

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

Page 1 of 5

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

PHONE: 1-855-364-0974 For other lines of business: Please use other form.

1-855-734-9389

For Ohio MMP:

FAX:

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

Please indicate: Start of treatment: Start dat				·	
☐ Continuation of therapy: Da Precertification Requested By:	ate of last treatment/	Phone:		Fax:	
A. PATIENT INFORMATION		1 Hone.		ı ax	
First Name:	Last Name:			DOB:	
Address:	City:			State:	ZIP:
Home Phone: Work Pho		Cell Phone:		Email:	ZIF.
		1 -		EIIIaII.	
	Patient Height: inches	s orcms /	Allergies:		
B. INSURANCE INFORMATION	Deep well-sub-leave eth		7 V		
Aetna Member ID #: Group #:	Does patient have oth If yes, provide ID#:		☐ Yes ☐ No Carrier Name:		
Insured:	Insured:		<u></u>		_
C. PRESCRIBER INFORMATION					
First Name:	Last Name:		(Check one):	☐ M.D. ☐ D	.O. N.P. P.A.
Address:	City:			State:	ZIP:
Phone: Fax:	St Lic #:	NPI #:	DEA #:		UPIN:
Provider Email:	Office Contact Name:	•	Phone:		•
D. DISPENSING PROVIDER/ADMINISTRATION	INFORMATION				
Place of Administration:			ovider/Pharmac	-	
Self-administered Physician's Office		☐ Physician's		☐ Retail Phar ☐ Mail Order	macy
Outpatient Infusion Center Phone: Center Name:			паппасу	_	
		Name:			
Agency Name:		_ Address:			
Administration code(s) (CPT):				Fax: NPI:	
Address: E. PRODUCT INFORMATION		_ TIN:		NPI	
Granix (tbo-filgrastim) Dose:	Directions for L	Jse:			
Leukine (sargramostim) Dose:		Jse:			
☐ Nivestym (filgrastim-aafi) Dose:		Jse:			
□ Neupogen (filgrastim) Dose:		Jse:			
Releuko (filgrastim-ayow) Dose:		Jse:			
Zarxio (filgrastim-sndz) Dose:	Directions for U	Jse:			
F. DIAGNOSIS INFORMATION - Please indicate	primary ICD code and specit	fy any other where	applicable.		
Primary Indication:		Other:			
G. CLINICAL INFORMATION - Required clinical i	nformation must be complete	ed in its <u>entirety</u> for	all precertificatio	n requests.	
For All requests (clinical documentation required	for all requests):				
Please indicate the patient's absolute neutrophil coun					N (6)
Yes No Does the patient have a nadir count that requires an immediate need for Granix (tbo-filgrastim), Leukine (sargramostim), Neupogen (filgrastim), Nivestym (filgrastim-aafi), Releuko (filgrastim-ayow), or Zarxio (filgrastim-sndz)?					
☐ Yes ☐ No Is the requested dose less than 180 mcg (0.3 mL)?					
└────────────────────────────────────					
	☐ Yes ☐ No Is the patient o			egimen that red	uires current use of
, -	this medication	n to remain unchang	ed?		•
Yes No Will Granix (tbo-filgrastim), Leukine (sargramostim), Neupogen (filgrastim), Nivestym (filgrastim-aafi), Releuko (filgrastim-ayow), or Zarxio (filgrastim-sndz) be used with another colony stimulating factor?					
🗀 Yes 🔲 No´ls Granix (tbo-filgrastim), Leukine (sargramostim), Neupogen (filgrastim), Nivestym (filgrastim-aafi), Releuko (filgrastim-ayow),					
or Zarxio (filgrastim-sndz) part of a stem cell mobilization protocol?					
☐ Yes ☐ No Will Granix (tbo-filgrastim), Neupogen (filgrastim), Nivestym (filgrastim-aafi), Releuko (filgrastim-ayow), or Zarxio (filgrastim-sndz) be used in combination with Leukine (sargramostim)?					
Yes No Will Granix (tbo-filgrastim), Leukine (sargramostim), Neupogen (filgrastim), Nivestym (filgrastim-aafi), Releuko (filgrastim-ayow) or Zarxio (filgrastim-sndz) be used in the same chemotherapy cycle as another colony stimulating factor?					
Yes No Is the patient currently receiving concomitant chemotherapy and radiation therapy?			ovew) or 7		
☐ Yes ☐ No Will Granix (tbo-filgrastim), Leukine (sargramostim), Neupogen (filgrastim), Nivestym (filgrastim-aafi), Releuko (filgrastim-ayow) or Zarxio (filgrastim-sndz) be used within 7 days of Neulasta (pegfilgrastim)?					



