

MEDICARE FORM Filgrastim Precertification Request (Granix<sup>®</sup>, Leukine<sup>®</sup>, Neupogen<sup>®</sup>, Nivestym<sup>®</sup>, Nypozi<sup>™</sup>, Releuko<sup>®</sup>, Zarxio<sup>®</sup>) Page 1 of 6

(All fields must be completed and legible for precertification review.)

For Medicare Advantage Part B: For other lines of business: Please use commercial form.

Note: Granix, Leukine, Neupogen, Nivestym, Nypozi, and Releuko are non-preferred. The preferred product is Zarxio (Neupogen biosimilar). Zarxio does not require precertification

Would you like to use electronic prior authorization? Consider using **Availity**, our electronic prior authorization portal. Learn more about **Availity** from the links in the table below.

For phone or fax requests, refer to the table below for routing information. To determine which box to use, refer to the patient's Aetna ID card. State specific special needs and Medicare-Medicaid Plans may be designated on the front of the ID card or in the website URL on the back of the card. If you don't see your specific plan listed, call the number on the back of the member's ID card to confirm routing information.

|                        | a Medicare Advantage and Allina Health Aetna Medicare members send request to:                                     |
|------------------------|--|
|                        | <u>1-866-503-0857</u> (TTY: <u>711</u> )   |
| Fax:                   | <u>1-844-268-7263</u>  |
| Availity:              | https://www.aetna.com/health-care-professionals/resource-center/availity.html                                      |
| For Aetna<br>send requ | a Medicare Advantage <b>Virginia Dual Eligible Special Needs Plans</b> (HMO D-SNP)<br>uest to:                     |
| Phone:                 | <u>1-855-463-0933</u>  |
| Fax:                   | <u>1-833-280-5224</u>  |
| Availity:              | https://www.aetnabetterhealth.com/virginia-hmosnp/providers/portal   |
|                        | a Assure Premier Plus Medicare Advantage <b>New Jersey Dual Eligible Special Needs Plans</b> SNP) send request to: |
| Phone:                 | <u>1-844-362-0934</u>  |
| Fax:                   | <u>1-833-322-0034</u>  |
| Availity:              | https://www.aetnabetterhealth.com/new-jersey-hmosnp/providers/portal.html  |
| For Aetna              | a Better Health of Illinois Premier Medicare Medicaid Plan (MMP) send request to:                                  |
| Phone:                 | <u>1-866-600-2139</u>  |
| FAX:                   | <u>1-855-320-8445</u>  |
| Availity:              | https://www.aetnabetterhealth.com/illinois/providers/portal  |
| For Aetna              | a Better Health of Ohio Premier Medicare Medicaid Plan (MMP) send request to:                                      |
| Phone:                 | 1-855-364-0974   |
| Fax:                   | 1-855-734-9389   |
| Availity:              | https://www.aetnabetterhealth.com/ohio/providers/portal  |
| For Aetna              | a Better Health of Michigan Premier Medicare Medicaid Plan (MMP) send request to:                                  |
| Phone:                 | <u>1-855-676-5772</u>  |
| Fax:                   | 1-844-241-2495   |
| Availity.              | https://www.aetnabetterhealth.com/michigan/providers/portal.html   |

