AvMed

PHARMACY PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

<u>Directions</u>: <u>The prescribing physician must sign and clearly print name (preprinted stamps not valid)</u> on this request. All other information may be filled in by office staff; <u>fax to 1-305-671-0200</u>. No additional phone calls will be necessary if all information (<u>including phone and fax #s</u>) on this form is correct. <u>If the information provided is not complete, correct, or legible, the authorization process can be delayed.</u>

Drug Requested: COLONY STIMULATING FACTORS

[Form to be completed **ONLY** if the member is self-administering]

Short-actin	ng Granulocyte Colon	y-Stimulating	Factors (G-CSFs)
□ Granix [®] (tbo-filgrastin	n)	(filgrastim)	□ Nivestym [™] (filgrastim-aafi)
□ Nypozi [™] (filgrastim-tx	id)	ilgrastim-ayow)	□ Zarxio [®] (filgrastim-sndz)
Granulocy	te-macrophage Colony	y-Stimulating	Factor (GM-CSF)
□ Leukine ® (sargramosti	im)		
	ng Granulocyte Colony	y-Stimulating	Factors (G-CSFs)
□ Fulphila [™] (pegfilgrast	im-jmdb)	□ Ryzneuta	® (efbemalenograstim alfa-vuxw)
□ Fylnetra [™] (pegfilgrasi	tim-pbbk)	□ Stimufeno	d® (pegfilgrastim-fpgk)
□ Neulasta® (pegfilgrast	im)	□ Udenyca®	(pegfilgrastim-cbqv)
□ Nyvepria [™] (pegfilgras	stim-apgf)	□ Ziextenzo	(pegfilgrastim-bmez)
□ Rolvedon [™] (eflapegra	stim-xnst)		
MEMBER & PRESCH Member Name: Member AvMed #:			on may be delayed if incomplete. Date of Birth:
Prescriber Name:			
Prescriber Signature:			
Office Contact Name:			
Phone Number:		Fax Nui	mber:
NPI #:			
DRUG INFORMATIO	N : Authorization may be	delayed if incomp	olete.
Drug Name/Form/Strength	:		
Dosing Schedule:		Length of T	Therapy:
Diagnosis:		ICD Code,	if applicable:
Weight (if applicable):		Date v	veight obtained:

Maximum Daily Dose:

Fulphila 6 mg prefilled syringe: 1 syringe/14 days	Nypozi 300 mcg prefilled syringe: 3 syringes/1 day
Fylnetra 6 mg prefilled syringe: 1 syringe/14 days	Nypozi 480 mcg prefilled syringe: 3 syringes/1 day
Granix 300 mcg prefilled syringe: 3 syringes/1 day	Releuko 300 mcg vial: 3 vials/1 day
Granix 300 mcg single-dose vial: 3 vials/1 day	Releuko 300 mcg prefilled syringe: 3 syringes/1 day
Granix 480 mcg prefilled syringe: 3 syringes/1 day	Releuko 480 mcg vial: 3 vials/1 day
Granix 480 mcg single-dose vial: 3 vials/1 day	Releuko 480 mcg prefilled syringe: 3 syringes/1 day
Leukine 250 mcg vial: 28 vials/14 days	Rolvedon 13.2 mg prefilled syringe: 1 syringe/14 days
Neulasta 6 mg prefilled syringe: 1 syringe/14 days	Ryzneuta 20 mg prefilled syringe: 1 syringe/14 days
Neulasta 6 mg prefilled syringe kit: 1 kit/14 days	Stimufend 6 mg prefilled syringe: 1 syringe/14 days
Neupogen 300 mcg vial: 3 vials/1 day	Udenyca 6 mg prefilled syringe: 1 syringe/14 days
Neupogen 300 mcg SingleJect: 3 syringes/1 day	Udenyca 6 mg auto-injector: 1 injection/14 days
Neupogen 480 mcg vial: 3 vials/1 day	Udenyca 6 mg onbody (syringe, with wearable injector): 1 syringe/14 days
Nivestym 300 mcg prefilled syringe: 3 syringes/1 day	Zarxio 300 mcg prefilled syringe: 3 syringes/1 day
Nivestym 480 mcg vial: 3 vials/1 day	Zarxio 480 mcg prefilled syringe: 3 syringes/1 day
Nivestym 480 mcg prefilled syringe: 3 syringes/1 day	Ziextenzo 6 mg prefilled syringe: 1 syringe/14 days
Nyvepria 6 mg prefilled syringe: 1 syringe/14 days	

CLINICAL CRITERIA: Check below all that apply. All criteria must be met for approval. To support each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be provided or request may be denied.