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB
G. CLINICAL INFORMATION (continued) - F	Required clinical information must be comp	leted in its entirety for all precerti	ication requests.
For Initiation requests:	•		•
Note: Granix, Leukine, Neupogen, Nivestym ☐ Yes ☐ No Has the patient had prior theral Releuko (filgrastim-ayow) within the last 365 da ☐ Yes ☐ No Has the patient had a trial and Please explain if there are any other medical re	py with Granix (tbo-filgrastim), Leukine (sa ys? failure, intolerance, or contraindication to 2	rgramostim), Neupogen (filgrastir Zarxio (filgrastim-sndz)?	n), Nivestym (filgrastim-aafi), or
Granix (tbo-filgrastim):			
Yes No Does the patient have a solid to significant incidence of febrile r	umor or non-myeloid malignancy and will r neutropenia for primary or secondary propl		herapy associated with a clinically
Leukine (sargramostim):			
Acute myeloid leukemia Yes No Is the patient receiving inc Please indicate the regir Yes No Is the patient receiving co	nen:		
Please indicate the regir			
Adjunct to progenitor cell-transplantation Please indicate which type of transplant at			<u> </u>
☐ Advanced HIV infection Please indicate the myelosuppressive anti ☐ Yes ☐ No Is the patient neutropenic		ng:	-
☐ Bone Marrow Transplantation			
☐ Yes ☐ No Does the patient have a double of the patient have and ☐ Yes ☐ No Is the medication being red ☐ Yes ☐ No Is the patient undergoing ☐ Please identify if the treation of the patient of the patient of the patient have a double of the patie	equested to reduce the duration of neutropoly myeloablative chemotherapy? atment will be followed by: Autologous I	enia and neutropenia-related infe	ctious complications?
☐ Congenital, cyclic or idiopathic neutrope	nia		
Please identify which documented type of Yes No Is the patient currently syn		al neutropenia 🔲 cyclic neutrope	enia 🔲 idiopathic neutropenia
☐ Drug- induced agranulocytosis ☐ Yes ☐ No Is the agranulocytosis cau ☐ Please provide the medi	used by chemotherapy? cation(s) that caused the agranulocytosis:		
☐ Hematopoietic Subsyndrome of Acute Ra ☐ Yes ☐ No Is the medication being re	adiation Syndrome (H-ARS)		a radiological/nuclear incident?
☐ Intermittent use in patients with myelody ☐ Yes ☐ No Does the patient have syr			
☐ Yes ☐ No Has the patient been tested		_	
		Da	ate obtained: ///
Yes No Does the patient present of Yes No Has a serum erythropoieti			
	It of the test and date obtained:	Da	ate obtained://
☐ Neuroblastoma			
☐ Yes ☐ No Is the patient's disease co	nsidered high-risk?		

☐ Yes ☐ No Will the requested medication be used in combination with ALL of the following medications: dinutuximab (Unituxin), interleukin-2

→ ☐ Yes ☐ No Will the requested medication be used in combination with Naxitamab-gqgk (Danyelza)?

(Aldesleukin), (Proleukin), isotretinoin (13-cis-retinoic acid)?