|  |  |   |  | For Medicare Advantage Part B:<br>For other lines of business:<br>Please use commercial form.<br>Note: Granix, Leukine,<br>Neupogen, Nivestym, Nypozi,<br>and Releuko are non-preferred.<br>The preferred product is Zarxio<br>(Neupogen biosimilar). Zarxio<br>does not require precertification. |                                    |
|--|--|---|--|--|------------------------------------|
| A. PATIENT INFORMATION   |  |   |  |  |                                    |
| First Name:  | Last Name:   |   |  | DOB:   |                                    |
| Address:   | City:  |   |  | State:   | ZIP:                               |
| Home Phone: Work Phone:  | Cell Phor  | ne:   | Email:   | :  |                                    |
| Patient Current Weight: lbs or kgs Pat   | tient Height: inches d   | or cms Al   | lleraies:  |  |                                    |
| B. INSURANCE INFORMATION   |  |   |  |  |                                    |
| Aetna Member ID #:   | Does patient have other  | coverage?   | Yes 🗌 No   |  |                                    |
| Group #:   | If yes, provide ID#:   |   |  |  |                                    |
| Insured:   | Insured:   |   |  |  |                                    |
| C. PRESCRIBER INFORMATION  |  |   |  |  |                                    |
| First Name:  | Last Name:   |   | (Check one):   | 🗌 M.D. 🗌   | D.O. 🗌 N.P. 🗌 P.A.                 |
| Address:   | City:  |   |  | State:   | ZIP:                               |
| Phone: Fax:  | St Lic #:  | NPI #:  | DEA #:   | .1   | UPIN:                              |
| Provider Email: O  | ffice Contact Name:  |   | Phone:   |  | I                                  |
| D. DISPENSING PROVIDER/ADMINISTRATION INF  | ORMATION   |   |  |  |                                    |
| □ Self-administered       □ Physician's Office         □ Outpatient Infusion Center       Phone:         □ Center Name:  | Directions for Use<br>Directions for Use   | Name:   | harmacy  | Fax:NPI: _   |                                    |
| G. CLINICAL INFORMATION – Required clinical info   |  |   | all precertificatio  | n requests   |                                    |
| For All requests (clinical documentation required for a  | ·  | in its <u>entirety</u> for a  | all precertificatio  | in requests.   |                                    |
| Please indicate the patient's absolute neutrophil count:<br>☐ Yes ☐ No Does the patient have a nadir count that<br>Nivestym (filgrastim-aafi), Nypozi (filgra<br>☐ Yes ☐ No Is the requested dose less than 180 mcg<br>☐ Yes ☐ No Is the requested dose less than 180 mcg<br>☐ Yes ☐ No Has the patient tried Z<br>☐ Yes ☐ No Does | mm <sup>3</sup> Date obtained:<br>requires an immediate need t<br>istim-txid), Releuko (filgrastim<br>g (0.3 mL)?<br>arxio (filgrastim-sndz)?<br>the patient have a contraindic<br>′es ☐ No Is the patient com<br>this medication to<br>gramostim), Neupogen (filgras<br>rastim-sndz) be used with and | for Granix (tbo-filgr<br>n-ayow), or Zarxio (<br>cation to Zarxio (filg<br>ppleting an existing<br>p remain unchange<br>stim), Nivestym (filg<br>other colony stimula | (filgrastim-sndz)?<br>grastim-sndz)?<br>g chemotherapy re<br>d?<br>grastim-aafi), Nyp<br>ating factor? | egimen that re<br>bozi (filgrastir   | equires current use of<br>m-txid), |

