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Section:	Prescription Drugs	Effective Date:	July 1, 2026
Subsection:	Hematological Agents	Original Policy Date:	December 7, 2011
Subject:	Neulasta Fulphila Fylnetra Nyvepria Stimufend Udenyca Ziextenzo	Page:	1 of 6

Last Review Date: June 11, 2026

Neulasta Fulphila Fylnetra Nyvepria Stimufend Udenyca Ziextenzo

Description

Neulasta, Neulasta Onpro (pegfilgrastim), **Fulphila** (pegfilgrastim-jmdb), Fylnetra (pegfilgrastim-pbbk), Nyvepria (pegfilgrastim-apgf), Stimufend (pegfilgrastim-fpgk), **Udenyca**, **Udenyca Onbody** (pegfilgrastim-cbqv), Ziextenzo (pegfilgrastim-bmez)

Preferred products: Fulphila, Udenyca, Udenyca Onbody

Background

Neutropenia occurs when an individual has an abnormally low level of neutrophils, a type of white blood cell important in fighting off infections. Neutropenia and its complications, including febrile neutropenia and infection, remain major toxicities associated with myelosuppressive systemic cancer chemotherapy. Colony stimulating factors are medications used to stimulate the production of neutrophils. Neulasta (pegfilgrastim) and its biosimilars are granulocyte colony-stimulating factors (G-CSF) that act on hematopoietic cells by binding to specific cell surface receptors, thereby stimulating proliferation, differentiation, commitment, and end cell functional activation. Fulphila, Fylnetra, Nyvepria, Udenyca, and Ziextenzo are biosimilars to

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Neulasta. The FDA defines biosimilar as a biological product that is highly similar to and has no clinically meaningful differences from an existing FDA-approved reference product (1-8).

Regulatory Status

FDA-approved indications:

Neulasta and its biosimilars are leukocyte growth factors indicated: (2-8)

- To decrease the incidence of infection, as manifested by febrile neutropenia, in patients with non-myeloid malignancies receiving myelosuppressive anti-cancer drugs associated with a clinically significant incidence of febrile neutropenia

Neulasta is indicated: (2)

- To increase survival in patients acutely exposed to myelosuppressive doses of radiation

Neulasta and its biosimilars are not indicated for the mobilization of peripheral blood progenitor cells for hematopoietic stem cell transplantation (2-8).

Related policies

Leukine, Neupogen, Rolvedon

Policy

This policy statement applies to clinical review performed for pre-service (Prior Approval, Precertification, Advanced Benefit Determination, etc.) and/or post-service claims.

Neulasta and its biosimilars may be considered **medically necessary** if the conditions indicated below are met.

Neulasta and its biosimilars may be considered **investigational** for all other indications.

Prior-Approval Requirements

Diagnoses

Patient must have **ONE** the following:

1. Prophylaxis for chemotherapy induced febrile neutropenia
2. Treatment of chemotherapy induced febrile neutropenia

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3. Acute radiation syndrome

AND ALL of the following for **ALL** diagnoses:

- a. **NOT** used in combination with another granulocyte colony-stimulating factor (G-CSF)
- b. **Non-preferred medications only:** Inadequate treatment response, intolerance, or contraindication to **ONE** of the preferred products (Fulphila, Udenyca, Udenyca Onbody)

Prior – Approval *Renewal* Requirements

Diagnoses

Patient must have **ONE** the following:

1. Prophylaxis for chemotherapy induced febrile neutropenia
2. Treatment of chemotherapy induced febrile neutropenia
3. Acute radiation syndrome

AND the following for **ALL** diagnoses:

- a. **NOT** used in combination with another granulocyte colony-stimulating factor (G-CSF)

Policy Guidelines

Pre - PA Allowance

None

Prior - Approval Limits

Duration 6 months

Prior – Approval *Renewal* Limits

Same as above

Rationale

Summary

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Neutropenia occurs when an individual has an abnormally low level of neutrophils, a type of white blood cells (WBCs) important in fighting off infections. Neutropenia and its complications, including febrile neutropenia and infection, remain major toxicities associated with myelosuppressive systemic cancer chemotherapy. Colony stimulating factors are medications used to stimulate the production of neutrophils. Neulasta (pegfilgrastim) and its biosimilars are granulocyte colony-stimulating factors (G-CSF) that act on hematopoietic cells by binding to specific cell surface receptors, thereby stimulating proliferation, differentiation, commitment, and end cell functional activation. Fulphila, Fylnetra, Nyvepria, Udenyca, and Ziextenzo are biosimilars to Neulasta. The FDA defines biosimilar as a biological product that is highly similar to and has no clinically meaningful differences from an existing FDA-approved reference product (1-8).

Prior authorization is required to ensure the safe, clinically appropriate, and cost-effective use of Neulasta and its biosimilars while maintaining optimal therapeutic outcomes.