Continued on next page



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB
G. CLINICAL INFORMATION (continued) – F	Required clinical information must be comp	leted in its entirety for all precertit	ication requests.
Primary prophylaxis of neutropenia	•		·
	locumented diagnosis of non-myeloid malig	gnancy?	
Yes No Is the patient receiving my			
	of cancer the patient is being treated for:		
What is the expected percentage of febrile neu	chemotherapy regimen patient is currently by		
	10-19% (Intermediate risk)		
	to be at high risk for chemotherapy-induced		emplications?
T - :	f the following reasons that categorizes the	•	
☐ Active infections ☐	Age greater than or equal to 65 years	Bone marrow compromise	
	ement by tumor producing cytopenias 🔲 🤇		
•	atus 🔲 Previous chemotherapy 🔲 Prev	ious radiation therapy 🔲 Previo	us episodes of FN
Recent surgery	_	<u></u>	_
	rbidities: 🔲 Cardiovascular disease 🔲 l	HIV infection	n 🔲 Renal dysfunction
Other- Please explain	n:		
☐ Secondary prophylaxis of neutropenia			
	locumented diagnosis of non-myeloid malig e a febrile neutropenic complication from a		
	ropenic complication the patient experience		herany:
Neutropenic complicatio		ou nom the prior cycle of chamet	iorapy.
	cycle of chemotherapy that the patient rec	eived with the neutropenic comp	ication:
☐ Yes ☐ No Did the patient experience	e a dose-limiting neutropenic event (a nadir	or day of treatment count impac	ting the planned dose of
	or cycle of similar chemotherapy?		_
	e patient treated with the same dose and so		?
☐ Yes ☐ No Did the	patient receive primary prophylaxis against	t febrile neutropenia?	
☐ Therapeutic use in a high-risk, febrile ne			
Please indicate which of the following prog			
☐ Age greater than 65 g			
	the time of the development of fever e date of hospitalization: / /		
☐ Invasive fungal infect		_	
	of fungal infection and date infection occurre	ed·	Date: / /
☐ Pneumonia	or rangar innection and date innection ecount	<u> </u>	
——————————————————————————————————————	e date of pneumonia infection:/	/	
☐ Prior episodes of feb	rile neutropenia		
☐ Prolonged neutropen			
_ · — —	s Is the prolonged neutropenia expected to	o last greater than 10 days?	
☐ Profound neutropenia	a		
☐ Sepsis syndrome			
Other			
Please explain		(filamotim and).	
Neupogen (filgrastim), Nivestym (filgrastim-	aan), Releuko (filgrastim-ayow), Zarxio i	(nigrastim-sndz):	
☐ Acute lymphoblastic leukemia (ALL)☐ Yes ☐ No Has the first days of chem	notherany been completed?		
Yes No Is this the initial induction			
Yes No Is this the first post-remise			
	n and date started: Regimen:		Date started://
☐ Acute myeloid leukemia			
☐ Yes ☐ No Is the patient receiving inc	duction chemotherapy?		
> Please indicate the regin	men:		
Yes No Is the patient receiving co	nsolidation chemotherapy?		
Please indicate the regir	men: emotherapy for relapsed or refractory disea	2002	
Yes No is the patient receiving ch	lemomerapy for relapsed or retractory disea □ Refractory disease	45 C (
Please indicate the regin	men:		
	-		•



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB
G. CLINICAL INFORMATION (continued) – I	Required clinical information must be con	npleted in its entirety for all prece	ertification requests
Adjunct to progenitor cell-transplantatio			ranoaton roquocto.
Please indicate which type of transplant a			/ /
☐ Advanced HIV infection	_ 3 _	· <u> </u>	
Please indicate the myelosuppressive ant Yes No Is the patient neutropenic	•	ving:	
☐ Bone Marrow Transplantation			
Yes No Does the patient have a			
Yes No Is the medication being re		penia and neutropenia-related in	fectious complications?
Yes No Is the patient undergoing	atment will be followed by: Autologous	s hone marrow transplantation	
y 1 loade ladilary is allo all		bone marrow transplantation	
│ │	_		
Please identify which documented type of	f neutropenia that patient has: 🗌 congen	ital neutropenia 🔲 cyclic neutro	openia 🔲 idiopathic neutropenia
Yes No Is Granix (tbo-filgrastim),		astim), Nivestym (filgrastim-aafi)	, Releuko (filgrastim-ayow),
or Zarxio (filgrastim-sndz) being requested for chronic administration to reduce the incidence and duration of sequelae of neutropenia (e.g., fever, infections, oropharyngeal ulcers)?			
☐ Chronic Myeloid Leukemia			
Yes No Does the patient have res		stions?	
Yes No Is the neutropenia second	☐ Gleevec (imatinib) ☐ Iclusig (ponatir		Tasigna (nilotinih)
☐ Drug- induced agranulocytosis		iib)	ruoigna (mount)
☐ Yes ☐ No Is the agranulocytosis ca	used by chemotherapy? lication(s) that caused the agranulocytosi	s.	
☐ Glycogen storage disease (GSD) type 1	incation(e) that sadeca the agranales, test	o	
Yes No Does the patient have a	ow neutrophil count?		
☐ Hairy Cell Leukemia			
☐ Yes ☐ No Does the patient have cli		ing chemotherapy?	
☐ Increase dose intensity chemotherapy re		amanatratas that dags intensive	thereny produces improvement in
Yes No Is the patient being treated disease control?	ed in a setting in which clinical research d	emonstrates that dose-intensive	therapy 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 ☐ Previous chemotherapy ☐ Previous radiation therapy ☐ Previous episodes of FN			
☐ Recent surgery			
Other serious co-mo	rbidities: Cardiovascular disease Other- Please explain:	HIV infection	
☐ Intermittent use in patients with myelody			
☐ Yes ☐ No Does the patient have sy	mptomatic anemia?		
Yes No Has the patient been test			Data obtained:
Yes No Does the patient present	ult of the test and date obtained: with other cytogenetic abnormalities?		Date obtained/_/
Yes No Has a serum erythropoiet	tin test been completed?		
	ult of the test and date obtained:		Date obtained: / /
Lymphoma	that the nations is being to the standards	tive shamatharass /s = /D OLL	OD) vituvimah avalar barrida
	orednisone) or more aggressive regimens	?	
Please indicate the pati	ent's chemotherapy regimen:		