<u>PROVIDER PLEASE NOTE</u>: SUBMISSION OF APPLICABLE DOCUMENTATION IS NECESSARY (I.E. CHART NOTES, DISEASE HISTORY, CURRENT/PAST THERAPY RECORD, COMPLETE BLOOD COUNT OR OTHER LABORATORY RESULTS) FOR COMPLETION OF REQUEST

□ Short-acting Granulocyte Colony-Stimulating Factors (G-CSFs)

PLEASE SELECT ONE OF THE FOLLOWING INDICATIONS FOR USE:

- ☐ Medication will be used as primary prevention of febrile neutropenia in members with non-myeloid malignancy meeting <u>ONE</u> of the following [Length of authorization = 6 months]:
 - ☐ Member is undergoing myelosuppressive chemotherapy with an expected incidence of febrile neutropenia greater than 20%

	neı	ember is undergoing myelosuppressive chemotherapy with an expected incidence of febrile atropenia of 10% to < 20% AND one or more of the following co-morbidities (select all that oly):
		Age >65 years receiving full dose intensity chemotherapy
	_	Extensive prior exposure to chemotherapy
		Previous exposure of pelvis, or other areas of large amounts of bone marrow, to radiation
		Persistent neutropenia (ANC $\leq 1000/\text{mm}^3$)
		Bone marrow involvement by tumor
		Member has a condition that can potentially increase the risk of serious infection (i.e., HIV/AIDS with low CD4 counts)
		Recent surgery and/or open wounds
		Poor performance status
		Renal dysfunction (creatinine clearance <50 mL/min)
		Liver dysfunction (elevated bilirubin >2.0 mg/dL)
		Chronic immunosuppression in the post-transplant setting, including organ transplant
		<u>OR</u>
is n	eed	er is 18 years of age or older, has a diagnosis of acute myeloid leukemia, <u>AND</u> filgrastim therapy led shortly following completion of induction or consolidation chemotherapy [Length of rization = 6 months]
		<u>OR</u>
		er has been acutely exposed to myelosuppressive doses of radiation (Hematopoietic Acute ion Syndrome [H-ARS]) [Length of authorization = Date of service only]
		<u>OR</u>
		er has been diagnosed with a non-myeloid malignancy, <u>AND</u> will be receiving myeloablative therapy following a bone marrow transplant [Length of authorization = Date of service only]
		<u>OR</u>
		ation will be used in apheresis collection of autologous hematopoietic progenitor cells [Length of cization = Date of service only]
		<u>OR</u>
sho	wir	er has been diagnosed with congenital, cyclic, or idiopathic neutropenia, <u>AND</u> is currently ag symptoms and incidence of complications (e.g., fever, infections, oropharyngeal ulcers) the of authorization = 12 months
		<u>OR</u>
Tre	atn	nent with filgrastim is needed as adjunctive treatment of febrile neutropenia when primary

<u>OR</u>

authorization = 6 months]

prophylaxis with a long-acting granulocyte colony stimulating factor is not given [Length of

- Adjunctive treatment of febrile neutropenia is considered clinically appropriate when at least **ONE** of the following risk factors are present (in the absence of prior growth factor use within the same chemotherapy cycle of treatment) (select all that apply) [Length of authorization = 6 months]:
 - \square Age > 65 years
 - □ Neutrophil recovery is expected to be delayed (greater than 10 days)
 - \square Neutropenia is profound (less than 0.1 x 10⁹)
 - □ Active pneumonia
 - ☐ Sepsis syndrome (hypotension and/or multi-organ damage/dysfunction noted)
 - ☐ Invasive fungal or opportunistic infection
 - Onset of fever during inpatient stay

NOTE: Febrile neutropenia is defined as an oral temperature > 38.3°C (101.0°F) or 2 consecutive readings of 38.0°C (100.4°F) for 1 hour, with an absolute neutrophil count less than 500 cells/ μ L (0.5 x 10°/L) or less than 1000 cells/ μ L and expected to fall below 500 cells/ μ L over the next 48 hours.

<u>OR</u>

☐ Member has a diagnosis of primary myelodysplastic syndrome, <u>AND</u> filgrastim therapy will be used in combination with epoetin to treat anemia [Length of authorization = 6 months]

<u>OR</u>

□ Member has a diagnosis of non-Hodgkin lymphoma or multiple myeloma, <u>AND</u> filgrastim therapy will be used in combination with plerixafor for the collection of progenitor cells leading to subsequent autologous transplantation. [Length of authorization = Date of service only]

NOTE: Mozobil (plerixafor) requires prior authorization

□ Granulocyte-macrophage Colony-Stimulating Factor (GM-CSF) [Leukine]

PLEASE SELECT ONE OF THE FOLLOWING INDICATIONS FOR USE:

■ Member is 55 years of age or older, has a diagnosis of acute myeloid leukemia, <u>AND</u> sargramostim therapy is needed shortly after the completion of induction or repeat induction of chemotherapy [Length of authorization = 6 months]