References

1. NCCN Clinical Practice Guidelines in Oncology® Hematopoietic Growth Factors 2026. National Comprehensive Cancer Network, Inc. Accessed on May 1, 2026.
2. Neulasta [package insert]. Thousand Oaks, CA: Amgen Inc.; August 2025.
3. Fulphila [package insert]. Cambridge, MA: Biocon Biologics Inc.; June 2023.
4. Fylnetra [package insert]. Piscataway, NJ: Amneal Pharmaceuticals LLC; April 2025.
5. Nyvepria [package insert]. New York, NY: Pfizer Inc.; March 2023.
6. Stimufend [package insert]. Lake Zurich, IL: Fresenius Kabi USA, LLC; September 2023.
7. Udenyca [package insert]. Raleigh, NC: Accord BioPharma, Inc.; July 2025.
8. Ziextenzo [package insert]. Princeton, NJ: Sandoz Inc.; February 2024.

Policy History

Date	Reason
July 2010	ICD-9 code was removed for myelosuppressive chemotherapy, to decrease the incidence of infection as manifested by febrile neutropenia (various), bone marrow transplantation (996.85), peripheral blood progenitor cell collection (various), acceleration of myeloid recovery in patients with non-Hodgkin's lymphoma, ALL or Hodgkin's disease undergoing bone marrow transplantation (various), induction chemotherapy in acute myelogenous leukemia (various), mobilization and following transplantation of autologous PBPC (various), myeloid

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	reconstitution after allogenic bone marrow transplantation (various), severe chronic neutropenia (various) and bone marrow transplantation failure or engraftment delay (996.0-996.5). ICD-9 code was updated for bone marrow transplantation failure or engraftment delay (996.82). ICD-10 code was added for bone marrow transplantation failure or engraftment delay (T86.02).
November 2010	Separation of colony stimulating factors to improve functionality and workflow; remove non-FDA approved indications (including ICD-9 and 10 codes) as follows: Myelodysplastic Syndrome (MDS), Myeloid engraftment following bone marrow transplantation, Myeloid engraftment following hematopoietic stem cell transplantation, Congenital, Cyclic, or Idiopathic Neutropenia, Neutropenia associated with AIDS treatment, and Peripheral progenitor cell yield.
September 2011	Separation of the colony stimulating agents' criterion; Neulasta is not FDA approved for the same indications as Leukine and Neupogen. Removal of ICD-9 and 10 codes due to lack of specificity.
December 2011	Aligned with Medical Policy
December 2012	Annual Review-editorial updates
March 2014	Annual review and decreased approval and renewal limits to 6 months
March 2015	Annual editorial review and reference update Addition of not used in combination with another granulocyte colony-stimulating factor (G-CSF)
December 2015	Addition of new indication acute radiation syndrome
March 2016	Annual editorial review Policy number changed from 5.10.09 to 5.85.09
December 2016	Annual editorial review and reference update
September 2017	Annual review and reference update
July 2018	Addition of Fulphila biosimilar to criteria
September 2018	Annual review Addition of off-label indications to Fulphila per SME
November 2018	Annual review and reference update. Addition of Udenyca biosimilar to criteria
March 2019	Annual review. Revised regulatory status section to separate indications based on medication per SME
December 2019	Annual review. Addition of requirement to trial preferred products. Addition of Ziextenzo biosimilar to criteria. Renamed policy Neulasta Fulphila Udenyca Ziextenzo
March 2020	Annual review and reference update

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July 2020	Addition of Nyvepria biosimilar
September 2020	Annual review
December 2020	Annual review and reference update. Added Ziextenzo and Nyvepria as preferred products
March 2021	Annual editorial review and reference update. Revised background and summary sections. Clarification added to the t/f preferred products requirement indicating that it only applies to claims adjudicated through the pharmacy benefit
June 2021	Annual review and reference update
June 2022	Annual review and reference update. Addition of biosimilar Fylnetra to policy as preferred product
September 2022	Annual review. Addition of biosimilar Stimufend to policy as preferred product
March 2023	Annual review and reference update
June 2023	Annual review and reference update
December 2023	Annual review and reference update. Per FEP, changed preferred products to Neulasta, Neulasta Onpro, and Udenyca. Also removed Medex requirements. Added t/f requirement of ONE preferred agent to initiation
June 2024	Annual editorial review and reference update. Addition of Udenyca Onbody to policy as preferred product
September 2024	Annual editorial review and reference update. Per FEP, added Fulphila as a preferred product and removed Neulasta/Neulasta Onpro as preferred products for 2025
March 2025	Annual review and reference update
June 2025	Annual review and reference update
June 2026	Annual review and reference update

Keywords

This policy was approved by the FEP® Pharmacy and Medical Policy Committee on June 11, 2026 and is effective on July 1, 2026.