



## Drug Coverage Policy

Effective Date ..... 5/1/2026

Coverage Policy Number ..... 1403

Policy Title.....Oncology Medications

# Oncology Medications

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### **INSTRUCTIONS FOR USE**

*The following Coverage Policy applies to health benefit plans administered by Cigna Companies. Certain Cigna Companies and/or lines of business only provide utilization review services to clients and do not make coverage determinations. References to standard benefit plan language and coverage determinations do not apply to those clients. Coverage Policies are intended to provide guidance in interpreting certain standard benefit plans administered by Cigna Companies. Please note, the terms of a customer's particular benefit plan document [Group Service Agreement, Evidence of Coverage, Certificate of Coverage, Summary Plan Description (SPD) or similar plan document] may differ significantly from the standard benefit plans upon which these Coverage Policies are based. For example, a customer's benefit plan document may contain a specific exclusion related to a topic addressed in a Coverage Policy. In the event of a conflict, a customer's benefit plan document always supersedes the information in the Coverage Policies. In the absence of a controlling federal or state coverage mandate, benefits are ultimately determined by the terms of the applicable benefit plan document. Coverage determinations in each specific instance require consideration of 1) the terms of the applicable benefit plan document in effect on the date of service; 2) any applicable laws/regulations; 3) any relevant collateral source materials including Coverage Policies and; 4) the specific facts of the particular situation. Each coverage request should be reviewed on its own merits. Medical directors are expected to exercise clinical judgment where appropriate and have discretion in making individual coverage determinations. Where coverage for care or services does not depend on specific circumstances, reimbursement will only be provided if a requested service(s) is submitted in accordance with the relevant criteria outlined in the applicable Coverage Policy, including covered diagnosis and/or procedure code(s). Reimbursement is not allowed for services when billed for conditions or diagnoses that are not covered under this Coverage Policy (see "Coding Information" below). When billing, providers must use the most appropriate codes as of the effective date of the submission. Claims submitted for services that are not accompanied by covered code(s) under the applicable Coverage Policy will be denied as not covered. Coverage Policies relate exclusively to the administration of health benefit plans. Coverage Policies are not recommendations for treatment and should never be used as treatment guidelines. In certain markets, delegated vendor guidelines may be used to support medical necessity and other coverage determinations.*

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**This coverage policy addresses medications used for the primary treatment of cancer.**

The use of oncology agents for non-oncology uses are addressed in separate coverage policies.

For a list of medications included in the oncology medications coverage policy, refer to the [Cigna - Oncology Medication and Code List](#)

All products are approved for a duration of up to 12 months unless otherwise noted.

**Oncology Medications for any other use is considered not medically necessary. Criteria will be updated as new published data are available.**

Oncology Medications for uses that are an NCCN category 3 recommendation (unless the use is approved by the FDA) are not covered because they are considered not medically necessary.

## Coverage Policy

**Documentation:** Documentation is required where noted in the criteria as **[documentation required]**. Documentation may include, but is not limited to, chart notes, laboratory tests, claims records, prescription receipts and/or other information. All documentation must include patient-specific identifying information.

**Oncology Medications are considered medically necessary when BOTH of the following are met:**

1. **ONE** of the following criteria are met:
  - a. Use is an approved indication by the Food and Drug Administration (FDA)
  - b. Use is a category 1, 2A, or 2B recommendation by the National Comprehensive Cancer Network (NCCN) Clinical Practice Guidelines in Oncology (NCCN Guidelines®) or its derivative information product, The NCCN Drugs & Biologics Compendium (NCCN Compendium®)
  - c. For **Pediatric Oncology** use, **ALL** of the following criteria are met:
    - i. The drug is FDA approved for at least one indication
    - ii. The drug has not been contraindicated or not recommended by the FDA for the off-label use
    - iii. Supported by **ONE** of the following:
      1. Compendia recognized by federal Centers for Medicare and Medicare Services (CMS) as part of an anticancer chemotherapeutic regimen (AHFS, Clinical Pharmacology, DrugDex, etc.)
      2. Results of at least two different controlled clinical studies published in peer reviewed English-language, biomedical journal(s) analyzed as supporting the off-label use where consideration is given to quality of evidence, validity of the data, efficacy, and safety of the drug in specific patient populations and how well the study was designed to assess the intervention, including analysis of baseline patient characteristics, patient withdrawals, and meaningful clinical outcome
      3. Established as standard of care as analyzed in clinical practice guidelines from professional or medical specialty societies, national government supported evidence assessments or guidelines
2. If required, preferred product criteria are met as listed in the below table

Product	Criteria
<b>Abraxane intravenous infusion</b> (paclitaxel albumin-bound)	<p><b><u>Cigna Pathwell Specialty Drug List Plans</u></b></p> <p><b>Abraxane is considered medically necessary when BOTH of the following are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. <b>ONE</b> of the following:           <ol style="list-style-type: none"> <li>a. For <b>Breast Cancer</b>, <b>ONE</b> of the following:               <ol style="list-style-type: none"> <li>i. Patient is currently receiving Abraxane or paclitaxel albumin-bound intravenous infusion</li> <li>ii. Patient has tried paclitaxel intravenous infusion</li> <li>iii. Patient has a hypersensitivity to paclitaxel intravenous infusion or docetaxel intravenous infusion</li> </ol> </li> </ol> </li> </ol>

- iv. Patient had a contraindication to the standard pre-mediations (for example, dexamethasone, ranitidine, famotidine, diphenhydramine)
- b. For **Cervical Cancer**, **ONE** of the following:
  - i. Patient is currently receiving Abraxane or paclitaxel albumin-bound intravenous infusion
  - ii. Patient has tried paclitaxel intravenous infusion
  - iii. Patient has a hypersensitivity to paclitaxel intravenous infusion or docetaxel intravenous infusion
  - iv. Patient had a contraindication to the standard pre-mediations (for example, dexamethasone, ranitidine, famotidine, diphenhydramine)
- c. For **Endometrial Cancer**, **ONE** of the following:
  - i. Patient is currently receiving Abraxane or paclitaxel albumin-bound intravenous infusion
  - ii. Patient has tried paclitaxel intravenous infusion
  - iii. Patient has a hypersensitivity to paclitaxel intravenous infusion or docetaxel intravenous infusion
  - iv. Patient had a contraindication to the standard pre-mediations (for example, dexamethasone, ranitidine, famotidine, diphenhydramine)
- d. For **Melanoma**, **ONE** of the following:
  - i. Patient is currently receiving Abraxane or paclitaxel albumin-bound intravenous infusion
  - ii. Patient has tried paclitaxel intravenous infusion
  - iii. Patient had a hypersensitivity to paclitaxel intravenous infusion or docetaxel intravenous infusion
  - iv. Patient has a contraindication to the standard pre-mediations (for example, dexamethasone, ranitidine, famotidine, diphenhydramine)
- e. For **Non-Small Cell Lung Cancer**, **ONE** of the following:
  - i. Patient is currently receiving Abraxane or paclitaxel albumin-bound intravenous infusion
  - ii. Patient has tried paclitaxel intravenous infusion
  - iii. Patient had a hypersensitivity to paclitaxel intravenous infusion or docetaxel intravenous infusion
  - iv. Patient has a contraindication to the standard pre-mediations (for example, dexamethasone, ranitidine, famotidine, diphenhydramine)
  - v. Abraxane or paclitaxel albumin-bound intravenous infusion is being used as subsequent therapy in patients with advanced or metastatic disease
- f. For **Ovarian Cancer**, **ONE** of the following:
  - i. Patient is currently receiving Abraxane or paclitaxel albumin-bound intravenous infusion
  - ii. Patient has tried paclitaxel intravenous infusion
  - iii. Patient had a hypersensitivity to paclitaxel intravenous infusion or docetaxel intravenous infusion
  - iv. Patient has a contraindication to the standard pre-mediations (for example, dexamethasone, ranitidine, famotidine, diphenhydramine)

	<p>g. <b>All Other Conditions.</b> Approve Abraxane intravenous infusion if the patient meets the Oncology Medications criteria above the table</p>
<p><b>Akeega</b> (niraparib and abiraterone)</p>	<p><b><u>Employer Plans and Individual and Family Plan:</u></b></p> <p><b>Akeega is considered medically necessary when BOTH of the following are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. <b>ONE</b> of the following: <ol style="list-style-type: none"> <li>A. For <b><u>BRCA-mutated Prostate Cancer</u></b>, documentation of <b>ONE</b> of the following: <ol style="list-style-type: none"> <li>1. Trial of, contraindication, or intolerance to <b>Lynparza (olaparib), with or without, generic abiraterone</b> [may require prior authorization]</li> <li>2. Patient has BRCA2-mutated metastatic castration-sensitive prostate cancer</li> <li>3. Patient has already been started on therapy with Akeega</li> </ol> </li> <li>B. <b>All Other Conditions.</b> Approve Akeega if the patient meets the Oncology Medications criteria above the table</li> </ol> </li> </ol>
<p><b>Alunbrig</b> (brigatinib)</p>	<p><b><u>Individual and Family Plans</u></b></p> <p><b>Alunbrig (brigatinib) is considered medically necessary when BOTH of the following are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. <b>ONE</b> of the following: <ol style="list-style-type: none"> <li>A. For <b><u>Non-Small Cell Lung Cancer – Anaplastic Lymphoma Kinase (ALK)-positive</u></b>, documentation of <b>ONE</b> of the following: <ol style="list-style-type: none"> <li>i. Trial of, contraindication, or intolerance to <b>Alecensa (alectinib)</b> [may require prior authorization]</li> <li>ii. Patient has already been started on therapy with Alunbrig</li> </ol> </li> <li>B. <b>All Other Conditions.</b> Approve Alunbrig if the patient meets the Oncology Medications criteria above the table</li> </ol> </li> </ol>
<p><b>Alymys</b> (bevacizumab-maly)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Alymys is considered medically necessary when BOTH of the follow are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Patient meets <b>BOTH</b> of the following (A and B): <ol style="list-style-type: none"> <li>A. Patient has tried <b>BOTH</b> of the following [<b>documentation required</b>]: <ol style="list-style-type: none"> <li>i. <b>Mvasi (bevacizumab-awwb)</b> [may require prior authorization]</li> <li>ii. <b>Zirabev (bevacizumab-bvzr)</b> [may require prior authorization]</li> </ol> </li> <li>B. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction:</li> </ol> </li> </ol>

<p><b>Augtyro</b> (repotrectinib)</p>	<p><b><u>Employer Plans</u></b></p> <p><b>Augtyro is considered medically necessary when BOTH of the follow are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. <b>ONE</b> of the following: <ol style="list-style-type: none"> <li>A. <b>For <u>ROS1-positive non-small cell lung cancer</u>, ONE</b> of the following: <ol style="list-style-type: none"> <li>i. Trial of, contraindication, or intolerance to <b>Rozlytrek</b> (entrectinib)</li> <li>ii. If Augtyro has not been tried previously, approve if the patient has progression on Rozlytrek (entrectinib capsules and pellet packet), Xalkori (crizotinib capsules), or Ibtrozi (talectrectinib capsules)</li> <li>iii. Patient has congestive heart failure</li> <li>iv. According to the prescriber, the patient has a risk of QT prolongation</li> <li>v. Patient has already started on therapy with Augtyro</li> </ol> </li> <li>B. <b>All Other Conditions.</b> Approve Augtyro if the patient meets the Oncology Medications criteria above the table</li> </ol> </li> </ol>
<p><b>Avastin®</b> (bevacizumab)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Avastin is considered medically necessary when BOTH of the follow are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Patient meets <b>BOTH</b> of the following (A and B): <ol style="list-style-type: none"> <li>A. Patient has tried <b>BOTH</b> of the following [<b>documentation required</b>]: <ol style="list-style-type: none"> <li>i. <b>Mvasi</b> (bevacizumab-awwb) [may require prior authorization]</li> <li>ii. <b>Zirabev</b> (bevacizumab-bvzr) [may require prior authorization]</li> </ol> </li> <li>B. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.</li> </ol> </li> </ol>
<p><b>Avgamsi</b> (gemcitabine)</p>	<p><b><u>Employer Plans and Individual and Family Plan</u></b></p> <p><b>Avgamsi (gemcitabine) is considered medically necessary when BOTH of the follow are met (1 and 2):</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Patient meets <b>ONE</b> of the following (A or B): <ol style="list-style-type: none"> <li>A. Patient has tried one generic gemcitabine for injection product</li> <li>B. Patient is unable to obtain gemcitabine for injection</li> </ol> </li> </ol>
<p><b>Besremi</b> (ropeginterferon-alfa-2b-njft)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Besremi (ropeginterferon-alfa-2b-njft) is considered medically necessary when BOTH of the following are met:</b></p>