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| Patient First Name   | Patient Last Name   | Patient Phone   | Patient DOB                      |  |
|--|---|---|----------------------------------|--|
|  |   |   |                                  |  |
| G. CLINICAL INFORMATION (continued   | d) – Required clinical information must be comp   | leted in its <u>entirety</u> for all precertifica         | tion requests.                   |  |
| For All requests (clinical documentation   | n required for all requests):   |   |                                  |  |
| Releuko (filgrastim-ayow)  | ), Leukine (sargramostim), Neupogen (filgrastim<br>or Zarxio (filgrastim-sndz) be used in the same  | chemotherapy cycle as another cold                        |                                  |  |
|  | ceiving concomitant chemotherapy and radiatior  |   |                                  |  |
|  | ),Leukine (sargramostim), Neupogen (filgrastim)<br>or Zarxio (filgrastim-sndz) be used within 7 day |   | filgrastim-txid),                |  |
| For Initiation requests:   |   |   |                                  |  |
| Zarxio does not require precertification   |   |   | Neupogen biosimilar).            |  |
|  | therapy with the requested product within the la  | st 365 days?  |                                  |  |
| Yes No Has the patient had a trial   |   |   |                                  |  |
| Please describe the nati   | ure of the failure of Zarxio  |   |                                  |  |
| $\square$ Yes $\square$ No Has the patient had an ad   | lverse reaction to Zarxio (filgrastim-sndz)?  |   |                                  |  |
|  | s adverse reaction to Zarxio?   |   |                                  |  |
|  | ure of the adverse reaction to Zarxio   |   |                                  |  |
|  | ations or other medical reason(s) that the patier   |   |                                  |  |
|  |   | (3)   |                                  |  |
|  | olid tumor or non-myeloid malignancy and will r<br>brile neutropenia for primary or secondary prop  |   | apy associated with a clinically |  |
| Acute myeloid leukemia   |   |   |                                  |  |
| Yes No Is the patient receiving  | ng induction chemotherapy?  |   |                                  |  |
| $\square$ Please indicate the  | regimen:<br>ng consolidation chemotherapy?  |   |                                  |  |
| $\rightarrow$ Please indicate the  | regimen:  |   |                                  |  |
|  | tation [to mobilize peripheral-blood progenit<br>ant and date received:                             |   | 1                                |  |
| Advanced HIV infection   |   |   |                                  |  |
|  | e anti-retroviral medication the patient is receivi   | ng:   |                                  |  |
| Yes □ No Is the patient neutrop  | penic?  |   |                                  |  |
| Bone Marrow Transplantation  |   | -   |                                  |  |
|  | ve a documented diagnosis of non-myeloid mali   |   |                                  |  |
| Yes No is the medication be  | ing requested to reduce the duration of neutrop   | enia and neutropenia-related infectio                     | us complications?                |  |
|  | e treatment will be followed by: Autologous   | oone marrow transplantation<br>one marrow transplantation |                                  |  |
| Congenital, cyclic or idiopathic neut  | ropenia   |   |                                  |  |
| Please identify which documented type of neutropenia that patient has:      Congenital neutropenia      Cyclic neutropenia      idiopathic neutropenia     Yes      No Is the patient currently symptomatic? |   |   |                                  |  |
| Drug- induced agranulocytosis  |   |   |                                  |  |
| ☐ Yes ☐ No Is the agranulocytosis caused by chemotherapy?<br>→ Please provide the medication(s) that caused the agranulocytosis:   |   |   |                                  |  |
| Hematopoietic Subsyndrome of Acute Radiation Syndrome (H-ARS)  |   |   |                                  |  |
| Yes No Is the medication being requested for the treatment of radiation-induced myelosuppression following a radiological/nuclear incident?  |   |   |                                  |  |
| ☐ Intermittent use in patients with my   |   | , 5   | -                                |  |
| ☐ Yes ☐ No Does the patient have symptomatic anemia?   |   |   |                                  |  |
| 🖵 Yes 🔲 No Has the patient been tested for 5q gene deletion?   |   |   |                                  |  |
| Please indicate the result of the test and date obtained: Date obtained: Date obtained:/   |   |   |                                  |  |
| ☐ Yes ☐ No Does the patient present with other cytogenetic abnormalities?<br>☐ Yes ☐ No Has a serum erythropoietin test been completed?  |   |   |                                  |  |
| Please indicate the  | result of the test and date obtained:   | Date  | obtained: / /                    |  |

Continued on next page

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## MEDICARE FORM Filgrastim Precertification Request (Granix<sup>®</sup>, Leukine<sup>®</sup>, Neupogen<sup>®</sup>, Nivestym<sup>®</sup>, Nypozi<sup>™</sup>, Releuko<sup>®</sup>, Zarxio<sup>®</sup>)