<u>OR</u>

■ Member is 2 years of age or older, <u>AND</u> sargramostim therapy is needed for faster reconstitution of myeloid to prepare for allogeneic bone marrow transplant (NOTE: confirmation of HLA-matched donor status is required) [Length of authorization = 6 months]

<u>OR</u>

Member is 2 years of age or older, has undergone bone marrow transplant (allogeneic or autologous),
 AND sargramostim therapy is needed because there is delayed or failed neutrophil recovery [Length of authorization = 6 months]

<u>OR</u>

☐ Medication will be used in apheresis collection of autologous hematopoietic progenitor cells [Length of authorization = Date of service only]

OR

☐ Member is 2 years of age or older, has a diagnosis of acute lymphoblastic leukemia (ALL), Hodgkin lymphoma (HL), or non-Hodgkin lymphoma (NHL), <u>AND</u> sargramostim therapy is needed for faster reconstitution of myeloid following an autologous peripheral blood progenitor cell transplant or bone marrow transplant [Length of authorization = 6 months]

OR

☐ Member has been acutely exposed to myelosuppressive doses of radiation (Hematopoietic Acute Radiation Syndrome [H-ARS]) [Length of authorization = Date of service only]

<u>OR</u>

- ☐ Member has a diagnosis of high-risk neuroblastoma, <u>AND</u> sargramostim is needed for combination therapy with a with GD2-binding monoclonal antibody (i.e., dinutiximab or naxitamab) [Length of authorization = 6 months]
- □ Long-acting Granulocyte Colony-Stimulating Factors (G-CSFs)

PLEASE SELECT ONE OF THE FOLLOWING INDICATIONS FOR USE:

- ☐ Medication will be used as primary prevention of febrile neutropenia in members with non-myeloid malignancy meeting <u>ONE</u> of the following [Length of authorization = 6 months]:
 - ☐ Member is undergoing myelosuppressive chemotherapy with an expected incidence of febrile neutropenia greater than 20%
 - ☐ Member is undergoing myelosuppressive chemotherapy with an expected incidence of febrile neutropenia of 10% to < 20% **AND one or more** of the following co-morbidities (select all that apply):
 - ☐ Age >65 years receiving full dose intensity chemotherapy
 - □ Extensive prior exposure to chemotherapy
 - ☐ Previous exposure of pelvis, or other areas of large amounts of bone marrow, to radiation
 - \square Previous/persistent neutropenia (ANC $\leq 1000/\text{mm}3$)
 - □ Bone marrow involvement by tumor
 - □ Patient has a condition that can potentially increase the risk of serious infection (i.e., HIV/AIDS with low CD4 counts)
 - □ Recent surgery and/or open wounds
 - □ Poor performance status
 - □ Renal dysfunction (creatinine clearance <50 mL/min)
 - □ Liver dysfunction (elevated bilirubin >2.0 mg/dL)
 - Chronic immunosuppression in the post-transplant setting, including organ transplant

<u>OR</u>

Member has been acutely exposed to myelosuppressive doses of radiation (Hematopoietic Acute
Radiation Syndrome [H-ARS]) [Length of authorization = Date of service only]

OR

☐ Medication will be used as secondary prevention of febrile neutropenia in members with non-myeloid malignancy, <u>AND</u> having experienced a neutropenic complication from a prior cycle of the same chemotherapy [Length of authorization = 6 months]

OR

☐ Treatment with requested medication is needed as adjunctive treatment of febrile neutropenia when primary prophylaxis is not given [Length of authorization = 6 months]

OR

- Adjunctive treatment of febrile neutropenia is considered clinically appropriate when at least **ONE** of the following risk factors are present (in the absence of prior growth factor use within the same chemotherapy cycle of treatment) (select all that apply) [Length of authorization = 6 months]:
 - \Box Age > 65 years
 - □ Neutrophil recovery is expected to be delayed (greater than 10 days)
 - \Box Neutropenia is profound (less than 0.1 x 10⁹)
 - □ Active pneumonia
 - ☐ Sepsis syndrome (hypotension and/or multi-organ damage/dysfunction noted)
 - ☐ Invasive fungal or opportunistic infection
 - Onset of fever during inpatient stay

NOTE: Febrile neutropenia is defined as an oral temperature > 38.3°C (101.0°F) or 2 consecutive readings of 38.0°C (100.4°F) for 1 hour, with an absolute neutrophil count less than 500 cells/ μ L (0.5 x 10°/L) or less than 1000 cells/ μ L and expected to fall below 500 cells/ μ L over the next 48 hours

<u>OR</u>

☐ Treatment with requested medication is needed after bone marrow transplantation (BMT) failure or engraftment delay [Length of authorization = 6 months]

OR

	For medical necessity on a treatment purpose not listed, please provide clinical rationale
	and submit any chart notes/literature you feel would be pertinent in support of medical necessity:
Medicati	
·ICGICU	ion being provided by Specialty Pharmacy – Proprium Rx
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