	<ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met AND</li> <li>2. <b>ONE</b> of the following: <ol style="list-style-type: none"> <li><b>A. For Polycythemia Vera, ONE</b> of the following: <ol style="list-style-type: none"> <li>i. Patient has high risk polycythemia vera and documentation provided that the patient has <b>ONE</b> of the following: <ol style="list-style-type: none"> <li>1. Documentation provided that the patient has tried hydroxyurea</li> <li>2. Documentation provided that the patient is NOT a candidate for hydroxyurea therapy</li> </ol> </li> <li>ii. Documentation provided that the patient has low-risk polycythemia vera</li> <li>iii. Documentation provided that the patient is currently receiving Besremi</li> </ol> </li> <li><b>B. All Other Conditions.</b> Approve Besremi if the patient meets the Oncology Medications criteria above the table</li> </ol> </li> </ol>
<p><b>Beizray</b> (docetaxel albumin-bound intravenous infusion)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b><u>Beizray is considered medically necessary when BOTH of the following are met:</u></b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met AND</li> <li>2. <b>ONE</b> of the following: <ol style="list-style-type: none"> <li>A. Patient has tried generic docetaxel</li> <li>B. Patient with hypersensitivities to polysorbate 80</li> </ol> </li> </ol>
<p><b>Boruzu</b> (bortezomib injection)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b><u>Boruzu (bortezomib injection) is considered medically necessary when BOTH of the following are met:</u></b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met AND</li> <li>2. Patient has tried bortezomib injection (Velcade, generics)</li> </ol>
<p><b>Bosulif</b> (bosutinib tablets and capsules)</p>	<p><b><u>Employer Plans</u></b></p> <p><b><u>Bosulif is considered medically necessary when BOTH of the following are met (1 and 2):</u></b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. <b>ONE</b> of the following (A B, or C): <ol style="list-style-type: none"> <li>A. For <b><u>Chronic Myeloid Leukemia (CML)</u></b>, <b>ONE</b> the following: <ol style="list-style-type: none"> <li>i. Trial of, contraindication, or significant intolerance to <b>TWO</b> of the following: <ol style="list-style-type: none"> <li>1. <b>dasatinib</b></li> <li>2. <b>imatinib</b></li> <li>3. <b>nilotinib hcl</b></li> <li>4. <b>Danziten</b> [may require prior authorization]</li> <li>5. <b>Imkeldi</b> [may require prior authorization]</li> <li>6. <b>Scemblix</b> [may require prior authorization]</li> </ol> <p><u>Note:</u> Prior use of brand Gleevec, Phyrago, Sprycel, or Tasigna counts.</p> </li> <li>ii. Patient is currently receiving therapy with Bosulif</li> <li>iii. Patient meets <b>BOTH</b> of the following:</li> </ol> </li> </ol> </li> </ol>

	<ol style="list-style-type: none"> <li>1. Patient has intermediate- to high-risk chronic phase CML, accelerated phase CML or blast phase CML</li> <li>2. Patient has tried at least two other tyrosine kinase inhibitors for CML  Note: Examples of tyrosine kinase inhibitors include: dasatinib, nilotinib, Iclusig, and Scemblix; OR</li> <li>iv. Patient has a resistance mutation in which imatinib, dasatinib, nilotinib hcl, Danziten, Imkeldi, or Scemblix should not be used</li> </ol> <p><b>B. For Acute Lymphoblastic Leukemia: ONE</b> of the following:</p> <ol style="list-style-type: none"> <li>i. Trial of, contraindication, or significant intolerance to <b>ONE</b> of the following: imatinib or dasatinib; OR  <u>Note:</u> Prior use of Gleevec, Imkeldi, Phyrago, or Sprycel also counts.</li> <li>ii. Patient is currently receiving therapy with Bosulif; OR</li> <li>iii. Patient meets ONE of the following: <ol style="list-style-type: none"> <li>1. Patient has a history of serious, chronic lung disease or has had or is at risk of pleural effusion; OR  <u>Note:</u> Examples of lung diseases include pulmonary arterial hypertension and interstitial pneumonitis.</li> <li>2. Patient is at risk of bleeding; OR  <u>Note:</u> An example of a patient with an increased risk of bleeding is a patient with thrombocytopenia or with medication use that inhibits platelet function or anticoagulants.</li> <li>3. Patient has a prolonged QT interval or is at risk of developing QT interval prolongation; OR</li> </ol> </li> <li>iv. Patient has a mutation in which imatinib or dasatinib should not used,</li> </ol> <p><b>C. All Other Conditions.</b> Approve Bosulif if the patient meets the Oncology Medications criteria above the table</p> <p><b><u>Individual and Family Plans</u></b>  <b>Bosulif is considered medically necessary when BOTH of the following are met (1 and 2):</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. <b>ONE</b> of the following (A, B or C): <ol style="list-style-type: none"> <li>A. For <b><u>Chronic Myeloid Leukemia (CML)</u></b>, <b>ONE</b> the following (i or ii): <ol style="list-style-type: none"> <li>i. Patient meets BOTH of the following (1 and 2): <ol style="list-style-type: none"> <li>1. Patient meets ONE of the following: <ol style="list-style-type: none"> <li>a) Trial of, contraindication, or significant intolerance to imatinib; OR  <u>Note:</u> Prior use of brand Gleevec or Imkeldi also counts.</li> <li>b) Patient has intermediate-to-high risk chronic phase CML, accelerated phase CML, or blast phase CML; OR</li> <li>c) Patient has tried at least one other tyrosine kinase inhibitor for CML; OR</li> </ol> </li> </ol> </li> </ol> </li> </ol> </li> </ol>
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	<p><u>Note:</u> Examples of tyrosine kinase inhibitors include: dasatinib, nilotinib, Iclusig, and Scemblix.</p> <p>d) Patient has a resistance mutation in which imatinib should not be used; AND</p> <p>2. Patient meets ONE of the following:</p> <p>a) Trial of, contraindication, or significant intolerance to dasatinib; OR</p> <p>b) Patient has a history of a serious, chronic lung disease or has had or is at risk of pleural effusion; OR</p> <p><u>Note:</u> Examples of lung diseases are pulmonary arterial hypertension and interstitial pneumonitis.</p> <p>c) Patient is at risk of bleeding; OR</p> <p><u>Note:</u> An example of a patient with an increased risk of bleeding is a patient with thrombocytopenia or with medication use that inhibits platelet function or anticoagulants.</p> <p>d) Patient has a prolonged QT interval or is at risk of developing QT interval prolongation; OR</p> <p>e) Patient has tried at least two other tyrosine kinase inhibitors for CML; OR</p> <p><u>Note:</u> Examples of tyrosine kinase inhibitors include: dasatinib, nilotinib, Iclusig and Scemblix.</p> <p>f) Patient has a resistance mutation in which dasatinib should not be used; OR</p> <p>ii. Patient is currently receiving therapy with Bosulif.</p> <p>B. For <b>Acute Lymphoblastic Leukemia</b>, ONE of the following:</p> <p>i. Trial of, contraindication, or significant intolerance to ONE of the following: imatinib or dasatinib ; OR</p> <p><u>Note:</u> Prior use of Gleevec, Imkeldi, Phyrago, or Sprycel also counts.</p> <p>ii. Patient is currently receiving therapy with Bosulif; OR</p> <p>iii. Patient meets ONE of the following:</p> <ol style="list-style-type: none"> <li>1. Patient has a history of serious, chronic lung disease or has had or is at risk of pleural effusion; OR</li> <li><u>Note:</u> Examples of lung diseases include pulmonary arterial hypertension and interstitial pneumonitis.</li> <li>2. Patient is at risk of bleeding; OR</li> <li><u>Note:</u> An example of a patient with an increased risk of bleeding is a patient with thrombocytopenia or with medication use that inhibits platelet function or anticoagulants.</li> <li>3. Patient has a prolonged QT interval or is at risk of developing QT interval prolongation; OR</li> </ol> <p>iv. Patient has a mutation in which imatinib or dasatinib should not be used,</p> <p>C. <b>All Other Conditions.</b> Approve Bosulif if the patient meets the Oncology Medications criteria above the table</p>
<p><b>Braftovi®</b> (encorafenib)</p>	<p><b><u>Employer Plans and Individual and Family Plan</u></b></p>

	<p><b>Braftovi is considered medically necessary when BOTH of the following are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. <b>ONE</b> of the following: <ol style="list-style-type: none"> <li>A. For <b>Melanoma, Unresectable or Metastatic, Treatment of BRAF V600 Mutation-Positive Disease</b>, documentation of <b>ONE</b> of the following: <ol style="list-style-type: none"> <li>i. Trial of, contraindication, significant intolerance, or other exceptional clinical circumstance to <b>ONE</b> of the following: <ol style="list-style-type: none"> <li>1. <b>Tafinlar</b> [may require prior authorization]</li> <li>2. <b>Zelboraf</b> [may require prior authorization]</li> </ol> </li> <li>ii. Patient is currently receiving Braftovi</li> </ol> </li> <li>B. <b>All Other Conditions.</b> Approve Braftovi if the patient meets the Oncology Medications criteria above the table</li> </ol> </li> </ol>
<p><b>Cyclophosphamide tablets</b></p> <p><i>This applies to oncology and non-oncology uses of cyclophosphamide.</i></p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Cyclophosphamide tablets is considered medically necessary when BOTH of the following are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Documented trial of, contraindication, or intolerance to cyclophosphamide capsules</li> </ol>
<p><b>Danziten</b> (nilotinib tartrate tablets)</p>	<p><b><u>Employer Plans</u></b></p> <p><b>Danziten is considered medically necessary when BOTH of the following are met (A and B):</b></p> <ol style="list-style-type: none"> <li>A. When the Oncology Medications criteria above the table are met</li> <li>B. <b>ONE</b> of the following (a or b): <ol style="list-style-type: none"> <li>a. For <b>Chronic Myeloid Leukemia (CML)</b>, <b>ONE</b> the following: <ol style="list-style-type: none"> <li>A. Patient meets <b>ONE</b> of the following: <ol style="list-style-type: none"> <li>1. Trial of, contraindication, or significant intolerance to <b>ONE</b> of the following: imatinib or dasatinib <i>Note:</i> Prior use of brand Gleevec, Imkeldi, Phyrago, or Sprycel, also counts.</li> <li>2. Patient has tried AND cannot take generic nilotinib hcl capsules due to due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the non-preferred and preferred product which, per the prescriber, would result in a significant allergy or serious adverse reaction; OR</li> </ol> </li> <li>B. Patient is currently receiving therapy with Danziten</li> <li>C. Patient has a resistance mutation in which imatinib and dasatinib, should not be used</li> </ol> </li> <li>b. <b>All Other Conditions.</b> Approve Danziten if the patient meets the Oncology Medications criteria above the table</li> </ol> </li> </ol> <p><b><u>Individual and Family Plans</u></b></p>

	<p><b>Danziten is considered medically necessary when BOTH of the following are met (A and B):</b></p> <ul style="list-style-type: none"> <li>A. When the Oncology Medications criteria above the table are met</li> <li>B. <b>ONE</b> of the following (a or b): <ul style="list-style-type: none"> <li>a. For <b>Chronic Myeloid Leukemia (CML)</b>, <b>ONE</b> the following: <ul style="list-style-type: none"> <li>A. Trial of, contraindication, or significant intolerance to <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>1. dasatinib</li> <li>2. imatinib</li> </ul> <p><u>Note:</u> Prior use of brand Gleevec, Imkeldi, Phyrago, or Sprycel also counts</p> </li> <li>B. Patient is currently receiving therapy with Danziten</li> <li>C. Patient meets <b>BOTH</b> of the following: <ul style="list-style-type: none"> <li>1. Patient meets <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>a. Patient has intermediate- to high-risk chronic phase CML</li> <li>b. Patient has accelerated phase CML or blast phase CML</li> </ul> </li> <li>2. Patient meets <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>a. Patient has a history of serious, chronic lung disease or has had or is at risk of pleural effusion; OR</li> <li><u>Note:</u> Examples of lung disease are pulmonary arterial hypertension and interstitial pneumonitis.</li> <li>b. Patient is at risk of bleeding; OR</li> <li><u>Note:</u> An example of a patient with an increased risk of bleeding is a patient with thrombocytopenia or with medication use that inhibits platelet function or anticoagulants.</li> </ul> </li> </ul> </li> <li>D. Patient has a resistance mutation in which imatinib and dasatinib should not be used</li> </ul> </li> <li>b. <b>All Other Conditions.</b> Approve Danziten if the patient meets the Oncology Medications criteria above the table</li> </ul> </li> </ul>
<p><b>Docivyx</b> (docetaxel)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Docivyx is considered medically necessary when BOTH of the following are met:</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>A. Patient has tried generic docetaxel</li> <li>B. Patients with hypersensitivities to polysorbate 80</li> </ul> </li> </ul>
<p><b>Ensacove</b> (ensartinib hydrochloride)</p>	<p><b><u>Individual and Family Plans</u></b></p> <p><b>Ensacove is considered medically necessary when BOTH of the following are met:</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>A. For <b>Non-Small Cell Lung Cancer – anaplastic lymphoma kinase (ALK)-positive</b>, <b>ONE</b> of the following:</li> </ul> </li> </ul>