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| Patient First Name                                 | Patient Last Name   | Patient Phone                                | Patient DOB                      |  |  |
|--|---|--|----------------------------------|--|--|
|  |   |  |                                  |  |  |
|  | ued) – Required clinical information must be con  | npleted in its <u>entirety</u> for all prece | ertification requests.           |  |  |
| ■ Neuroblastoma<br>■ Yes ■ No Is the patient's dis | soaso considered high risk?   |  |                                  |  |  |
| Yes No Will the requested                          | I medication be used in combination with ALL of oleukin), isotretinoin (13-cis-retinoic acid)?                    | the following medications: dinutu            | ıximab (Unituxin), interleukin-2 |  |  |
|  | ] No Will the requested medication be used in   | combination with Naxitamab-gog               | k (Danvelza)?                    |  |  |
| Primary prophylaxis of neutropenia                 |   | 515  |                                  |  |  |
|  | nave a documented diagnosis of non-myeloid ma   | alignancy?                                   |                                  |  |  |
|  | eiving myelosuppressive chemotherapy?   |  |                                  |  |  |
|  | the type of cancer the patient is being treated for   |  |                                  |  |  |
|  | e exact chemotherapy regimen patient is currentl<br>ected percentage of febrile neutropenia incidence             |  | -2                               |  |  |
|  | sk)   |  | 12                               |  |  |
|  | sidered to be at high risk for chemotherapy-induc   |  | complications?                   |  |  |
|  | which of the following reasons that categorizes th  |  |                                  |  |  |
|  | ons 🔲 Age greater than or equal to 65 years   |  |                                  |  |  |
|  | <i>i</i> nvolvement by tumor producing cytopenias   |  |                                  |  |  |
|  | ance status   |  |                                  |  |  |
| Other serious                                      | s co-morbidities: 🔲 Cardiovascular disease [  | HIV Infection Liver dysfund                  | ction [] Renal dysfunction       |  |  |
| Secondary prophylaxis of neutro                    |   |  |                                  |  |  |
|  | nave a documented diagnosis of non-myeloid ma   | alignancy?                                   |                                  |  |  |
|  | perience a febrile neutropenic complication from  |  |                                  |  |  |
|  | the neutropenic complication the patient experies   | nced from the prior cycle of chem            | notherapy:                       |  |  |
| Neutropenic cor<br>Please indicate                 | the prior cycle of chemotherapy that the patient r  | eceived with the neutropenic cor             | nolication.                      |  |  |
|  | perience a dose-limiting neutropenic event (a na  |  |                                  |  |  |
|  | om a prior cycle of similar chemotherapy?   |  |                                  |  |  |
|  | Was the patient treated with the same dose and  |  | cle?                             |  |  |
|  | Did the patient receive primary prophylaxis again   | nst febrile neutropenia?                     |                                  |  |  |
| Therapeutic use in a high-risk, fe                 | brile neutropenic patient<br>/ing prognostic factors pertains to the patient:                                     |  |                                  |  |  |
|  |   |  |                                  |  |  |
| Being hospit                                       | alized at the time of the development of fever  |  |                                  |  |  |
| Pleas  | e provide date of hospitalization: / /  |  |                                  |  |  |
|  | յal infection<br>le type of fungal infection and date infection օշշւ  | urrod :                                      | Date: / /                        |  |  |
|  | le type of fungal infection and date infection occu   |  |                                  |  |  |
|  | e provide date of pneumonia infection: /  | /  |                                  |  |  |
|  | es of febrile neutropenia   |  |                                  |  |  |
|  | eutropenia<br>s     No   Is the prolonged neutropenia expected  | to lost greater than 10 days?                |                                  |  |  |
|  |   | to last greater than to days?                |                                  |  |  |
|  | Sepsis syndrome   |  |                                  |  |  |
| 🗌 Other  |   |  |                                  |  |  |
|  | e explain:  |  |                                  |  |  |
|  | <u>rrastim-aafi), Releuko (filgrastim-ayow), Zarxi</u>  | <u>o (filgrastim-sndz):</u>                  |                                  |  |  |
| Acute lymphoblastic leukemia (A                    | LL) of chemotherapy been completed?   |  |                                  |  |  |
| ☐ Yes ☐ No Is this the initial in                  |   |  |                                  |  |  |
|  | st-remission course of chemotherapy?  |  |                                  |  |  |
| Please provide the chemotherapy                    | regimen and date started: Regimen:  |  | Date started: / /                |  |  |
| C Acute myeloid leukemia                           |   |  |                                  |  |  |
| Yes No Is the patient rece                         |   |  |                                  |  |  |
| Yes No Is the patient rece                         | eiving consolidation chemotherapy?  |  |                                  |  |  |
| Yes No Is the patient rece                         | Please indicate the regimen: ☐ Yes ☐ No Is the patient receiving chemotherapy for relapsed or refractory disease? |  |                                  |  |  |
| Relapsed dis                                       | → ☐ Relapsed disease ☐ Refractory disease   |  |                                  |  |  |
| Please indicate                                    | the regimen:  |  |                                  |  |  |

