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-acting Granulocyte Colony-Stimulating Factors (G-CSFs)									
	Granix® (tbo-filgrastim)		Neupogen® (fi		rastim)		Nivestym ¹	(filgrastim-aafi)	
	Releuko® (filgrastim-ayow)		Zarxio® (filgras		m-sndz)				
	Granulocyte-macrophage Colony-Stimulating Factor (GM-CSF)								
□ Leukine® (sargramostim)									
	Long-acting Granulocyte Colony-Stimulating Factors (G-CSFs)								
	Fulphila [™] (pegfilgrastim-jmdb)			□ Ryzneuta [®] (efbemalenograstim alfa-vuxw)					
□ Fylnetra [™] (pegfilgrastim-pbbk)			□ Stimufend® (pegfilgrastim-fpgk)						
□ Neulasta [®] (pegfilgrastim)			□ Udenyca® (pegfilgrastim-cbqv)						
				□ Ziextenzo [™] (pegfilgrastim-bmez)					
□ Rolvedon [™] (eflapegrastim-xnst)									
MEMBER & PRESCRIBER INFORMATION: Authorization may be delayed if incomplete. Member Name: Member AvMed #: Prescriber Name: Prescriber Signature: Date:									
Pho	Office Contact Name: Fax Number: DEA OR NPI #:								
DRUG INFORMATION: Authorization may be delayed if incomplete.									
Drug Form/Strength:									
Dosing Schedule:					Length of Therapy:				
Diagnosis:					ICD Code:				
Weight:					ate:				

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Maximum Daily Dose:

Fulphila 6 mg prefilled syringe: 1 syringe/14 days	Nyvepria 6 mg prefilled syringe: 1 syringe/14 days
Fylnetra 6 mg prefilled syringe: 1 syringe/14 days	Releuko 300 mcg vial: 3 vials/1 day
Granix 300 mcg prefilled syringe: 4 syringes/1 day	Releuko 300 mcg prefilled syringe: 3 syringes/1 day
Granix 300 mcg single-dose vial: 4 vials/1 day	Releuko 480 mcg vial: 3 vials/1 day
Granix 480 mcg prefilled syringe: 3 syringes/1 day	Releuko 480 mcg prefilled syringe: 3 syringes/1 day
Granix 480 mcg single-dose vial: 3 vials/1 day	Rolvedon 13.2 mg prefilled syringe: 1 syringe/14 days
Leukine 250 mcg vial: 28 vials/14 days	Ryzneuta 20 mg prefilled syringe: 1 syringe/14 days
Neulasta 6 mg prefilled syringe: 1 syringe/14 days	Stimufend 6 mg prefilled syringe: 1 syringe/14 days
Neulasta 6 mg prefilled syringe kit: 1 kit/14 days	Udenyca 6 mg prefilled syringe: 1 syringe/14 days
Neupogen 300 mcg vial: 3 vials/1 day	Udenyca 6 mg auto-injector: 1 injection/14 days
Neupogen 300 mcg SingleJect: 3 syringes/1 day	Udenyca 6 mg onbody (syringe, with wearable injector): 1 syringe/14 days
Neupogen 480 mcg vial: 3 vials/1 day	Zarxio 300 mcg prefilled syringe: 3 syringes/1 day
Nivestym 300 mcg prefilled syringe: 3 syringes/1 day	Zarxio 480 mcg prefilled syringe: 3 syringes/1 day
Nivestym 480 mcg vial: 3 vials/1 day	Ziextenzo 6 mg prefilled syringe: 1 syringe/14 days
Nivestym 480 mcg prefilled syringe: 3 syringes/1 day	

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%
 - ☐ 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 Persistent neutropenia (ANC $\leq 1000/\text{mm}^3$)
 - □ Bone marrow involvement by tumor

PA GCSF (AvMed) (Continued from previous page)

	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>
needed	er is 18 years of age or older, has a diagnosis of acute myeloid leukemia, <u>AND</u> filgrastim therapy is I 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 rization = Date of service only]
	<u>OR</u>
sympto	er has been diagnosed with congenital, cyclic, or idiopathic neutropenia, <u>AND</u> is currently showing oms and incidence of complications (e.g., fever, infections, oropharyngeal ulcers) [Length of rization = 12 months]
	<u>OR</u>
prophy	nent with filgrastim is needed as adjunctive treatment of febrile neutropenia when primary vlaxis with a long-acting granulocyte colony stimulating factor is not given [Length of rization = 6 months]
	<u>OR</u>
follow cycle o	ctive treatment of febrile neutropenia is considered clinically appropriate when at least <u>ONE</u> of the ing risk factors are present (in the absence of prior growth factor use within the same chemotherapy of treatment) (select all that apply) [Length of authorization = 6 months]: 10
_	utrophil recovery is expected to be delayed (greater than 10 days)
	utropenia is profound (less than 0.1 x 10 ⁹)
	tive pneumonia
	osis syndrome (hypotension and/or multi-organ damage/dysfunction noted)
_ 50	(Continued on next page)
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☐ Member has a condition that can potentially increase the risk of serious infection (i.e., HIV/AIDS

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- ☐ 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.

OR

☐ 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]

OR

☐ 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]

OR

■ 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]

OR

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

<u>OR</u>

☐ 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]

<u>OR</u>

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 $\le 1000/\text{mm}3$)
 - □ 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

OR

☐ 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]

<u>OR</u>

Treatment with requested medication is needed as adjunctive treatment of febrile neutropenia when primary prophylaxis is not given [Length of authorization = 6 months]
<u>OR</u>
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)
□ Neutropenia is profound (less than 0.1×10^9)
□ 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 ⁹ /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]
<u>OR</u>
Medication will be used in apheresis collection of autologous hematopoietic progenitor cells [Length of authorization = Date of service only]
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:
edication being provided by Specialty Pharmacy – Proprium Rx **Use of samples to initiate therapy does not meet step edit/ preauthorization criteria. **
Ose of samples to initiate incrupy aves not meet step eath predutitorization Criteria. ""

*Previous therapies will be verified through pharmacy paid claims or submitted chart notes. *

^{*}Approved by Pharmacy and Therapeutics Committee: 2/16/2023; 3/21/2024
REVISED/UPDATED/REFORMATTED: 2/9/2009; 6/14/2011; 8/19/2011; 1/23/2012; 1/14/2014; 4/9/2014; 5/7/2014; 5/28/2014; 8/13/2014; 1/31/2014; 5/21/2015; 12/27/2015; 6/9/2016; 8/19/2016; 9/22/2016; 12/11/2016; 8/3/2017; 5/14/2019; 8/6/2019; 12/20/2021; 1/12/2022; 2/23/2022; 3/23/2022;03/09/2023; 10/26/2023; 6/14/2024