	<p><b>1.</b> Patient has tried or is unable to take Alecensa  <u>Note:</u> A trial of Lorbrena would also meet an approval.</p> <p><b>2.</b> Patient has already been started on therapy with Ensacove.</p> <p><b>B. All Other Conditions.</b> Approve Ensacove if the patient meets the Oncology Medications criteria above the table</p>
<p><b>Fruzaqla™</b>  (fruquintinib)</p>	<p><b><u>Individual and Family Plans:</u></b></p> <p><b>Fruzaqla (fruquintinib) is considered medically necessary when BOTH of the following are met (1 and 2):</b></p> <p>1. When the Oncology Medications criteria above the table are met</p> <p>2. <b>ONE</b> of the following:</p> <p>A. For <b><u>Appendiceal, Colon or Rectal Cancer in an individual 18 years of age or older, ONE</u></b> of the following:</p> <p>i. Trial of, contraindication, or intolerance to <b>Lonsurf (trifluridine-tipiracil) tablets</b> [may require prior authorization]</p> <p>ii. According to the prescriber, the patient has or is at risk of myelosuppression</p> <p>iii. Patient has already been started on therapy with Fruzaqla</p> <p><b>B. All Other Conditions.</b> Approve Fruzaqla if the patient meets the Oncology Medications criteria above the table</p>
<p><b>Gleevec®</b>  (imatinib)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Gleevec is considered medically necessary when BOTH of the following are met:</b></p> <p>1. When the Oncology Medications criteria above the table are met</p> <p>2. Patient meets <b>BOTH</b> of the following (A <u>and</u> B):</p> <p>A) Patient has tried generic imatinib tablets ; AND</p> <p>B) Patient cannot take generic imatinib tablets due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the brand and bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction.</p>
<p><b>Gleostine</b>  (lomustine capsules)</p>	<p><b><u>Employer Plans</u></b></p> <p><b>Gleostine is considered medically necessary when BOTH of the following are met:</b></p> <p>1. When the Oncology Medications criteria above the table are met</p> <p>2. The patient has tried the bioequivalent generic product AND cannot take due to a formulation difference in the inactive ingredient(s) [for example, difference in dyes, fillers, preservatives] between the brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction.</p>
<p><b>Herceptin®</b>  (trastuzumab)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Herceptin (trastuzumab) is considered medically necessary when BOTH of the following are met:</b></p> <p>1. When the Oncology Medications criteria above the table are met</p>

	<p>2. Documentation of <b>ONE</b> of the following:</p> <p>A. Trial of AND cannot continue to use the alternative(s) due to a formulation difference in the inactive ingredient(s) which, according to the prescriber, would result in a significant allergy or serious adverse reaction to <b>ALL</b> of the following:</p> <ul style="list-style-type: none"> <li>i. <b>Kanjinti (trastuzumab-anns)</b> [may require prior authorization]</li> <li>ii. <b>Ogivri (trastuzumab-dkst)</b> [may require prior authorization]</li> <li>iii. <b>Trazimera (trastuzumab-qyyp)</b> [may require prior authorization]</li> </ul>
<p><b>Herceptin Hylecta™</b> (trastuzumab and hyaluronidase-oysk)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Herceptin Hylecta (trastuzumab and hyaluronidase-oysk) is considered medically necessary when BOTH of the following are met:</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Documentation of <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>A. Patient has trial of, contraindication, or intolerance to <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>i. <b>Kanjinti (trastuzumab-anns)</b> [may require prior authorization]</li> <li>ii. <b>Ogivri (trastuzumab-dkst)</b> [may require prior authorization]</li> <li>iii. <b>Trazimera (trastuzumab-qyyp)</b> [may require prior authorization]</li> </ul> </li> <li>B. Patient is unable to obtain or maintain intravenous access</li> <li>C. Currently receiving Herceptin Hylecta</li> </ul> </li> </ul>
<p><b>Hercessi</b> (trastuzumab-strf)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Hercessi (trastuzumab-strf) is considered medically necessary when BOTH of the following are met (1 and 2):</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Documentation of <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>A. Trial of AND cannot continue to use the alternative(s) due to a formulation difference in the inactive ingredient(s) which, according to the prescriber, would result in a significant allergy or serious adverse reaction to <b>ALL</b> of the following: <ul style="list-style-type: none"> <li>i. <b>Kanjinti (trastuzumab-anns)</b> [may require prior authorization]</li> <li>ii. <b>Ogivri (trastuzumab-dkst)</b> [may require prior authorization]</li> <li>iii. <b>Trazimera (trastuzumab-qyyp)</b> [may require prior authorization]</li> </ul> </li> </ul> </li> </ul>
<p><b>Herzuma®</b> (trastuzumab-pkrb)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Herzuma (trastuzumab-pkrb) is considered medically necessary when BOTH of the following are met:</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Documentation of <b>ONE</b> of the following:</li> </ul>

	<p>A. Trial of AND cannot continue to use the alternative(s) due to a formulation difference in the inactive ingredient(s) which, according to the prescriber, would result in a significant allergy or serious adverse reaction to <b>ALL</b> of the following:</p> <ul style="list-style-type: none"> <li>i. <b>Kanjinti (trastuzumab-anns)</b> [may require prior authorization]</li> <li>ii. <b>Ogivri (trastuzumab-dkst)</b> [may require prior authorization]</li> <li>iii. <b>Trazimera (trastuzumab-qyyp)</b> [may require prior authorization]</li> </ul>
<p><b>Ibrance®</b> (palbociclib)</p>	<p><b><u>Employer Plans</u></b></p> <p><b>Ibrance (palbociclib) is considered medically necessary when BOTH of the following are met:</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>A. <b>For Breast Cancer</b>, documentation of <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>i. Trial of, contraindication, or intolerance to <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>a. <b>Kisqali (ribociclib)</b> [may require prior authorization]</li> <li>b. <b>Verzenio (abemaciclib)</b> [may require prior authorization]</li> </ul> </li> <li>ii. For premenopausal patients using in combination with fulvestrant as subsequent therapy (not initial therapy), approve if the patient has tried Verzenio</li> <li>iii. Patient will be using Ibrance in combination with Itovebi</li> <li>iv. Currently receiving Ibrance</li> </ul> </li> <li>B. <b>All Other Conditions.</b> Approve Ibrance if the patient meets the Oncology Medications criteria above the table</li> </ul> </li> </ul>
<p><b>Iclusig</b> (ponatinib tablets)</p>	<p><b><u>Employer Plans</u></b></p> <p><b>Iclusig is considered medically necessary when BOTH of the following are met (1 and 2):</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Meets <b>ONE</b> of the following (A, B, C, or D): <ul style="list-style-type: none"> <li>A. <b>For Chronic Myeloid Leukemia (CML), ONE</b> of the following: <ul style="list-style-type: none"> <li>i. Trial of, contraindication, significant intolerance to <b>TWO</b> of the following: <ul style="list-style-type: none"> <li>1) <b>dasatinib</b></li> <li>2) <b>imatinib</b></li> <li>3) <b>nilotinib hcl</b></li> <li>4) <b>Danziten</b> [may require prior authorization]</li> <li>5) <b>Imkeldi</b> [may require prior authorization]</li> <li>6) <b>Scemblix</b> [may require prior authorization]</li> </ul> <p><u>Note:</u> Prior use of brand Gleevec, Phyrago, Sprycel, or Tassigna also counts.</p> </li> <li>ii. Patient meets <b>BOTH</b> of the following: <ul style="list-style-type: none"> <li>1) Patient has intermediate- to high-risk chronic phase CML, accelerated phase CML, or blast phase CML</li> <li>2) Patient has tried at least two other tyrosine kinase inhibitors for CML</li> </ul> </li> </ul> </li> </ul> </li> </ul>

Note: Examples of tyrosine kinase inhibitors include: dasatinib, Bosulif, nilotinib, and Scemblix.

- iii. Patient has a resistance mutation in which imatinib, dasatinib, nilotinib hcl, Danziten, Imkeldi, and Scemblix should not be used
- iv. Patient has the *T315I* mutation and is in accelerated or blast phase
- v. Patient is currently receiving therapy Iclusig

B. For **Acute Lymphoblastic Leukemia (ALL)**, **ONE** of the following:

- i. Trial of, contraindication, significant intolerance to **ONE** of the following:
  - 1) **dasatinib**
  - 2) **imatinib**

Note: Prior use of Gleevec , Imkeldi, Phyrago, or Sprycel also counts.

- ii. Patient is  $\geq 18$  years of age, has newly diagnosed disease, and is taking the requested medication with chemotherapy; OR
- iii. Patient has T315I-positive mutation; OR
- iv. Patient has a history of serious, chronic lung disease or has had or is a risk of pleural effusion

Note: Examples of lung diseases include pulmonary arterial hypertension and interstitial pneumonitis.

- v. Patient has a resistance mutation in which imatinib or dasatinib should not be used
- vi. Patient is currently receiving therapy with Iclusig

**C. All Other Conditions.** Approve Iclusig if the patient meets the Oncology Medications criteria above the table

### **Individual and Family Plans**

**Iclusig is considered medically necessary when BOTH of the following are met (1 and 2):**

- 1. When the Oncology Medications criteria above the table are met
- 2. Meets **ONE** of the following (A, B, or C):

A. For **Chronic Myeloid Leukemia (CML)**, **ONE** of the following:

- i. Patient meets **BOTH** of the following:

1) Patient meets **ONE** of the following:

- a. Trial of, contraindication, significant intolerance to **imatinib**
- b. Patient has intermediate- to high-risk chronic phase CML, accelerated phase CML, or blast phase CML
- c. Patient has tried at least one other tyrosine kinase inhibitor for CML

Note: Examples of tyrosine kinase inhibitors include: dasatinib, Bosulif, nilotinib, and Scemblix.

	<ul style="list-style-type: none"> <li>d. Patient has a resistance mutation in which imatinib not be used</li> <li>2) Patient meets <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>a. Trial of, contraindication, significant intolerance to <b>dasatinib</b> <u>Note:</u> Prior use of Phyrago or Sprycel also counts.</li> <li>b. Patient has tried at least two other tyrosine kinase inhibitors for CML <u>Note:</u> Examples of tyrosine kinase inhibitors include: dasatinib, Bosulif, nilotinib, and Scemblix.</li> <li>c. Patient has a history of serious, chronic lung disease or has had or is a risk of pleural effusion <u>Note:</u> Examples of lung diseases include pulmonary arterial hypertension and interstitial pneumonitis.</li> <li>d. Patient has a resistance mutation in which dasatinib should not be used</li> </ul> </li> <li>ii. Patient has the <i>T315I</i> mutation</li> <li>iii. Patient is currently receiving therapy with Iclusig</li> </ul> <p>B. For <b>Acute Lymphoblastic Leukemia (ALL)</b>, <b>ONE</b> of the following:</p> <ul style="list-style-type: none"> <li>i. Trial of, contraindication, significant intolerance to <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>1) <b>imatinib</b></li> <li>2) <b>dasatinib</b></li> </ul> <u>Note:</u> Prior use of Gleevec, Imkeldo, Phyrago, or Sprycel also counts. </li> <li>ii. Patient is <math>\geq 18</math> yrs of age, has newly diagnosed disease, and is taking the requested medication with chemotherapy; OR</li> <li>iii. Patient has T315I mutation; OR</li> <li>iv. Patient has a history of serious, chronic lung disease or has had or is a risk of pleural effusion <u>Note:</u> Examples of lung diseases include pulmonary arterial hypertension and interstitial pneumonitis.</li> <li>v. Patient has a resistance mutation in which imatinib or dasatinib should not be used</li> <li>vi. Patient is currently receiving therapy with Iclusig</li> </ul> <p>C. <b>All Other Conditions.</b> Approve Iclusig if the patient meets the Oncology Medications criteria above the table</p>
<p><b>Imkeldi</b> (imatinib oral solution)</p>	<p><b><u>Employer Plans</u></b></p> <p><b>Imkeldi is considered medically necessary when BOTH of the following is met (1 and 2):</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Meets <b>ONE</b> of the following (A or B): <ul style="list-style-type: none"> <li>A. For <b>Chronic Myeloid Leukemia (CML)</b>, patient meets <b>BOTH</b> of the following:</li> </ul> </li> </ul>

	<ul style="list-style-type: none"> <li>i. Patient is <math>\geq</math> 18 years old; AND</li> <li>ii. Patient meets ONE of the following: <ul style="list-style-type: none"> <li>1. Patient meets BOTH of the following: <ul style="list-style-type: none"> <li>a. Patient has tried imatinib tablets; AND <u>Note:</u> Prior use of Gleevec also counts.</li> <li>b. Patient cannot take generic imatinib due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the brand and bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction; OR</li> </ul> </li> <li>2. Patient is unable to swallow or has difficulty swallow tablets.</li> </ul> </li> </ul> <p><b>B. All Other Conditions.</b> Approve Imkeldi if the patient meets the Oncology Medications criteria above the table</p> <p><b><u>Individual and Family Plans</u></b></p> <p><b>Imkeldi is considered medically necessary when BOTH of the following is met (1 and 2):</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Patients meets ONE of the following (A or B): <ul style="list-style-type: none"> <li>A. Patient meets BOTH of the following: <ul style="list-style-type: none"> <li>i. Patient has tried imatinib tablets ; AND <u>Note:</u> Prior use of Gleevec also counts.</li> <li>ii. Patient cannot take generic imatinib due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the brand and bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction; OR</li> </ul> </li> <li>B. Patient is unable to swallow or has difficulty swallowing tablets.</li> </ul> </li> </ul>
<p><b>Infugem™</b> (gemcitabine)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Infugem (gemcitabine) is considered medically necessary when BOTH of the following is met:</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Documented trial of, contraindication, or intolerance to generic gemcitabine</li> </ul>
<p><b>Ivra</b> (melphalan)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Ivra (melphalan) is considered medically necessary when BOTH of the following are met:</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Patient has tried and cannot use melphalan injection (Alkeran, generics).</li> </ul>
<p><b>Jemperli™</b> (dostarlimab)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p>