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| Patient First Name                                     | Patient Last Name  | Patient Phone                                       | Patient DOB                     |
|--|--|---|---------------------------------|
|  |  |   |                                 |
| G. CLINICAL INFORMATION (continu                       | red) – Required clinical information must be com   | npleted in its <u>entirety</u> for all precertifica | tion requests.                  |
|  | antation [to mobilize peripheral-blood progen  |   |                                 |
|  | plant and date received:  Autologous   |   | 1                               |
| Advanced HIV infection                                 | plant and date received.  Autologous Alto  | geneic Date of transplant. /                        |                                 |
|  | ive anti-retroviral medication the patient is recei  | vina  |                                 |
| $\square$ Yes $\square$ No Is the patient neutr        |  | ving  |                                 |
|  | openie   |   |                                 |
| Bone Marrow Transplantation                            | ave a documented diagnosis of non-myeloid ma   | lignonov2   |                                 |
|  | being requested to reduce the duration of neutro   |   | ue complications?               |
|  | rgoing myeloablative chemotherapy?   | penia and neutropenia-related infectio              | us complications?               |
|  | the treatment will be followed by:  Autologous   | s hone marrow transplantation                       |                                 |
|  |  | bone marrow transplantation                         |                                 |
|  |  |   |                                 |
| Congenital, cyclic or idiopathic ne                    |  |   |                                 |
|  | type of neutropenia that patient has: 🗌 congeni  | ital neutropenia. 🔲 cyclic neutropenia              | idiopathic neutropenia          |
| Yes No Is the patient curre                            |  |   |                                 |
|  | astim), Leukine (sargramostim), Neupogen (filgr  | astim) Nivestym (filgrastim-aafi) Nype              | ozi (filgrastim-txid)           |
| Releuko (filgrastim                                    | n-ayow), or Zarxio (filgrastim-sndz) being reques  | ted for chronic administration to reduc             | e the incidence and duration of |
|  | penia (e.g., fever, infections, oropharyngeal ulc  |   |                                 |
| Chronic Myeloid Leukemia                               |  |   |                                 |
| Yes No Does the patient ha                             | ave resistant neutropenia?   |   |                                 |
|  | secondary to use of any of the following medica  | itions?   |                                 |
|  | inib) 🔲 Gleevec (imatinib) 🔲 Iclusig (ponatir  |   | na (nilotinib)                  |
| Drug- induced agranulocytosis                          | , _ ,  | , , ,   | · · · ·                         |
| Yes No Is the agranulocyto                             | osis caused by chemotherapy?   |   |                                 |
|  | ne medication(s) that caused the agranulocytosis   | S:  |                                 |
| ☐ Glycogen storage disease (GSD) t                     |  |   |                                 |
| Yes No Does the patient ha                             |  |   |                                 |
| ☐ Hairy Cell Leukemia                                  | •  |   |                                 |
|  | ave clinical evidence of neutropenic fever follow  | ing chemotherapy?                                   |                                 |
| ☐ Increase dose intensity chemother                    |  | 5 13  |                                 |
|  | g treated in a setting in which clinical research d  | emonstrates that dose-intensive thera               | by produces improvement in      |
| disease control?                                       | ,  | ·   |                                 |
|  | ne type of cancer the patient is being treated for   |   |                                 |
|  | exact chemotherapy regimen patient is currently  |   |                                 |
|  | f febrile neutropenia incidence from the chemot  |   |                                 |
|  | k) ☐ 10-19% (Intermediate risk) ☐ 20% or g   |   |                                 |
|  | idered to be at high risk for chemotherapy-induc   |   | lications?                      |
|  | which of the following reasons that categorizes the  |   |                                 |
|  | ns Age greater than or equal to 65 years   |   |                                 |
|  | involvement by tumor producing cytopenias  |   |                                 |
| ·  | ance status 🔲 Previous chemotherapy 🗌 Pre  | evious radiation therapy U Previous                 | episodes of FN                  |
| ☐ Recent surger  |  |   |                                 |
| ☐ Other serious  | co-morbidities:  Cardiovascular disease  |   |                                 |
|  | ☐ Other- Please explain:   |   |                                 |
| ☐ Intermittent use in patients with m                  |  |   |                                 |
| Yes No Does the patient ha                             |  |   |                                 |
| Yes No Has the patient be                              |  | Dete  | abtainadu / /                   |
|  | ne result of the test and date obtained:<br>resent with other cytogenetic abnormalities?         |   | obtailleu. / /                  |
| $\square$ Yes $\square$ No $\square$ Has a serum eryth |  |   |                                 |
|  | ne result of the test and date obtained:   | Date  | obtained / /                    |
|  |  | Date  |                                 |
|  |  |   |                                 |
|  | dence that the patient is being treated with cura  | tive chemotherapy (e.g. (R- CHOP) rit               | tuximab cyclophosphamide        |
| Yes No Is there clinical evidence                      | dence that the patient is being treated with cura stine, prednisone) or more aggressive regimens | tive chemotherapy (e.g. (R- CHOP) rit<br>?          | tuximab, cyclophosphamide,      |

