

## FLORIDA MEDICAID PRIOR AUTHORIZATION

## **COLONY STIMULATING FACTORS**

Preferred: Leukine®, Neupogen®, Nyvepria™

Clinical PA required (Non-Preferred): Fulphila™/Fylnetra®/Granix®/Neulasta®/Nivestym®/Releuko®/Rolvedon™/Stimufend®/Udenyca®/Zarxio®/Ziextenzo™

Note: Form must be completed in full. An incomplete form may be returned.

Recipient's Medicaid ID #										Date of Birth (MM/DD/YYYY)																			
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Recipient's Full Name																			ı		ı	_							
Pre	Prescriber's Full Name																												
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	<ol> <li>What is the diagnosis or the indication for the product? Please check below AND submit supporting documentation indicating the diagnosis.</li> </ol>															iiig													
	Cancer patient receiving myelosuppressive chemotherapy																												
	Cancer patient receiving myelosuppressive chemotherapy  Cancer patient receiving bone marrow transplant																												
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☐ Patient receiving induction or consolidated chemotherapy for acute myeloid leukemia (AML)																													
	<ul><li>☐ Peripheral blood progenitor cell collection and therapy in cancer patient</li><li>☐ Acute exposure to myelosuppressive doses of radiation in patient</li></ul>																												
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Mail or Fax Information to:

Magellan Medicaid Administration, Inc. Prior Authorization P. O. Box 7082

Tallahassee, FL 32314-7082 Phone: 877-553-7481 Fax: 877-614-1078 **Confidentiality Notice:** The documents accompanying this transmission contain confidential health information that is legally privileged. If you are not the intended recipient, you are hereby notified that any disclosure, copying, distribution, or action taken in reliance on the contents of these documents is strictly prohibited. If you have received this information in error, please notify the sender (via return fax) immediately and arrange for the return or destruction of these documents. Distribution, reproduction or any other use of this transmission by any party other than the intended recipient is strictly prohibited.



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Recip	ienť	s Fu	ıll N	ame	•																						
2.	This is: New therapy OR Continuation of therapy																										
3.	Can the prescriber attest the disease state or prescribed regimen is high risk (> 20%) for febrile neutropenia?    Yes    No																										
4.		Lab test date: cells/mm <sup>3</sup>																									
5.	Wh	What is the date range of therapy? Begin date:														End date:											
6.	What will be the dosage and frequency of dosing?																										
Prescriber's Signature: Date:																											

REQUIRED FOR REVIEW: Copies of medical records (i.e., diagnostic evaluations and recent chart notes) and the most recent copies of related labs. The provider must retain copies of all documentation for five years.

Phone: 877-553-7481 Fax: 877-614-1078

# WATE OF FLORIDA

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Stimufend<sup>®</sup>/Udenyca<sup>®</sup>/Zarxio<sup>®</sup>/Ziextenzo<sup>™</sup>

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# Approved Indications for Zarxio® and Nivestym®

- Cancer patients (Note that they do not have to meet ANC criteria. If they have the indication, approve.):
  - If patient has not yet undergone chemotherapy but it has been prescribed, no ANC is required
  - Cancer patients receiving myelosuppressive chemotherapy (approve up to 12 months)
  - Cancer patients receiving bone marrow transplants (approve up to 12 months)
  - Patient receiving induction or consolidated chemotherapy for AML (approve up to 12 months)
  - Peripheral blood progenitor cell collection and therapy in cancer patients (approve up to 12 months)
- Severe chronic neutropenia ANC now required
  - All lab documentation must be on official lab letterhead handwritten labs are not acceptable
  - The ANC is 1500 or less
  - Congenital, cyclic, or idiopathic (approve up to 12 months)
- AIDS ANC required
  - Severe neutropenia in AIDS patients on antiretroviral therapy
  - Initial Therapy: ANC is 1000 or less
  - Continuation of Therapy: ANC is 1600 or less
  - All lab documentation must be on official lab letterhead handwritten labs are not acceptable.
     (Approve for 6 months)

# Approved Indications for Releuko®

- Cancer patients (Note that they do not have to meet ANC criteria. If they have the indication, approve.):
  - If patient has not yet undergone chemotherapy but it has been prescribed, no ANC is required
  - Cancer patients receiving myelosuppressive chemotherapy (approve up to 12 months)
  - Cancer patients receiving bone marrow transplants (approve up to 12 months)
  - Patient receiving induction or consolidated chemotherapy for AML (approve up to 12 months)
- Severe chronic neutropenia ANC now required
  - All lab documentation must be on official lab letterhead handwritten labs are not acceptable
  - The ANC is 1500 or less
  - Congenital, cyclic, or idiopathic (approve up to 12 months)

# WATE OF FLORIDA

### FLORIDA MEDICAID PRIOR AUTHORIZATION

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# Fulphila™/FyInetra®/Granix®/Neulasta®/Nivestym®/Releuko®/Rolvedon™/ Stimufend®/Udenyca®/Zarxio®/Ziextenzo™

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# Approved Indications for Udenyca<sup>®</sup>, Neulasta<sup>®</sup>, Ziextenzo<sup>™</sup>, Fulphila<sup>™</sup>, Fylnetra<sup>®</sup>, Rolvedon<sup>™</sup>, and Stimufend<sup>®</sup>

- Chemotherapy-induced neutropenia
  - Cancer patient with non-myeloid malignancies receiving myelosuppressive chemotherapy (approve up to 12 months)
- Dosage: 6 mg subcutaneous once per chemotherapy cycle
- Patient acutely exposed to myelosuppressive doses of radiation (hematopoietic syndrome of acute radiation syndrome) (Neulasta® and Udenyca® only)
- Dosage: Two doses, 6 mg subcutaneous, each one week apart

### Note:

- Do not administer in the period 14 days before and 24 hours after administration of cytotoxic chemotherapy.
- Documentation of the ANC and/or lab values is not required.
- Not indicated for severe chronic neutropenia.
- Not indicated for neutropenia associated with human immunodeficiency virus (HIV)/AIDS.

# Approved Indications for Granix®

- Chemotherapy-induced neutropenia:
  - Cancer patient with non-myeloid malignancies receiving myelosuppressive chemotherapy (approve up to 12 months)
  - Dosage: 5 mcg/kg/day subcutaneously

## Note:

- Do not administer in the period 24 hours before and 24 hours after administration of cytotoxic chemotherapy.
- Documentation of the ANC and/or lab values is not required.
- Not indicated for severe chronic neutropenia.
- Not indicated for neutropenia associated with HIV/AIDS.