	<p><b>Jemperli is considered medically necessary when BOTH of the following are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. <b>ONE of the following:</b> <ol style="list-style-type: none"> <li>A. For <b><u>Mismatch Repair Deficient (dMMR) or Microsatellite Instability-High (MSI-H) Endometrial Cancer - Monotherapy, ONE</u></b> of the following:           <ol style="list-style-type: none"> <li>i. Patient has tried Keytruda [may require prior authorization]</li> <li>ii. Patient has already been started on therapy with Jemperli</li> </ol> </li> <li>B. For <b><u>Mismatch Repair Deficient (dMMR) or Microsatellite Instability-High (MSI-H) Solid Tumors - Monotherapy, ONE</u></b> of the following:  <u>Note:</u> Examples of solid tumors include ampullary adenocarcinoma, biliary tract cancer, breast cancer, esophageal and esophagogastric junction cancer, gastric cancer, hepatocellular cancer, and ovarian cancer.           <ol style="list-style-type: none"> <li>i. Patient has tried Keytruda [may require prior authorization]</li> <li>ii. Patient has already been started on therapy with Jemperli</li> </ol> </li> <li>C. <b>All Other Conditions</b> (e.g., rectal cancer). Approve Jemperli if the patient meets the Oncology Medications criteria above the table</li> </ol> </li> </ol>
<p><b>Jobevne</b> (bevacizumab-nwgd)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Jobevne is considered medically necessary when BOTH of the following are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Patient meets <b>BOTH</b> of the following (A and B):       <ol style="list-style-type: none"> <li>A. Patient has tried <b>BOTH</b> of the following <b>[documentation required]:</b> <ol style="list-style-type: none"> <li>i. <b>Mvasi (bevacizumab-awwb)</b> [may require prior authorization]</li> <li>ii. <b>Zirabev (bevacizumab-bvzr)</b> [may require prior authorization]</li> </ol> </li> <li>B. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.</li> </ol> </li> </ol>
<p><b>Keytruda IV</b> (pembrolizumab intravenous infusion)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Keytruda IV is considered medically necessary when BOTH of the following are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. <b>ONE</b> of the following:       <ol style="list-style-type: none"> <li>A. For, <b><u>Nasopharyngeal Carcinoma, ONE</u></b> of the following:           <ol style="list-style-type: none"> <li>i. According to the prescriber, the patient has tried, and has had inadequate efficacy or significant intolerance or patient has a contraindication to Loqtorzi; OR</li> </ol> </li> </ol> </li> </ol>

	<ul style="list-style-type: none"> <li>ii. Patient has been started on Keytruda IV or Keytruda Qlex; OR</li> <li>iii. Patient has a diagnosis of head and neck squamous cell carcinoma other than nasopharyngeal carcinoma.</li> </ul> <p>B. <b>All Other Conditions.</b> Approve Keytruda if the patient meets the Oncology Medications criteria above the table</p>
<p><b>Keytruda Qlex</b> (pembrolizumab and berahyaluronidase alpha-pmpH subcutaneous injection)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Keytruda Qlex is considered medically necessary when BOTH of the following are met:</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>A. Patient has tried and cannot take Keytruda intravenous (IV) [may require prior authorization]</li> <li>B. Patient is unable to obtain IV access.</li> </ul> </li> </ul>
<p><b>Khazory™</b> (levoleucovorin)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Khazory (levoleucovorin) is considered medically necessary when BOTH of the following are met:</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Meets <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>A. Patient has tried one generic levoleucovorin calcium injection or generic leucovorin injection</li> <li>B. If the patient has already been started on therapy with Khazory, patient has tried generic levoleucovorin calcium injection</li> </ul> </li> </ul>
<p><b>Krazati</b> (adagrasib)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Krazati (adagrasib) is considered medically necessary when BOTH of the following are met:</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>A. For <b><u>KRAS G12C-mutated Non-Small Cell Lung Cancer</u></b>, documentation of <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>i. Trial of, contraindication, or intolerance to <b>sotorasib (Lumakras)</b> [may require prior authorization]</li> <li>ii. Patient has brain metastases</li> <li>iii. Patient has already been started on therapy with Krazati</li> </ul> </li> <li>B. <b>All Other Conditions.</b> Approve Krazati if the patient meets the Oncology Medications criteria above the table</li> </ul> </li> </ul>
<p><b>Kyxata</b> (carboplatin)</p>	<p><b><u>Employer Plans and Individual and Family Plan</u></b></p> <p><b>Kyxata is considered medically necessary when BOTH of the following are met:</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Generic Carboplatin is not available on the market per a verifiable source (for example, FDA shortage database or ASHP Drug Shortage list)</li> </ul>

<p><b>Mektovi®</b> (binimetinib)</p>	<p><b><u>Employer Plans and Individual and Family Plan</u></b></p> <p><b>Mektovi is considered medically necessary when BOTH of the following are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. <b>ONE</b> of the following: <ol style="list-style-type: none"> <li>A. For <b>Melanoma, Unresectable or Metastatic, Treatment of BRAF V600 Mutation-Positive Disease</b>, documentation of <b>ONE</b> of the following: <ol style="list-style-type: none"> <li>i. Trial of, contraindication, significant intolerance, or other exceptional clinical circumstance to <b>ONE</b> of the following: <ol style="list-style-type: none"> <li>1. <b>Cotellic</b> [may require prior authorization]</li> <li>2. <b>Mekinist</b> [may require prior authorization]</li> </ol> </li> <li>ii. Patient is currently receiving Mektovi</li> </ol> </li> <li>B. <b>All Other Conditions.</b> Approve Mektovi if the patient meets the Oncology Medications criteria above the table</li> </ol> </li> </ol>
<p><b>Nexavar</b> (sorafenib)</p>	<p><b><u>Employer Plans</u></b></p> <p><b>Nexavar (sorafenib) is considered medically necessary when BOTH of the following are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Documented trial of <b>sorafenib</b> (the bioequivalent generic product) AND cannot take due to a formulation difference in the inactive ingredient(s) which would result in a significant allergy or serious adverse reaction</li> </ol> <p><b><u>Individual and Family Plans</u></b></p> <p><b>Nexavar (sorafenib) is considered medically necessary when BOTH of the following are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Documented trial of <b>sorafenib</b> (the bioequivalent generic product) AND cannot take due to a formulation difference in the inactive ingredient(s) which would result in a significant allergy or serious adverse reaction</li> </ol>
<p><b>Nilandron®</b> (nilutamide)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Nilandron (nilutamide) is considered medically necessary when BOTH of the following are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Documented trial of <b>nilutamide</b> (the bioequivalent generic product) AND cannot take due to formulation difference in the inactive ingredient(s) which would result in a significant allergy or serious adverse reaction</li> </ol>
<p><b>Nilotinib hcl</b></p>	<p><b><u>Individual and Family Plans</u></b></p> <p><b>Nilotinib hcl is considered medically necessary when BOTH of the following are met (1 and 2):</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. <b>ONE</b> of the following (A or B): <ol style="list-style-type: none"> <li>A. For <b>Chronic Myeloid Leukemia (CML)</b>, <b>ONE</b> of the following:</li> </ol> </li> </ol>

	<ul style="list-style-type: none"> <li>i. Trial of, contraindication, significant intolerance to <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>a. <b>imatinib</b></li> <li>b. <b>dasatinib</b></li> </ul> <p><u>Note:</u> Prior use of brand Gleevec, Imkeldi, Phyrago or Sprycel counts.</p> </li> <li>ii. Patient is currently receiving nilotinib hcl</li> <li>iii. Patient is less than 18 years of age with accelerated phase CML</li> <li>iv. Patient meets <b>BOTH</b> of the following: <ul style="list-style-type: none"> <li>a. Patient meets <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>i. Patient has intermediate- to high-risk disease chronic phase CML</li> <li>ii. Patient has accelerated phase CML or blast phase CML</li> </ul> </li> <li>b. Patient meets <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>i. Patient has a history of serious, chronic lung disease or has had or is at risk of pleural effusion <u>Note:</u> Examples of lung disease, pulmonary arterial hypertension, and interstitial pneumonitis.</li> <li>ii. Patient is at risk of bleeding <u>Note:</u> An example of a patient with an increased risk of bleeding is a patient with thrombocytopenia or with a medication that inhibits platelet function or anticoagulants.</li> </ul> </li> </ul> </li> <li>v. Patient has a resistance mutation in which imatinib and dasatinib should not be used</li> </ul> <p>B. <b>All Other Conditions.</b> Approve nilotinib if the patient meets the Oncology Medications criteria above the table</p>
<p><b>Nilotinib d-tartrate</b></p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Nilotinib d-tartrate is considered medically necessary when BOTH of the following are met (1 and 2):</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Patient has tried and cannot take generic nilotinib hcl capsules [may require prior authorization] due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the non-preferred and preferred product which, per the prescriber, would result in a significant allergy or serious adverse reaction.</li> </ul>
<p><b>Onivyde</b> (irinotecan liposomal intravenous infusion)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Onivyde (irinotecan liposomal intravenous infusion) is considered medically necessary when BOTH of the following are met:</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. <b>ONE</b> of the following (A, B, or C): <ul style="list-style-type: none"> <li>A. For <b>Pancreatic Adenocarcinoma</b>, documentation of <b>ONE</b> of the following:</li> </ul> </li> </ul>

	<ul style="list-style-type: none"> <li>i. According to the prescriber, the patient has experienced an inadequate response or significant intolerance, has a contraindication for irinotecan intravenous infusion; OR</li> <li>ii. According to the prescriber, the patient has baseline neuropathy; OR</li> <li>iii. Patient has been started on Onivyde; OR</li> <li>iv. Patient meets <b>BOTH</b> of the following: <ul style="list-style-type: none"> <li>a. Medication will be used for subsequent therapy</li> <li>b. Patient meets <b>ONE</b> of the following (1, 2, or 3): <ul style="list-style-type: none"> <li>1. According to the prescriber, patient is Eastern Cooperative Oncology Group performance status of 2; OR</li> <li>2. According to prescriber, patient is Eastern Cooperative Oncology Group Performance Status 0 or 1 and has tried a gemcitabine-based regimen; OR</li> <li>3. According to prescriber, patient is Eastern Cooperative Oncology Group Performance Status 0 or 1 and has tried a fluoropyrimidine-based regimen but without prior irinotecan.</li> </ul> </li> </ul> </li> </ul> <p>B. For <b>Ampullary Adenocarcinoma</b>, documentation of <b>ONE</b> of the following:</p> <ul style="list-style-type: none"> <li>i. According to the prescriber, the patient has experienced an inadequate response, significant intolerance, or has a contraindication for irinotecan intravenous infusion; OR</li> <li>ii. According to the prescriber, the patient has baseline neuropathy; OR</li> <li>iii. Patient has been started on Onivyde.</li> </ul> <p>C. <b>All Other Conditions.</b> Approve Onivyde if the patient meets the Oncology Medications criteria above the table</p>
<p><b>Ontruzant®</b> (trastuzumab-dttb)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Ontruzant (trastuzumab-dttb) is considered medically necessary when BOTH of the following are met:</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Documentation of <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>A. Trial of AND cannot continue to use the alternative(s) due to a formulation difference in the inactive ingredient(s) which, according to the prescriber, would result in a significant allergy or serious adverse reaction to <b>ALL</b> of the following: <ul style="list-style-type: none"> <li>i. <b>Kanjinti (trastuzumab-anns)</b> [may require prior authorization]</li> <li>ii. <b>Ogivri (trastuzumab-dkst)</b> [may require prior authorization]</li> <li>iii. <b>Trazimera (trastuzumab-qyyp)</b> [may require prior authorization]</li> </ul> </li> </ul> </li> </ul>
<p><b>Opdivo Intravenous</b> (nivolumab)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Opdivo (nivolumab) is considered medically necessary when BOTH of the following are met:</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> </ul>

	<p>2. <b>ONE</b> of the following:</p> <p>A. For, <b><u>Nasopharyngeal Carcinoma</u></b>, documentation of <b>ONE</b> of the following:</p> <ul style="list-style-type: none"> <li>i. Patient has been started on Opdivo</li> <li>ii. Patient meets <b>ALL</b> of the following: <ul style="list-style-type: none"> <li>a. Patient has recurrent, unresectable, oligometastatic, or metastatic disease</li> <li>b. The medication is used in combination with cisplatin and gemcitabine</li> <li>c. According to the prescriber, the patient has inadequate efficacy, contraindication, or significant intolerance to <b>Loqtorzi (toripalimab intravenous infusion)</b> [may require prior authorization]</li> </ul> </li> <li>iii. Patient meets <b>BOTH</b> of the following: <ul style="list-style-type: none"> <li>a. Patient has recurrent or metastatic non-keratinizing disease; AND</li> <li>b. Medication is used for subsequent therapy.</li> </ul> </li> </ul> <p>B. <b>All Other Conditions.</b> Approve Opdivo if the patient meets the Oncology Medications criteria above the table</p>
<p><b>Opdivo Qvantig</b> (nivolumab and hyaluronidase-nvhy)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Opdivo Qvantig (nivolumab and hyaluronidase-nvhy) is considered medically necessary when BOTH of the following are met:</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Documentation provided that the patient has <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>A. Patient has tried and cannot take Opdivo intravenous (IV) [may require prior authorization]</li> <li>B. Patient is unable to obtain IV access.</li> </ul> </li> </ul>
<p><b>Orgovyx®</b> (relugolix)</p>	<p><b><u>Individual and Family Plans</u></b></p> <p><b>Orgovyx (relugolix) is considered medically necessary when BOTH of the following are met:</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>A. For, <b><u>Prostate Cancer</u></b>, documentation of <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>i. Patient has tried <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>a. <b>Eligard</b> [may require prior authorization]</li> <li>b. <b>Firmagon</b> [may require prior authorization]</li> <li>c. <b>Trelstar</b> [may require prior authorization]</li> </ul> </li> <li>ii. According to the prescriber, is at risk of cardiovascular disease</li> <li>iii. Using for intermittent androgen deprivation therapy</li> <li>iv. Currently receiving Orgovyx</li> </ul> </li> <li>B. <b>All Other Conditions.</b> Approve Orgovyx if the patient meets the Oncology Medications criteria above the table</li> </ul> </li> </ul>
<p><b>Paclitaxel albumin-bound intravenous infusion</b></p>	<p><b><u>Cigna Pathwell Specialty Drug List Plans</u></b></p> <p><b>Paclitaxel albumin-bound intravenous infusion is considered medically necessary when BOTH of the following are met:</b></p>