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| Patient First Name                           | Patient Last N  | lame                               | Patient Phone                        | Patient DOB                                   |
|--|---|------------------------------------|--------------------------------------|---|
|  |   |                                    |                                      |   |
| G. CLINICAL INFORM                           | <b>IATION</b> (continued) – Required o                                    | linical information must be comp   | oleted in its <u>entirety</u> for al | precertification requests.                    |
| Primary prophylax                            | kis of neutropenia  |                                    |                                      |   |
|  | Does the patient have a documente   |                                    | gnancy?                              |   |
|  | s the patient receiving myelosuppre                                       |                                    |                                      |   |
| $ \longrightarrow $                          | Please indicate the type of cancer<br>Please enter the exact chemother    |                                    |                                      |   |
| What is the exper                            | cted percentage of febrile neutrope                                       |                                    |                                      |   |
|  | □ 0-9% (Low risk) □ 10-19% (I   |                                    |                                      |   |
|  | s the patient considered to be at his                                     |                                    |                                      | ctious complications?                         |
| $  \longrightarrow$                          | Please indicate which of the follow                                       | ing reasons that categorizes the   | e patient to be at high risk         | c.  |
|  | Active infections Age great   |                                    |                                      |   |
|  |   |                                    |                                      | stent neutropenia 🔲 Poor nutritional status   |
|  | Poor performance status   | Previous chemotherapy              | vious radiation therapy              | Previous episodes of FN                       |
|  | <ul> <li>Recent surgery</li> <li>Other serious co-morbidities:</li> </ul> |                                    | LUN/infaction Diverd                 | vefunction Donal dysfunction                  |
|  |   | Other- Please explain:             |                                      |   |
| Radiation therapy                            |   |                                    |                                      |   |
|  | Are prolonged delays in radiation th                                      | erapy expected due to neutrope     | nia?                                 |   |
| Secondary prophy                             | ylaxis of neutropenia   |                                    |                                      |   |
|  | Does the patient have a documente   | d diagnosis of non-myeloid mal     | gnancy?                              |   |
|  | Did the patient experience a febrile                                      |                                    |                                      |   |
|  | Please indicate the neutropenic co  | emplication the patient experience | ed from the prior cycle o            | f chemotherapy:                               |
|  | Neutropenic complication:<br>Please indicate the prior cycle of c         | homotherapy that the nationt re    | aniwood with the neutrone            | aia complication.                             |
|  | Did the patient experience a dose-li                                      |                                    |                                      |   |
| 1 1  | chemotherapy) from a prior cycle of                                       | • · · ·                            | i of day of treatment cod            |   |
|  | Yes No Was the patient tre  |                                    | hedule planned for curre             | nt cycle?                                     |
| [  | Yes No Did the patient rece   | eive primary prophylaxis against   | febrile neutropenia?                 | -   |