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB
G. CLINICAL INFORMATION (continued) –	Required clinical information must be comp	l eleted in its <u>entirety</u> for all pre	certification requests.
☐ Primary prophylaxis of neutropenia		· · · · · · · · · · · · · · · · · · ·	•
	documented diagnosis of non-myeloid mali	gnancy?	
Yes No Is the patient receiving r			
	e of cancer the patient is being treated for:		
	chemotherapy regimen patient is currently le neutropenia incidence from the chemothe		
] 10-19% (Intermediate risk) \square 20% or gr		
	I to be at high risk for chemotherapy-induce		us complications?
	of the following reasons that categorizes the	-	
	Age greater than or equal to 65 years		
	vement by tumor producing cytopenias		
☐ Poor performance s	status	nous radiation therapy \square P	revious episodes of FIN
	orbidities: Cardiovascular disease	HIV infection □ Liver dysfu	unction
_ Guidi dellede de lin	Other- Please explain:		
☐ Radiation therapy alone			
☐ Yes ☐ No Are prolonged delays in	radiation therapy expected due to neutrope	nia?	
☐ Secondary prophylaxis of neutropenia			
	documented diagnosis of non-myeloid mali		2
	ce a febrile neutropenic complication from a utropenic complication the patient experience		
Neutropenic complicati		ica irom the prior by old or on	smoundaby.
	or cycle of chemotherapy that the patient re	ceived with the neutropenic of	complication:
	ce a dose-limiting neutropenic event (a nad	r or day of treatment count ir	npacting the planned dose of
	rior cycle of similar chemotherapy?		
	ne patient treated with the same dose and se patient receive primary prophylaxis agains		cycle?
		st lebrile fleutroperila :	
Therapeutic use in a high-risk, febrile n Please indicate which of the following pro			
☐ Age greater than 65			
☐ Being hospitalized a	at the time of the development of fever		
	de date of hospitalization://	<u> </u>	
Invasive fungal infe			D
Provide type	of fungal infection and date infection occur	red:	Date: /
	de date of pneumonia infection:/	1	
☐ Prior episodes of fe		<u> </u>	
□ Prolonged neutrope	enia		
	No Is the prolonged neutropenia expected	to last greater than 10 days?	
☐ Profound neutroper	nia		
☐ Sepsis syndrome ☐ Other			
T '	·		
•	iin:		
☐ Treatment of high-risk neuroblastoma ☐ Treatment for radiation injury			
Please indicate the radiation dose that ca	aused the injury grays (Gv)		
For Continuation requests:	g.a.y. (3)/		
Yes No Is this continuation request a	result of the patient receiving samples of G	ranix (tbo-filgrastim), Leukine	e (sargramostim), Neupogen (filgrastim),
Nivestym (filgrastim-aafi), Re	leuko (filgrastim-ayow), or Zarxio (filgrastim	-sndz)?	
Yes No Is the patient continuing to re		argramostim), Neupogen (filg	rastim), Nivestym (filgrastim-aafi), Releuko
(filgrastim-ayow), or Zarxio (fi	igrastim-sndz) tnerapy?		
H. ACKNOWLEDGEMENT			
Request Completed By (Signature Requ	ired):		Date: /
Any person who knowingly files a request for			
insurance company by providing materiall insurance act, which is a crime and subject			se of misleading, commits a fraudulent

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