1. When the Oncology Medications criteria above the table are met
2. **ONE** of the following:
  - A. For **Breast Cancer**, **ONE** of the following:
    - i. Patient is currently receiving Abraxane or paclitaxel albumin-bound intravenous infusion
    - ii. Patient has tried paclitaxel intravenous infusion
    - iii. Patient had a hypersensitivity to paclitaxel intravenous infusion or docetaxel intravenous infusion
    - iv. Patient has a contraindication to the standard pre-medications (for example, dexamethasone, ranitidine, famotidine, diphenhydramine)
  - B. For **Cervical Cancer**, **ONE** of the following:
    - i. Patient is currently receiving Abraxane or paclitaxel albumin-bound intravenous infusion
    - ii. Patient has tried paclitaxel intravenous infusion
    - iii. Patient had a hypersensitivity to paclitaxel intravenous infusion or docetaxel intravenous infusion
    - iv. Patient has a contraindication to the standard pre-medications (for example, dexamethasone, ranitidine, famotidine, diphenhydramine)
  - C. For **Endometrial Cancer**, **ONE** of the following:
    - i. Patient is currently receiving Abraxane or paclitaxel albumin-bound intravenous infusion
    - ii. Patient has tried paclitaxel intravenous infusion
    - iii. Patient had a hypersensitivity to paclitaxel intravenous infusion or docetaxel intravenous infusion
    - iv. Patient has a contraindication to the standard pre-medications (for example, dexamethasone, ranitidine, famotidine, diphenhydramine)
  - D. For **Melanoma**, **ONE** of the following:
    - i. Patient is currently receiving Abraxane or paclitaxel albumin-bound intravenous infusion
    - ii. Patient has tried paclitaxel intravenous infusion
    - iii. Patient had a hypersensitivity to paclitaxel intravenous infusion or docetaxel intravenous infusion
    - iv. Patient has a contraindication to the standard pre-medications (for example, dexamethasone, ranitidine, famotidine, diphenhydramine)
  - E. For **Non-Small Cell Lung Cancer**, **ONE** of the following:
    - i. Patient is currently receiving Abraxane or paclitaxel albumin-bound intravenous infusion
    - ii. Patient has tried paclitaxel intravenous infusion
    - iii. Patient had a hypersensitivity to paclitaxel intravenous infusion or docetaxel intravenous infusion
    - iv. Patient has a contraindication to the standard pre-medications (for example, dexamethasone, ranitidine, famotidine, diphenhydramine)
    - v. Abraxane or paclitaxel albumin-bound intravenous infusion is being used as subsequent therapy in patients with advanced or metastatic disease
  - F. For **Ovarian Cancer**, **ONE** of the following:
    - i. Patient is currently receiving Abraxane or paclitaxel albumin-bound intravenous infusion

	<ul style="list-style-type: none"> <li>ii. Patient has tried paclitaxel intravenous infusion</li> <li>iii. Patient had a hypersensitivity to paclitaxel intravenous infusion or docetaxel intravenous infusion</li> <li>iv. Patient has a contraindication to the standard pre-medications (for example, dexamethasone, ranitidine, famotidine, diphenhydramine)</li> </ul> <p>G. <b>All Other Conditions.</b> Approve paclitaxel albumin-bound intravenous infusion if the patient meets the Oncology Medications criteria above the table</p>
<p><b>Pazopanib 400 mg</b></p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Pazopanib 400 mg is considered medically necessary when BOTH of the following are met (1 and 2):</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Patient has tried and experienced inadequate efficacy or significant intolerance with pazopanib 200 mg tablets.</li> </ul>
<p><b>Phyrago</b> (dasatinib tablets)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Phyrago is considered medically necessary when BOTH of the following are met (1 and 2):</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. <b>ONE of the following:</b> <ul style="list-style-type: none"> <li>A. Patient has tried AND cannot take generic dasatinib tablets due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the preferred and non-preferred product which, per the prescriber, would result in a significant allergy or serious adverse reaction; OR</li> <li>B. According to the prescriber, the patient requires co-administration with a histamine-2 (H2) antagonist or proton pump inhibitor (PPI). <b>Note:</b> Examples of H2 antagonists include famotidine, cimetidine, nizatidine. Examples of PPIs include omeprazole, esomeprazole, pantoprazole, rabeprazole.</li> </ul> </li> </ul>
<p><b>Revlimid</b> (lenalidomide)</p>	<p><b><u>Employer Plans</u></b></p> <p><b>Revlimid is considered medically necessary when BOTH of the following are met (1 and 2):</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. <b>BOTH</b> of the following (A <u>and</u> B): <ul style="list-style-type: none"> <li>A) Patient has tried generic lenalidomide [<b>documentation required</b>]; AND</li> <li>B) Patient cannot take generic lenalidomide due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the brand and bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [<b>documentation required</b>].</li> </ul> </li> </ul>
<p><b>Rituxan®</b> (rituximab)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Rituxan (rituximab) is considered medically necessary when BOTH of the following are met:</b></p>

	<ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Documentation provided that the patient has the following: <ol style="list-style-type: none"> <li>A. Trial of AND cannot continue to use the alternative(s) due to a formulation difference in the inactive ingredient(s) which, according to the prescriber, would result in a significant allergy or serious adverse reaction for <b>ALL</b> of the following: <ol style="list-style-type: none"> <li>i. <b>Riabni (rituximab-arrx)</b> [may require prior authorization]</li> <li>ii. <b>Ruxience (rituximab-pvvr)</b> [may require prior authorization]</li> <li>iii. <b>Truxima (rituximab-abbs)</b> [may require prior authorization]</li> </ol> </li> </ol> </li> </ol>
<p><b>Rituxan Hycela™</b> (rituximab and hyaluronidase human)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Rituxan Hycela (rituximab and hyaluronidase human) is considered medically necessary when BOTH of the following are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Documentation of <b>ONE</b> of the following: <ol style="list-style-type: none"> <li>A. <b>BOTH</b> of the following: <ol style="list-style-type: none"> <li>i. Has received at least one dose of intravenous rituximab</li> <li>ii. Trial of AND cannot continue to use the alternative(s) due to a formulation difference in the inactive ingredient(s) which, according to the prescriber, would result in a significant allergy or serious adverse reaction for <b>ALL</b> of the following: <ol style="list-style-type: none"> <li>a. <b>Riabni (rituximab-arrx)</b> [may require prior authorization]</li> <li>b. <b>Ruxience (rituximab-pvvr)</b> [may require prior authorization]</li> <li>c. <b>Truxima (rituximab-abbs)</b> [may require prior authorization]</li> </ol> </li> </ol> </li> <li>B. Currently receiving Rituxan Hycela</li> </ol> </li> </ol>
<p><b>Scemblix</b> (asciminib tablets)</p>	<p><b><u>Individual and Family Plans</u></b></p> <p><b>Scemblix is considered medically necessary when BOTH of the following are met (1 and 2):</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. <b>ONE</b> of the following (a or b): <ol style="list-style-type: none"> <li>a. For <b><u>Chronic Myeloid Leukemia (CML)</u></b>, <b>ONE</b> of the following: <ol style="list-style-type: none"> <li>A. Patient meets <b>BOTH</b> of the following: <ol style="list-style-type: none"> <li>i. Patient meets <b>ONE</b> of the following: <ol style="list-style-type: none"> <li>a. Trial of, contraindication, significant intolerance to <b>imatinib</b> <i>Note:</i> Prior use of brand Gleevec and Imkeldi also counts.</li> <li>b. Patient has newly diagnosed disease</li> <li>c. Patient has intermediate- to high-risk chronic phase CML, accelerated CML or blast phase CML</li> <li>d. Patient has tried at least one other tyrosine kinase inhibitor for CML</li> </ol> </li> </ol> </li> </ol> </li> </ol> </li> </ol>

	<p><u>Note:</u> Examples of tyrosine inhibitors include: dasatinib, Bosulif, nilotinib, and Iclusig.</p> <p>e. Patient has a resistance mutation in which imatinib should not be used</p> <p>ii. Patient meets <b>ONE</b> of the following:</p> <p>a. Trial of, contraindication, significant intolerance to <b>dasatinib</b></p> <p><u>Note:</u> Prior use of Phyrago or Sprycel also counts.</p> <p>b. Patient has tried at least two other tyrosine kinase inhibitors for CML</p> <p><u>Note:</u> Examples of tyrosine kinase inhibitors include: dasatinib, Bosulif, nilotinib, and Iclusig.</p> <p>c. Patient has a history of serious, chronic lung disease or has had or is at risk of pleural effusion</p> <p><u>Note:</u> Examples of lung disease are pulmonary arterial hypertension and interstitial pneumonitis.</p> <p>d. Patient is at risk of bleeding</p> <p><u>Note:</u> Examples of increased risk of bleeding are if a patient has thrombocytopenia or is receiving a medication that inhibits platelet function or anticoagulants.</p> <p>e. Patient has a resistance mutation in which dasatinib should not be used</p> <p>B. Patient is currently receiving Scemblix</p> <p>C. Patient has the <i>T315I</i> mutation</p> <p>b. <b>All Other Conditions.</b> Approve Scemblix if the patient meets the Oncology Medications criteria above the table</p>
<p><b>Sprycel</b> (dasatinib tablets)</p>	<p><b><u>Employer Plans and Individual and Family Plan</u></b></p> <p><b>Sprycel is considered medically necessary when BOTH of the following are met:</b></p> <p>1. When the Oncology Medications criteria above the table are met</p> <p>2. <b>BOTH</b> of the following (A <u>and</u> B):</p> <p>A) Patient has tried generic dasatinib ; AND</p> <p>B) Patient cannot take generic dasatinib due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the brand and bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction.</p>
<p><b>Sutent</b> (sunitinib)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Sutent (sunitinib) is considered medically necessary when BOTH of the following are met:</b></p> <p>1. When the Oncology Medications criteria above the table are met</p> <p>2. Documented trial of <b><u>sunitinib</u></b> (the bioequivalent generic product) AND cannot take due to a formulation difference in the inactive ingredient(s) which would result in a significant allergy or serious adverse reaction</p>
<p><b>Talzenna®</b> (talazoparib)</p>	<p><b><u>Employer Plans and Individual and Family Plan:</u></b></p>

	<p><b>Talzenna (talazoparib) is considered medically necessary when BOTH of the following are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. <b>ONE</b> of the following: <ol style="list-style-type: none"> <li>A. For <b><u>BRCA-mutated, recurrent or metastatic Breast Cancer</u></b>, <b>ONE</b> of the following: <ol style="list-style-type: none"> <li>i. Documented trial of, contraindication, intolerance to <b>Lynparza (olaparib)</b> [may require prior authorization]</li> <li>ii. Currently receiving Talzenna</li> </ol> </li> <li>B. For <b><u>Prostate Cancer</u></b>, <b>ONE</b> of the following: <ol style="list-style-type: none"> <li>i. For BRCA-mutated prostate cancer, documented trial of, contraindication, intolerance to <b>Lynparza (Olaparib)</b> [may require prior authorization]</li> <li>ii. Patient has a homologous recombination repair (HRR) mutation OTHER THAN a BRCA-mutation (i.e., patient does not have a BRCA mutation)</li> <li>iii. Currently receiving Talzenna</li> </ol> </li> <li>C. <b>All Other Conditions.</b> Approve Talzenna if the patient meets the Oncology Medications criteria above the table</li> </ol> </li> </ol>
<p><b>Tarceva®</b> (erlotinib)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Tarceva (erlotinib) is considered medically necessary when BOTH of the following are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Documented trial of <b><u>erlotinib</u></b> (the bioequivalent generic product) [may require prior authorization] AND cannot take due to a formulation difference in the inactive ingredient(s) which would result in a significant allergy or serious adverse reaction</li> </ol>
<p><b>Targretin®</b> (bexarotene)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Targretin (bexarotene) is considered medically necessary when BOTH of the following are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Documented trial of <b><u>bexarotene</u></b> (the bioequivalent generic product) AND cannot take due to a formulation difference in the inactive ingredient(s) which would result in a significant allergy or serious adverse reaction</li> </ol>
<p><b>Tasigna</b> (nilotinib hcl capsules)</p>	<p><b><u>Employer Group Plans</u></b></p> <p><b>Tasigna is considered medically necessary when BOTH of the following are met (1 and 2):</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Patient meets <b>BOTH</b> of the following (A <u>and</u> B): <ol style="list-style-type: none"> <li>A) Patient has tried generic nilotinib hcl capsules; AND</li> <li>B) Patient cannot take generic nilotinib hcl capsules due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the brand and bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction.</li> </ol> </li> </ol>