| Therapeutic use in                           | n a high-risk, febrile neutropenic  | patient                            |                                      |   |
|  | hich of the following prognostic fac                                      |                                    |                                      |   |
|  | Age greater than 65 years   |                                    |                                      |   |
| [  | Being hospitalized at the time of   |                                    |                                      |   |
| г  | Please provide date of h  | ospitalization: / /                |                                      |   |
| L  | ☐ Invasive fungal infection<br>└────────────────────────────────────      | fection and date infection occur   | red.                                 | Date: / /                                     |
| Г  | ☐ Pneumonia   |                                    |                                      | Date: /                                       |
|  |   | neumonia infection: /              | 1                                    |   |
| [  | Prior episodes of febrile neutrop   |                                    |                                      |   |
| [  | Prolonged neutropenia   |                                    |                                      |   |
| -  | Yes No Is the p   | rolonged neutropenia expected      | to last greater than 10 da           | ys?   |
|  | Profound neutropenia  |                                    |                                      |   |
|  | ☐ Sepsis syndrome<br>☐ Other  |                                    |                                      |   |
| L  |   |                                    |                                      |   |
| ☐ Treatment of high                          | Please explain:   |                                    |                                      |   |
| Treatment for radi                           |   |                                    |                                      |   |
|  | ne radiation dose that caused the ir                                      | ijury: grays (Gy)                  |                                      |   |
| For Continuation requ                        |   |                                    |                                      |   |
| ☐ Yes ☐ No Is this                           | continuation request a result of the                                      | e patient receiving samples of G   | ranix (tbo-filgrastim), Leu          | kine (sargramostim), Neupogen (filgrastim),   |
| Nivest                                       | tym (filgrastim-aafi), Nypozi (filgras                                    | stim-txid), Releuko (filgrastim-a  | /ow), or Zarxio (filgrastim          | -sndz)?                                       |
|  |   |                                    |                                      | (filgrastim), Nivestym (filgrastim-aafi),     |
| Nypoz  | zi (filgrastim-txid), Releuko (filgras                                    | tim-ayow), or Zarxio (filgrastim-  | sndz) therapy?                       |   |
| H. ACKNOWLEDGE                               | MENT  |                                    |                                      |   |
| Request Completed                            | By (Signature Required):  |                                    |                                      | Date: /                                       |
| Any person who know                          |   |                                    |                                      | with the intent to injure, defraud or deceive |
| any insurance compa                          | any by providing materially false   | information or conceals mater      | ial information for the p            | urpose of misleading, commits a fraudulent    |
| insurance act, which                         | is a crime and subjects such per  | son to criminal and civil penalt   | ies.                                 |   |
| The sub-sub-sub-sub-sub-sub-sub-sub-sub-sub- | additional information or algoritics                                      |                                    |                                      |   |

The plan may request additional information or clarification, if needed, to evaluate requests. GR-69389-3 (3-25)