancy?</li><li>☐ Yes</li><li>☐ No Is the medication being requested to reduce the duration of neutropenia and neutropenia-related infectious complications?</li></ul>				
☐ Yes ☐ No Is the patient undergoing	myeloablative chemotherapy?		·		
> Please identify if the tre	eatment will be followed by: Autologous				
	☐ Allogeneic b	one marrow transplantation			
☐ Congenital, cyclic or idiopathic neutrope					
Please identify which documented type of	f neutropenia that patient has: 🗌 congenita	al neutropenia 🔲 cyclic neutro	openia 🔲 idiopathic neutropenia		
Yes No Is the patient currently sy	/mptomatic? Leukine (sargramostim), Neupogen (filgras	stim) Nivostym (filarastim aafi)	Polouko (filgrastim avow)		
or Zarxio (filgrastim-sndz	<ul><li>being requested for chronic administration</li></ul>	n to reduce the incidence and	duration of sequelae of neutropenia		
(e.g., fever, infections, or					
☐ Chronic Myeloid Leukemia ☐ Yes ☐ No Does the patient have re-	aistant nautranonia?				
	dary to use of any of the following medicati	ons?			
	☐ Gleevec (imatinib) ☐ Iclusig (ponatinib		Tasigna (nilotinib)		
☐ Drug- induced agranulocytosis					
Yes No Is the agranulocytosis ca	used by chemotherapy? lication(s) that caused the agranulocytosis:				
☐ Glycogen storage disease (GSD) type 1	incation(s) that caused the agrandiocytosis.				
Yes No Does the patient have a low neutrophil count?					
☐ Hairy Cell Leukemia					
	nical evidence of neutropenic fever followin	g chemotherapy?			
☐ Increase dose intensity chemotherapy re	<b>egimens</b> ed in a setting in which clinical research der	monstrates that dose-intensive	therapy produces improvement in		
disease control?			merapy produces improvement in		
> Please indicate the type of cancer the patient is being treated for:  Please enter the exact chemotherapy regimen patient is currently being treated with:					
What is the expected percentage of febrile neutropenia incidence from the chemotherapy regimen?					
□ 0-9% (Low risk) □ 10-19% (Intermediate risk) □ 20% or greater (high risk)					
☐ Yes ☐ No Is the patient considered to be at high risk for chemotherapy-induced febrile neutropenia infectious complications?  → Please indicate which of the following reasons that categorizes the patient to be at high risk:					
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  H	IIV infaction D Liver dysfunct	ion		
Other serious co-mo	Other- Please explain:				
☐ Intermittent use in patients with myelody			_		
Yes No Does the patient have sy					
Yes No Has the patient been test  Please indicate the resu	ted for 5q gene deletion'? ult of the test and date obtained:		Date obtained: / /		
☐ Yes ☐ No Does the patient present	with other cytogenetic abnormalities?		<del></del>		
	Yes No Has a serum erythropoietin test been completed?  Please indicate the result of the test and date obtained:				
Lymphoma	uit of the test and date obtained:		Date obtained. / /		
☐ Yes ☐ No Is there clinical evidence	that the patient is being treated with curative		OP ) rituximab, cyclophosphamide,		
	orednisone) or more aggressive regimens?				
Please indicate the pati	ient's chemotherapy regimen:				



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB		
G. CLINICAL INFORMATION (continued) –	Paguired clinical information mud	t he completed in its entirety for all n	resertification requests		
☐ Primary prophylaxis of neutropenia	Required clinical information mus	it be completed in its <u>entirety</u> for all pr	recertification requests.		
Yes No Does the patient have a	documented diagnosis of non-my	eloid malignancy?			
☐ Yes ☐ No Is the patient receiving m	yelosuppressive chemotherapy?				
> Please indicate the type					
		currently being treated with:			
What is the expected percentage of febril  □ 0-9% (Low risk) □					
	☐ 0-9% (Low risk) ☐ 10-19% (Intermediate risk) ☐ 20% or greater (high risk) ☐ Yes ☐ No Is the patient considered to be at high risk for chemotherapy-induced febrile neutropenia infectious complications?				
Please indicate which o	of the following reasons that categ	orizes the patient to be at high risk:			
		years Bone marrow compromise			
		•	nt neutropenia		
☐ Poor performance si	tatus 🔲 Previous chemotherapy	Previous radiation therapy	Previous episodes of FN		
	orbidities:	ease	function		
		ain:			
☐ Radiation therapy alone					
Yes No Are prolonged delays in I	radiation therapy expected due to	neutropenia?			
☐ Secondary prophylaxis of neutropenia					
Yes No Does the patient have a comparison of the patient experience.	•		w2		
		experienced from the prior cycle of c			
Neutropenic complication	on:		• •		
		patient received with the neutropenic			
☐ Yes ☐ No Did the patient experience			impacting the planned dose of		
	rior cycle of similar chemotherapy	? ose and schedule planned for current	t evelo?		
	e patient receive primary prophyla		. cycle:		
☐ Therapeutic use in a high-risk, febrile ne		,			
Please indicate which of the following pro		tient:			
☐ Age greater than 65					
	t the time of the development of t				
☐ Invasive fungal infec	de date of hospitalization:/				
		ion occurred:	Date: / /		
☐ Pneumonia	or rangar imposion and date impos				
	de date of pneumonia infection: _	1 1			
☐ Prior episodes of feb	•				
Prolonged neutrope			2		
└────────────────────────────────────					
☐ Sepsis syndrome					
☐ Other					
Please explain	in:				
☐ Treatment of high-risk neuroblastoma					
☐ Treatment for radiation injury					
Please indicate the radiation dose that ca	used the injury: grays (Gy	)			
For Continuation requests:					
Yes No Is this continuation request a I	result of the patient receiving sam euko (filgrastim-ayow), or Zarxio		ne (sargramostim), Neupogen (filgrastim),		
Yes No Is the patient continuing to res			larastim), Nivestvm (filarastim-aafi), Releuko		
(filgrastim-ayow), or Zarxio (fil		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	,, , , , , , , , , , , , , , , , , , ,		
H. ACKNOWLEDGEMENT					
Request Completed By (Signature Requi	red):		Date: / /		
Any person who knowingly files a request fo					
insurance company by providing materially					
insurance act, which is a crime and subjects			<del>-</del>		

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