	<p><b><u>Individual and Family Plans:</u></b></p> <p><b>Tasigna is considered medically necessary when BOTH of the following are met (1 and 2):</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. <b>ONE</b> of the following (A or B):       <ol style="list-style-type: none"> <li>A. For <b><u>Chronic Myeloid Leukemia (CML)</u></b>, <b>ONE</b> of the following:           <ol style="list-style-type: none"> <li>i. Trial of, contraindication, significant intolerance to <b>ONE</b> of the following:               <ol style="list-style-type: none"> <li>a. <b>imatinib</b></li> <li>b. <b>dasatinib</b></li> </ol> <p><u>Note:</u> Prior use of brand Gleevec, Imkeldi Phyrago, or Sprycel counts.</p> </li> <li>ii. Patient is currently receiving Tasigna</li> <li>iii. Patient is less than 18 years of age with accelerated phase CML</li> <li>iv. Patient meets <b>BOTH</b> of the following:               <ol style="list-style-type: none"> <li>a. Patient meets <b>ONE</b> of the following:                   <ol style="list-style-type: none"> <li>i. Patient has intermediate- to high-risk disease chronic phase CML</li> <li>ii. Patient has accelerated phase CML or blast phase CML</li> </ol> </li> <li>b. Patient meets <b>ONE</b> of the following:                   <ol style="list-style-type: none"> <li>i. Patient has a history of serious, chronic lung disease or has had or is at risk of pleural effusion</li> </ol> <p><u>Note:</u> Examples of lung disease, pulmonary arterial hypertension, and interstitial pneumonitis.</p> </li> <li>ii. Patient is at risk of bleeding</li> </ol> <p><u>Note:</u> An example of a patient with an increased risk of bleeding is a patient with thrombocytopenia or with a medication that inhibits platelet function or anticoagulants.</p> </li> <li>v. Patient has a resistance mutation in which imatinib and dasatinib should not be used</li> </ol> </li> </ol> </li> <li>B. <b>All Other Conditions.</b> Approve Tasigna if the patient meets the Oncology Medications criteria above the table</li> </ol>
<p><b>Temodar®</b> (temozolomide)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Temodar (temozolomide) is considered medically necessary when BOTH of the following are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Documented trial of <b><u>temozolomide</u></b> (the bioequivalent generic product) AND cannot take due to a formulation difference in the inactive ingredient(s) which would result in a significant allergy or serious adverse reaction</li> </ol>
<p><b>Tepylute</b> (thiotepa)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p>

	<p><b>Tepylute (thiotepa) is considered medically necessary when BOTH of the following are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Patient has tried and cannot take one of the following: generic thiotepa injection or Tepadina</li> </ol>
<p><b>Tykerb®</b> (lapatinib)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Tykerb (lapatinib) is considered medically necessary when BOTH of the following are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Documented trial of <b>lapatinib</b> (the bioequivalent generic product) AND cannot take due to a formulation difference in the inactive ingredient(s) which would result in a significant allergy or serious adverse reaction</li> </ol>
<p><b>Vabrinty</b> (leuprolide acetate for injectable suspension)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Vabrinty (leuprolide acetate for injectable suspension) is considered medically necessary when BOTH of the following are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Eligard is not available on the market per a verifiable source (for example, FDA Drug Shortage database or ASHP Drug Shortage list).</li> </ol>
<p><b>Vectibix</b> (panitumumab intravenous infusion)</p>	<p><b><u>Cigna Pathwell Specialty Drug List Plans</u></b></p> <p><b>Vectibix is considered medically necessary when BOTH of the following are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. ONE of the following: <ol style="list-style-type: none"> <li>A. For <b>Colon or Rectal Cancer</b>, documentation of <b>ONE</b> of the following: <ol style="list-style-type: none"> <li>i. According to the prescriber, the patient has experienced an inadequate response or significant intolerance, has a contraindication for Erbitux (cetuximab intravenous infusion); OR</li> <li>ii. Patient has been started on Vectibix or has already been started on therapy with Lumakras; OR</li> <li>iii. Patient had a serious infusion reaction to Erbitux; OR</li> <li>iv. <b>ONE</b> of the following (i or ii): <ol style="list-style-type: none"> <li>a. According to the prescriber, patient lives in high endemic rates of alpha-gal; OR</li> <li>b. Patient has known alpha-gal positivity</li> </ol> </li> </ol> </li> <li>B. <b>All Other Conditions.</b> Approve Vectibix if the patient meets the Oncology Medications criteria above the table</li> </ol> </li> </ol>
<p><b>Vegzelma</b> (bevacizumab-adcd)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Vegzelma is considered medically necessary when BOTH of the following are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Patient meets <b>BOTH</b> of the following (A and B):</li> </ol>

	<p>A. Patient has tried <b>BOTH</b> of the following [<b>documentation required</b>]</p> <ul style="list-style-type: none"> <li>i. <b>Mvasi (bevacizumab-awwb)</b> [may require prior authorization]</li> <li>ii. <b>Zirabev (bevacizumab-bvzr)</b> [may require prior authorization]</li> </ul> <p>B. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction</p>
<p><b>Votrient®</b> (pazopanib)</p>	<p><b><u>Employer Plans and Individual and Family Plan</u></b></p> <p><b>Votrient (pazopanib) is considered medically necessary when BOTH of the following are met:</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Documented trial of <b>pazopanib</b> (the bioequivalent generic product) [may require prior authorization] AND cannot take due to a formulation difference in the inactive ingredient(s) which would result in a significant allergy or serious adverse reaction</li> </ul>
<p><b>Xeloda®</b> (capecitabine)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Xeloda (capecitabine) is considered medically necessary when BOTH of the following are met:</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Documented trial of <b>capecitabine</b> (the bioequivalent generic product) AND cannot take due to a formulation difference in the inactive ingredient(s) which would result in a significant allergy or serious adverse reaction</li> </ul>
<p><b>Yonsa®</b> (abiraterone)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Yonsa (abiraterone) is considered medically necessary when BOTH of the following are met:</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>A. For <b><u>Prostate Cancer – Metastatic, Castration-Resistant,</u></b> documentation of <b>ONE</b> of the following: <ul style="list-style-type: none"> <li>i. Documented trial of, contraindication, or intolerance to <b>generic abiraterone</b></li> <li>ii. Patient has been started on therapy with Yonsa</li> </ul> </li> <li>B. <b>All Other Conditions.</b> Approve Yonsa if the patient meets the Oncology Medications criteria above the table</li> </ul> </li> </ul>
<p><b>Zykadia</b> (ceritinib)</p>	<p><b><u>Employer Plans:</u></b></p> <p><b>Zykadia (ceritinib) is considered medically necessary when BOTH of the following are met:</b></p> <ul style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. ONE of the following:</li> </ul>

	<p>A. For <b><u>Non-Small Cell Lung Cancer – Anaplastic Lymphoma Kinase (ALK)-positive</u></b>, documentation of <b>ONE</b> of the following:</p> <ol style="list-style-type: none"> <li>1. Patient has tried Alecensa or Alunbrig</li> <li>2. Patient is currently receiving Zykadia</li> </ol> <p>B. <b>All Other Conditions.</b> Approve Zykadia if the patient meets the Oncology Medications criteria above the table</p>
<p><b>Zytiga®</b> (abiraterone)</p>	<p><b><u>Employer Plans and Individual and Family Plans</u></b></p> <p><b>Zytiga (abiraterone) is considered medically necessary when BOTH of the following are met:</b></p> <ol style="list-style-type: none"> <li>1. When the Oncology Medications criteria above the table are met</li> <li>2. Documented trial of <b><u>abiraterone</u></b> (the bioequivalent generic product) AND cannot take due to a formulation difference in the inactive ingredient(s) which would result in a significant allergy or serious adverse reaction</li> </ol>

When coverage is available and medically necessary, the dosage, frequency, duration of therapy, and site of care should be reasonable, clinically appropriate, and supported by evidence-based literature and adjusted based upon severity, alternative available treatments, and previous response to therapy.

Receipt of sample product does not satisfy any criteria requirements for coverage.

## Background

### FDA Approved Indication

- **Drugs**  
Drugs@FDA.  
<http://www.accessdata.fda.gov/scripts/cder/drugsatfda/>
- **Biologics**  
Licensed Biological Products with Supporting Documents.  
<http://www.fda.gov/BiologicsBloodVaccines/ucm133705.htm>

### Professional Societies/Organizations

- National Comprehensive Cancer Network (NCCN) Clinical Practice Guidelines in Oncology (NCCN Guidelines®)  
[Available with free subscription]  
[http://www.nccn.org/professionals/physician\\_gls/f\\_guidelines.asp](http://www.nccn.org/professionals/physician_gls/f_guidelines.asp)
- The NCCN Drugs & Biologics Compendium (NCCN Compendium®)  
[available with paid subscription]  
[http://www.nccn.org/professionals/drug\\_compendium/content/contents.asp](http://www.nccn.org/professionals/drug_compendium/content/contents.asp)

The National Comprehensive Cancer Network® (NCCN®) is an organization of cancer centers, developing treatment guidelines for most cancers. The NCCN Clinical Practice Guidelines in Oncology (NCCN Guidelines®) present evidenced-based recommendations for the diagnosis and treatment of cancer and cancer care supportive therapies. NCCN provides the following definitions for their categories of recommendations:<sup>2</sup>

- Category 1: Based upon high-level evidence, there is uniform NCCN consensus that the intervention is appropriate;

- Category 2A: Based upon lower-level evidence, there is uniform NCCN consensus that the intervention is appropriate;
- Category 2B: Based upon lower-level evidence, there is NCCN consensus that the intervention is appropriate;
- Category 3: Based upon any level of evidence, there is major NCCN disagreement that the intervention is appropriate.

For the 'uniform NCCN consensus' defined in Category 1 and Category 2A, a majority Panel vote of at least 85% is required. For the 'NCCN consensus' defined in Category 2B, a Panel vote of at least 50% (but less than 85%) is required. Lastly, for recommendations where there is strong Panel disagreement regardless of the quality of the evidence, NCCN requires a vote from at least three Panel Members (representing at least three different Member Institutions) to include and designate a recommendation as Category 3. The large majority of the recommendations put forth in the Guidelines are Category 2A. Where categories are not specified within the Guidelines, the default designation for the recommendation is Category 2A.<sup>2</sup>

The NCCN Drugs & Biologics Compendium (NCCN Compendium®) includes FDA Approved Indications of cancer and cancer support medications and recommended non-FDA approved uses based upon the recommendations contained within the NCCN Guidelines.

## References

1. U.S. Food and Drug Administration. Drugs@FDA. U.S. Department of Health & Human Services: <http://www.accessdata.fda.gov/scripts/cder/drugsatfda/>.
2. National Comprehensive Cancer Network. Retrieved from <https://www.nccn.org>.
3. The NCCN Chronic Myeloid Leukemia Clinical Practice Guidelines in Oncology. © 2025 National Comprehensive Cancer Network, Inc. Available at: <http://www.nccn.org>.
4. U.S. Food and Drug Administration. Licensed Biological Products with Supporting Documents. U.S. Department of Health & Human Services: <http://www.fda.gov/BiologicsBloodVaccines/ucm133705.htm>.

## Revision Details

Type of Revision	Summary of Changes	Date
Selected Revision	<p><b>Fruzaqla</b> Appendiceal, Colon or Rectal Cancer: <b>Added</b> preferred product step requirement through Lonsurf for Employer Plans</p> <p><b>Krazati</b> <b>Added</b> <i>has brain metastases</i> exception to the sotorasib (Lumakras) preferred product step requirement</p>	5/15/2024
Selected Revision	<p><b>Augtyro</b> ROS1-positive non-small cell lung cancer: <b>Added</b> preferred product step requirement through Rozlytrek for Employer Plans</p> <p><b>Abraxane and Paclitaxel albumin-bound</b></p>	6/1/2024

	<b>Updated</b> Abraxane and Paclitaxel albumin-bound preferred product requirement criteria on Cigna Pathwell Specialty Drug List Plans	
Selected Revision	<p><b>Alunbrig/Zykadia</b> Non-Small Cell Lung Cancer – anaplastic lymphoma kinase (<i>ALK</i>)-positive: <b>Added</b> preferred product step requirement through Alecensa for Employer and Individual and Family Plans</p> <p><b>Votrient</b> <b>Added</b> preferred product step requirement through generic pazopanib for Employer Plans</p> <p><b>Braftovi</b> Melanoma, unresectable or metastatic, treatment of <i>BRAF</i> V600 mutation-positive: <b>Added</b> preferred product step requirement through Tafinlar or Zelboraf on Employer plans</p> <p><b>Mektovi</b> Melanoma, unresectable or metastatic, treatment of <i>BRAF</i> V600 mutation-positive: <b>Added</b> preferred product step requirement through Cotellic or Mekinist for Employer plans</p> <p><b>Bosulif</b> Chronic Myeloid Leukemia (CML), Philadelphia Chromosome Positive: <b>Updated</b> preferred product criteria to add Scemblix and Tasigna as step requirement options, <b>Updated</b> preferred product step requirement exceptions</p> <p><b>Gleevec</b> <b>Updated</b> preferred product step through generic imatinib requirement criteria</p> <p><b>Iclusig</b> Chronic Myeloid Leukemia, Philadelphia Chromosome Positive: <b>Updated</b> preferred product criteria to add Scemblix and Tasigna as step requirement options, <b>Updated</b> step requirement from requiring “ONE” to requiring “TWO” preferred products for Employer and Individual and Plans, <b>Updated</b> exceptions to the step requirement for Employer plans and Individual and Family Plans Acute Lymphoblastic Leukemia, Philadelphia Chromosome Positive: <b>Added</b> preferred product step requirement through generic imatinib or Sprycel for Iclusig</p> <p><b>Scemblix</b> Chronic Myeloid Leukemia (CML), Philadelphia Chromosome Positive: <b>Removed</b> Scemblix</p>	7/1/2024

