

| Field Name | Field Description |
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| Prior Authorization Group Description | White Blood Cell Stimulators |
| Drugs | <p><u>FORMULARY STATUS</u></p> <p><u>Short-Acting G-CSFs:</u> Nivestym (filgrastim-aafi) – PREFERRED AGENT Granix (TBO-filgrastim) Neupogen (filgrastim) Releuko (filgrastim-ayow) Zarxio (filgrastim-sndz) Or any newly market agent</p> <p><u>Long-Acting G-CSFs:</u> Ziextenzo (pegfilgrastim-bmez) – PREFERRED AGENT Fylnetra (pegfilgrastim-pbbk) – PREFERRED AGENT Nyvepria (pegfilgrastim-apgf) Fulphila (pegfilgrastim-jmdb) Udenyca (pegfilgrastim-cbqv) Neulasta (pegfilgrastim) Neulasta Onpro (pegfilgrastim) Rolvedon (eflapegrastim-xnst) Stimufend (pegfilgrastim-fpgk) Or any newly market agent</p> <p><u>Additional Agents:</u> Mozobil (Plerixafor) Leukine (Sargramostim) Or any newly marketed agent</p> |
| Covered Uses | Medically accepted indications are defined using the following sources: the Food and Drug Administration (FDA), Micromedex, American Hospital Formulary Service (AHFS), United States Pharmacopeia Drug Information for the Healthcare Professional (USPDI), the Drug Package Insert (PPI), or disease state specific standard of care guidelines. |
| Exclusion Criteria | N/A |
| Required Medical Information | See “Other Criteria” |
| Age Restrictions | N/A |
| Prescriber Restrictions | Prescriber must be a hematologist, an oncologist, or an infection disease specialist |
| Coverage Duration | Initial authorization requests for all indications will be approved for 12 weeks. Reauthorization requests for all indications, with the exception of chronic neutropenia, will be approved for 12 weeks. Reauthorization requests for chronic neutropenia will be approved for 24 weeks. |
| Other Criteria | <u>Initial Authorization:</u> |

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| <p>Revision/Review Date: 1/2023</p> | <ul style="list-style-type: none"> • The drug is being used for an appropriate indication at an appropriate dose per “Covered Uses”. • <i>For ALL requests for treatment or prophylaxis of febrile neutropenia:</i> Documentation of the patient’s absolute neutrophil count (ANC) within the last 30 day has been provided. <p><u>Short-Acting G-CSFs:</u></p> <ul style="list-style-type: none"> • <i>For all requests for non-preferred agents:</i> The patient has a documented treatment failure {i.e. failure to reach and/or maintain target ANC, prolonged febrile neutropenia, unplanned hospitalization or infection requiring prolonged anti-infective use} with an adequate trial (including dates, doses of therapy) of Nivestym or has another documented medical reason (intolerance, hypersensitivity, stem cell collection, etc.) for not using Nivestym to treat their medical condition. <p><u>Long-Acting G-CSFs:</u></p> <ul style="list-style-type: none"> • <i>For Fulphila, Udenyca, Nyvepria, Rolvedon, or Stimufend requests:</i> The patient has a documented treatment failure (i.e. failure to reach and/or maintain target ANC, prolonged febrile neutropenia or infection requiring prolonged anti-infection use) with the use of Ziextenzo or Fylnetra, or has a documented medical reason (intolerance, hypersensitivity, stem cell collection, etc.) for not using Ziextenzo or Fylnetra • <i>For Neulasta or Neulasta Onpro requests:</i> The patient has a documented treatment failure (i.e. failure to reach and/or maintain target ANC, prolonged febrile neutropenia or infection requiring prolonged anti-infection use) with the use of Ziextenzo AND Fylnetra PLUS either Fulphila, Udenyca, or Nyvepria or has a documented medical reason (intolerance, hypersensitivity, stem cell collection, etc.) for not using these therapies <p><u>Additional Agents:</u></p> <ul style="list-style-type: none"> • <i>For Leukine requests:</i> Documentation is submitted of the patient’s current diagnosis, current body weight, body surface area (within 30 days of the request). • <i>For Mozobil requests:</i> Documentation is submitted of the patient’s current diagnosis, current body weight, and that the patient is using Mozobil in combination with a granulocyte-colony stimulating factor (G-CSF) agent (e.g. Zarxio, Nivestym) <p>Medical Director/clinical reviewer must override criteria when, in his/her professional judgement, the requested item is medically necessary.</p> |
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