	<p>preferred product step requirement on Employer Plans, <b>Updated</b> step requirement from requiring "ONE" to requiring "TWO" preferred products for Individual and Plans, <b>Updated</b> exceptions to the step requirement for Individual and Family Plans</p> <p><b>Tasigna</b> Chronic Myeloid Leukemia (CML), Philadelphia Chromosome Positive: <b>Updated</b> exceptions to the step requirement for Employer and Individual and Family Plans</p>	
Selected Revision	<p><b>Tecentriq</b> Non-Small Cell Lung Cancer – Advanced or Metastatic, Squamous or Non-Squamous Cell Disease: <b>Updated</b> Tecentriq preferred product criteria: changed "initial therapy" to "first-line therapy"; added "patient has a performance status of 3" as an exception to the preferred product Keytruda step requirement</p>	8/1/2024
Selected Revision	<p><b>Pomalyst</b> <b>Removed</b> Pomalyst preferred product criteria requirement.</p>	9/1/2024
Selected Revision	<p><b>Anktiva</b> <b>Added</b> Anktiva preferred product criteria requirement for Employer Plans and Individual and Family Plans</p> <p><b>Besremi</b> <b>Updated from</b> "Employer Plans and" to "Employer Plans and Individual and Family Plans"</p> <p><b>Docivyx</b> <b>Added</b> Docivyx preferred product criteria requirement for Employer Plans and Individual and Family Plans</p> <p><b>Yonsa</b> <b>Added "Prostate Cancer – Metastatic, Castration-Resistant"</b> to the preferred product criteria for Employer Plans and Individual and Family Plans</p> <p><b>Sandostatin LAR Depot.</b> <b>Removed</b> criteria for Sandostatin LAR Depot for Employer Plans and Individual and Family Plan</p> <p><u>Effective 1/1/2025:</u> <b>Keytruda</b> <b>Added</b> Keytruda preferred product criteria requirement for Cigna Pathwell Specialty Drug List Plans.</p>	10/15/2024

	<p><b>Lupron Depot</b>  <b>Added</b> Lupron Depot preferred product criteria for Employer Plans and Individual and Family Plans.</p> <p><b>Onivyde</b>  <b>Added</b> Onivyde preferred product criteria for Employer Plans and Individual and Family Plans.</p> <p><b>Opdivo</b>  <b>Added</b> Opdivo preferred product criteria requirement for Cigna Pathwell Specialty Drug List Plans.</p> <p><b>Orgovyx</b>  <b>Updated from</b> "Trial of, contraindication, or intolerance to ONE of the following: Eligard [may require prior authorization], Firmagon [may require prior authorization], Lupron Depot [may require prior authorization], Trelstar [may require prior authorization]" to "Patient has tried ONE of the following: Eligard [may require prior authorization], Firmagon [may require prior authorization], Trelstar [may require prior authorization]" for Individual and Family Plans.</p> <p><b>Vectibix</b>  <b>Added</b> Vectibix preferred product criteria requirement for Cigna Pathwell Specialty Drug List Plans.</p> <p><b>Votrient</b>  <b>Added</b> Votrient preferred product criteria for Individual and Family Plans.</p>	
Selected Revision	<p><b>Ibrance.</b>  <b>Added</b> "Patient will be using Ibrance in combination with Itovebi"</p> <p><b>Effective 1/15/2025</b>  <b>Scemblix.</b>  Added "Patient has newly diagnosed disease" option under generic imatinib criteria</p>	12/5/2024
Selected Revision	<p><b>Bosulif</b>  <b>Employer Plans:</b>  <b>Updated from</b> 'Sprycel [may require prior authorization]' <b>to</b> 'generic dasatinib'  <b>Updated from</b> '<u>Note</u>: Prior use of Gleevec or Phyrago (dasatinib) counts.' <b>to</b> '<u>Note</u>: Prior use of Gleevec (imatinib), Phyrago (dasatinib), or Sprycel (dasatinib) counts.'</p> <p><b>Individual and Family Plans:</b></p>	1/1/2025

	<p><b>Added</b> `generic dasatinib or' to `Sprycel [may require prior authorization]`</p> <p><b>Keytruda</b>  <b>Updated</b> preferred product criteria <b>from</b> "Cigna Pathwell Specialty Drug List Plans" <b>to</b> "Employer Plans and Individual and Family Plans"</p> <p><b>Updated from</b> "Tumor is tumor mutational burden-high (TMB-H) [<math>\geq 10</math> mutations/megabase]" <b>to</b> "Tumor is tumor mutational burden-high (TMB-H) [<math>\geq 10</math> mutations/megabase] "</p> <p><b>Added</b> patient has recurrent or metastatic disease, tumor is programmed death-ligand 1 positive (combined positive score [CPS] <math>\geq 1</math>), and medication is used as subsequent therapy as new option for approval.</p> <p><b>Iclusig</b>  <b>Employer Plans:</b>  <b>Updated from</b> `Sprycel [may require prior authorization]` <b>to</b> `generic dasatinib`  <b>Updated from</b> `Note: Prior use of Gleevec or Phyrago (dasatinib) counts.` <b>to</b> `Note: Prior use of Gleevec (imatinib), Phyrago (dasatinib), or Sprycel (dasatinib) counts.`  <b>Individual and Family Plans: Added</b> `generic dasatinib` or' <b>to</b> `Sprycel [may require prior authorization]`</p> <p><b>Lanreotide acetate (by Cipla)</b>  Added "Effective 1/1/2025 through 2/15/2025" to criteria</p> <p><b>Opdivo</b>  <b>Updated</b> preferred product criteria <b>from</b> "Cigna Pathwell Specialty Drug List Plans" <b>to</b> "Employer Plans and Individual and Family Plans"  <b>Added</b> patient has recurrent or metastatic non-keratinizing disease and medication is used for subsequent therapy as new option for approval</p> <p><b>Scemblix</b>  <b>Individual and Family Plans: Updated from</b> Trial of, contraindication, significant intolerance to <b>Sprycel</b> <b>to</b> 'Trial of, contraindication, significant intolerance to <b>generic dasatinib or Sprycel</b>'</p> <p><b>Sprycel</b>  <b>Added</b> preferred product preferencing criteria for Employer Plans</p> <p><b>Tasigna</b>  <b>Employer Plans:</b></p>	
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	<p><b>Updated from</b> `Sprycel [may require prior authorization]` <b>to</b> `generic dasatinib`  <b>Updated from</b> `Note: Prior use of Gleevec or Phyrago (dasatinib) counts.` <b>to</b> `Note: Prior use of Gleevec (imatinib), Phyrago (dasatinib), or Sprycel (dasatinib) counts.`  <b>Individual and Family Plans: Added</b> `generic dasatinib or` <b>to</b> `Sprycel [may require prior authorization]`</p> <p><b>Vectibix</b>  <b>Added</b> "According to the prescriber, patient lives in high endemic rates of alpha-gal" or "patient has known alpha-gal positivity"</p>	
Selected Revision	<p><b>Abraxane intravenous infusion.</b>  <b>Added</b> "All Other Conditions. Approve Abraxane intravenous infusion if the patient meets the Oncology Medications criteria above the table"</p> <p><b>Akeega.</b>  <b>Added</b> "All Other Conditions. Approve Akeega if the patient meets the Oncology Medications criteria above the table"</p> <p><b>Alunbrig.</b>  <b>Added</b> "All Other Conditions. Approve Alunbrig if the patient meets the Oncology Medications criteria above the table"</p> <p><b>Anktiva.</b>  <b>Added</b> "All Other Conditions. Approve Anktiva if the patient meets the Oncology Medications criteria above the table"</p> <p><b>Augtyro.</b>  <b>Added</b> "All Other Conditions. Approve Augtyro if the patient meets the Oncology Medications criteria above the table"</p> <p><b>Besremi.</b>  <b>Removed,</b> for polycythemia vera, Pegasys as an alternative option  <b>Added</b> "Documentation provided that the patient has:" to polycythemia vera criteria  <b>Added</b> "All Other Conditions. Approve Besremi if the patient meets the Oncology Medications criteria above the table"</p> <p><b>Bosulif.</b>  <b>Added</b> "All Other Conditions. Approve Bosulif if the patient meets the Oncology Medications criteria above the table"</p>	4/15/2025

	<p><b>Braftovi.</b>  <b>Added "All Other Conditions.</b> Approve Braftovi if the patient meets the Oncology Medications criteria above the table"</p> <p><b>Fruzaqla</b>  <b>Added "All Other Conditions.</b> Approve Fruzaqla if the patient meets the Oncology Medications criteria above the table"</p> <p><b>Fusilev.</b>  <b>Removed</b> criteria for Fusilev.</p> <p><b>Herceptin.</b>  <b>Removed</b> "Currently receiving Herceptin"</p> <p><b>Herceptin Hylecta.</b>  <b>Updated from</b> "Trial of AND cannot continue to use the alternative(s) due to a formulation difference in the inactive ingredient(s) which, according to the prescriber, would result in a significant allergy or serious adverse reaction to ONE of the following:Kanjinti (trastuzumab-anns) [may require prior authorization], Ogivri (trastuzumab-dkst) [may require prior authorization], Trazimera (trastuzumab-qyyp) [may require prior authorization]" <b>to</b> "Trial of, contraindication, or intolerance to ONE of the following: Kanjinti (trastuzumab-anns) [may require prior authorization], Ogivri (trastuzumab-dkst) [may require prior authorization], Trazimera (trastuzumab-qyyp) [may require prior authorization]"</p> <p><b>Updated from</b> "Unable to obtain or maintain intravenoud access" <b>to</b> "Patient is unable to obtain or maintain intravenous access"</p> <p><b>Herzuma.</b>  <b>Removed</b> "Currently receiving Herzuma"</p> <p><b>Ibrance.</b>  <b>Added</b> "For premenopausal patients using in combination with fulvestrant as subsequent therapy (not initial therapy), approve if the patient has tried Verzenio"  <b>Added "All Other Conditions.</b> Approve Ibrance if the patient meets the Oncology Medications criteria above the table"</p> <p><b>Iclusig.</b>  <b>Added "All Other Conditions.</b> Approve Iclusig if the patient meets the Oncology Medications criteria above the table"</p>	
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	<p><b>Jemperli.</b>  <b>Added</b>, for <u>Mismatch Repair Deficient (dMMR) or Microsatellite Instability-High (MSI-H) Solid Tumors – Monotherapy</u>, “Note: Examples of solid tumors include ampullary adenocarcinoma, biliary tract cancer, breast cancer, esophageal and esophagogastric junction cancer, gastric cancer, hepatocellular cancer, and ovarian cancer.”  <b>Added</b> “All Other Conditions (e.g., rectal cancer). Approve Jemperli if the patient meets the Oncology Medications criteria above the table”</p> <p><b>Keytruda.</b>  <b>Added</b> “<b>All Other Conditions.</b> Approve Keytruda if the patient meets the Oncology Medications criteria above the table”</p> <p><b>Khapzory.</b>  <b>Updated from</b> “Inability to obtain leucovorin injection due to a documented drug shortage (Food and Drug Administration [FDA] Drug Shortage database or American Society of Health-Systems Pharmacists [ASHP] Drug Shortage list” <b>to</b> “Meets ONE of the following: Patient has tried one generic levoleucovorin calcium injection or generic leucovorin injection; If the patient has already started on therapy with Khapzory, patient has tried generic levoleucovorin calcium injection.”</p> <p><b>Krazati.</b>  <b>Added</b> “<b>All Other Conditions.</b> Approve Krazati if the patient meets the Oncology Medications criteria above the table”</p> <p><b>Lanreotide acetate (by Cipla).</b>  <b>Removed</b> criteria for lanreotide acetate (by Cipla)</p> <p><b>Lupron Depot 7.5 mg, 22.5 mg, 30 mg, 45 mg.</b>  <b>All Other Conditions.</b> Approve Lupron Depot 7.5 mg, 22.5 mg, 30 mg, 45 mg if the patient meets the Oncology Medications criteria above the table</p> <p><b>Mektovi.</b>  <b>Added</b> “<b>All Other Conditions.</b> Approve Mektovi if the patient meets the Oncology Medications criteria above the table”</p> <p><b>Nilandron.</b>  <b>Removed</b> “trial of, contraindication, or intolerance to ONE of the following: Bicalutamide, Flutamide”</p> <p><b>Orgovyx.</b></p>	
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	<p><b>Added</b> to criteria "For Prostate Cancer"</p> <p><b>Onivyde.</b>  <b>Added "All Other Conditions.</b> Approve Onivyde if the patient meets the Oncology Medications criteria above the table"</p> <p><b>Ontruzant.</b>  <b>Removed</b> "Currently receiving Ontruzant"</p> <p><b>Opdivo.</b>  <b>Added "All Other Conditions.</b> Approve Opdivo if the patient meets the Oncology Medications criteria above the table"</p> <p><b>Orgovyx.</b>  <b>Added "All Other Conditions.</b> Approve Orgovyx if the patient meets the Oncology Medications criteria above the table"</p> <p><b>Paclitaxel albumin-bound intravenous infusion.</b>  <b>Added "All Other Conditions.</b> Approve paclitaxel albumin-bound intravenous infusion if the patient meets the Oncology Medications criteria above the table"</p> <p><b>Provenge.</b>  <b>Added "All Other Conditions.</b> <u>Approve Provenge if the patient meets the Oncology Medications criteria above the table"</u></p> <p><b>Scemblix.</b>  For the exception to the requirement of a trial of Sprycel, the requirement that the patient has tried at least "two" other tyrosine kinase inhibitors for CML was changed to at least "one" other tyrosine kinase inhibitor for CML.  <b>Added "All Other Conditions.</b> Approve Scemblix if the patient meets the Oncology Medications criteria above the table"</p> <p><b>Talzenna.</b>  <b>Updated from</b> "For <u>BRCA-mutated Prostate Cancer</u>, ONE of the following: Documented trial of, contraindication, intolerance to Lynparza (olaparib) [may require prior authorization], Currently receiving Talzenna" <b>to</b> "For BRCA-mutated prostate cancer, documented trial of, contraindication, intolerance to Lynparza (Olaparib) [may require prior authorization], Patient has a homologous recombination repair (HRR) mutation OTHER THAN</p>	
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	<p>a BRCA-mutation (i.e., patient does not have a BRCA mutation), Currently receiving Talzenna”  <b>Added “All Other Conditions.</b> Approve Talzenna if the patient meets the Oncology Medications criteria above the table”</p> <p><b>Tasigna.</b>  <b>Added “All Other Conditions.</b> Approve Tasigna if the patient meets the Oncology Medications criteria above the table”</p> <p><b>Tecentriq.</b>  <b>Added “All Other Conditions.</b> Approve Tecentriq if the patient meets the Oncology Medications criteria above the table”</p> <p><b>Vectibix</b>  <b>Added “All Other Conditions.</b> Approve Vectibix if the patient meets the Oncology Medications criteria above the table”</p> <p><b>Yonsa</b>  <b>Added “All Other Conditions.</b> Approve Yonsa if the patient meets the Oncology Medications criteria above the table”</p>	
Selected Revision	<p><b>Alunbrig.</b>  <b>Updated</b> from “Patient is currently receiving Alunbrig” <b>to</b> “Patient has already been started on therapy with Alunbrig”</p> <p><b>Boruzu.</b>  <b>Added</b> criteria for Boruzu</p> <p><b>Lupron Depot.</b>  <b>Removed</b> criteria for Lupron Depot.</p> <p><b>Tecentriq.</b>  <b>Removed</b> criteria for Tecentriq (Effective 4/15/2025)</p>	5/15/2025
Selected Revision	<p><b>Afinitor.</b>  <b>Removed</b> Afinitor criteria</p> <p><b>Bosulif.</b>  <b>Employer Plans</b>  <b>Updated from</b> “For <u>Chronic Myeloid Leukemia (CML), Philadelphia Chromosome Positive</u>, documentation of ONE the following: Trial of, contraindication, or significant intolerance to ONE of the following: Generic dasatinib, Generic imatinib, Scemblix [may require prior authorization],</p>	6/15/2025

	<p>Tasigna [may require prior authorization] <u>Note</u>: Prior use of Gleevec (imatinib), Phyrago (dasatinib), or Sprycel (dasatinib) counts" to "For <u>Chronic Myeloid Leukemia (CML), Philadelphia Chromosome Positive</u>, ONE the following: Trial of, contraindication, or significant intolerance to ONE of the following: dasatinib, imatinib, Danziten [may require prior authorization], Imkeldi [may require prior authorization], Scemblix [may require prior authorization], Tasigna [may require prior authorization]<u>Note</u>: Prior use of brand Gleevec, Phyrago, or Sprycel counts."</p> <p><b>Added</b> "Patient has a history of a serious, chronic lung disease or has had or is at risk of pleural effusion; <u>Note</u>: Examples of lung disease are pulmonary arterial hypertension and interstitial pneumonitis."</p> <p><b>Added</b> Danziten, Imkeldi to "Patient has a resistant mutation" criteria</p> <p><b>Bosulif</b>  <b>Individual and Family Plans</b>  <b>Updated from</b> "For <u>Chronic Myeloid Leukemia (CML), Philadelphia Chromosome Positive</u>, documentation of ONE the following: Trial of, contraindication, or significant intolerance to ONE of the following: generic imatinib [may require prior authorization], generic dasatinib or Sprycel [may require prior authorization]  <u>Note</u>: Prior use of Gleevec (imatinib), Imkeldi, or Phyrago (dasatinib) counts" to "For <u>Chronic Myeloid Leukemia (CML), Philadelphia Chromosome Positive</u>, ONE the following: Trial of, contraindication, or significant intolerance to ONE of the following: imatinib [may require prior authorization], dasatinib or Sprycel [may require prior authorization]  <u>Note</u>: Prior use of brand Gleevec, Imkeldi, or Phyrago also counts"</p> <p><b>Danziten.</b>  <b>Added</b> Danziten criteria.</p> <p><b>Hercessi.</b>  <b>Added</b> Hercessi criteria</p> <p><b>Iclusig.</b>  <b>Employer Plans</b>  <b>Updated from</b> "For <u>Chronic Myeloid Leukemia (CML), Philadelphia Chromosome Positive</u>, documentation of ONE of the following: Trial of, contraindication, significant intolerance to TWO of the following:</p>	
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	<p>Generic Dasatinib, generic imatinib, Scemblix [may require prior authorization], Tassigna [may require prior authorization] <u>Note</u>: Prior use of Gleevec (imatinib), Phyrago (dasatinib), or Sprycel (dasatinib) counts." <b>To</b> "</p> <p>For <u>Chronic Myeloid Leukemia (CML), Philadelphia Chromosome Positive</u>, ONE of the following: Trial of, contraindication, significant intolerance to TWO of the following: dasatinib, imatinib, Danziten [may require prior authorization], Imkeldi [may require prior authorization], Scemblix [may require prior authorization], Tassigna [may require prior authorization] <u>Note</u>: Prior use of brand Gleevec, Phyrago, or Sprycel also counts."</p> <p><b>Added</b> Danziten to tyrosine kinase inhibitor examples</p> <p><b>Added</b> Danziten, Imkeldi to "Patient has a resistant mutation" criteria</p> <p><b>Updated from</b> "For <u>Acute Lymphoblastic Leukemia (ALL), Philadelphia Chromosome-Positive</u>, documentation of ONE of the following: Trial of, contraindication, significant intolerance to ONE of the following: generic dasatinib, generic imatinib <u>Note</u>: Prior use of Gleevec (Imatiib), Phyrago (dasatinib), or Sprycel (dasatinib) counts." <b>To</b> "Trial of, contraindication, significant intolerance to ONE of the following: dasatinib, imatinib <u>Note</u>: Prior use of Gleevec , Imkeldi, Phyrago, or Sprycel also counts."</p> <p><b>Removed</b> examples of dasatinib products</p> <p><b>Added</b> "Patient is currently receiving therapy with Iclusig</p> <p><b>Iclusig</b>  <b>Individual and Family Plans.</b>  <b>Updated form</b> "For <u>Chronic Myeloid Leukemia (CML), Philadelphia Chromosome Positive</u>, documentation of ONE of the following:  Patient meets BOTH of the following: Patient meets ONE of the following: Trial of, contraindication, significant intolerance to generic imatinib [may require prior authorization] <u>Note</u>: Prior use of Gleevec (Imatinib) also counts." <b>To</b> "For <u>Chronic Myeloid Leukemia (CML), Philadelphia Chromosome Positive</u>, ONE of the following:  Patient meets BOTH of the following: Patient meets ONE of the following:  Trial of, contraindication, significant intolerance to imatinib [may require prior authorization] <u>Note</u>: Prior use of brand Gleevec or Imkeldi also counts.  <b>Added</b> Danziten to examples of tyrosine kinase inhibitors</p>	
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**Removed** "Patient is at risk of bleeding" with note  
**Updated from** "For Acute Lymphoblastic Leukemia (ALL), Philadelphia Chromosome-Positive, documentation of ONE of the following: According to the prescriber, patient has had a trial of, contraindication, significant intolerance to ONE of the following: generic imatinib [may require prior authorization], generic dasatinib or Sprycel [may require prior authorization] Note: Prior use of Gleevec (imatinib), or Phyrago (dasatiib) counts."  
**To** "For Acute Lymphoblastic Leukemia (ALL), Philadelphia Chromosome-Positive, ONE of the following: According to the prescriber, patient has had a trial of, contraindication, significant intolerance to ONE of the following: imatinib [may require prior authorization], dasatinib or Sprycel [may require prior authorization] Note: Prior use of Gleevec, Imkeldo, or Phyrago also counts."

**Imkeldi.**

**Added** Imkeldi criteria

**Scemblix.**

**Individual and Family Plan.**

**Updated from** "For Chronic Myeloid Leukemia (CML), Philadelphia Chromosome Positive, documentation of ONE of the following: Patient meets BOTH of the following: Patient meets ONE of the following: Trial of, contraindication, significant intolerance to generic imatinib Note: Prior use of Gleevec (imatinib) counts." **To** "For Chronic Myeloid Leukemia (CML), Philadelphia Chromosome Positive, ONE of the following: Patient meets BOTH of the following: Patient meets ONE of the following: Trial of, contraindication, significant intolerance to imatinib Note: Prior use of brand Gleevec and Imkeldi also counts."

**Added** Danziten to tyrosine kinase examples

**Tasigna**

**Employer Group Plans**

**Updated from** "For Chronic Myeloid Leukemia (CML), Philadelphia Chromosome Positive, documentation of ONE of the following: Trial of, contraindication, significant intolerance to ONE of the following: generic dasatinib, generic imatinib Note: Prior use of Gleevec (imatinib), Phyrago (dasatinib), or Sprycel (dasatinib) counts." **To** "For Chronic Myeloid Leukemia (CML), Philadelphia Chromosome Positive, ONE of the following: Trial of, contraindication, significant intolerance to ONE of the following: dasatinib, imatinib Note: Prior

	<p>use of brand Gleevec, Phyrago, or Sprycel also counts.”</p> <p><b>Tasigna</b>  <b>Individual and Family Plans</b>  <b>Updated from</b> “For <u>Chronic Myeloid Leukemia (CML), Philadelphia Chromosome Positive</u>, documentation of ONE of the following: Trial of, contraindication, significant intolerance to ONE of the following: generic imatinib [may require prior authorization], generic dasatinib or Sprycel [may require prior authorization] <u>Note</u>: Prior use of Gleevec (imatinib), or Phyrago (dasatinib) counts.” <b>To</b> “For <u>Chronic Myeloid Leukemia (CML), Philadelphia Chromosome Positive</u>, ONE of the following: Trial of, contraindication, significant intolerance to ONE of the following: imatinib [may require prior authorization], dasatinib or Sprycel [may require prior authorization] <u>Note</u>: Prior use of brand Gleevec, Imkeldi or Phyrago counts.”</p> <p><b>Ziihera.</b>  <b>Added</b> Ziihera criteria</p>	
Selected Revision	<p><b>Danziten.</b>  <b>Updated from</b> “Patient meets <b>ONE</b> of the following: Patient is at risk of bleeding <u>Note</u>: An example of a patient with an increased risk of bleeding as if a patient has thrombocytopenia or is receiving a medication that inhibits platelet function or anticoagulants; Patient has a prolonged QT interval or is at risk of developing QT interval prolongation” <b>to</b> “Patient meets ONE of the following: Patient has a history of serious, chronic lung disease or has had or is at risk of pleural effusion; OR <u>Note</u>: Examples of lung disease are pulmonary arterial hypertension and interstitial pneumonitis; Patient is at risk of bleeding; OR <u>Note</u>: An example of a patient with an increased risk of bleeding is a patient with thrombocytopenia or with medication use that inhibits platelet function or anticoagulants.”</p> <p><b>Opdivo Qvantig.</b>  <b>Added</b> Opdivo Qvantig criteria</p> <p><b>Rituxan.</b>  <b>Removed</b> “Currently receiving Rituxan”</p> <p><b>Ziihera.</b>  <b>Added</b> ‘may require prior authorization’ to: trastuzumab plus Perjeta, trastuzumab plus Tukysa</p>	7/1/2025
Selected Revision	<p><b>Anktiva.</b></p>	7/15/2025

	<p><b>Removed</b> Anktiva criteria.</p> <p><b>Alymsys.</b> <b>Removed</b> "Currently receiving Alymsys"</p> <p><b>Avastin.</b> <b>Removed</b> "Currently receiving Avastin"</p> <p><b>Vegzelma</b> <b>Removed</b> "Currently receiving Vegzelma"</p>	
Selected Revision	<p><b>Ensacove.</b> <b>Added</b> criteria for Ensacove for Individual and Family Plan</p> <p><b>Onivyde.</b> <b>Pancreatic Adenocarcinoma:</b> Removed the requirement that patient has been previously treated with fluoropyrimidine-based therapy without irinotecan. Added "according to prescriber patient is Eastern Cooperative Oncology Group Performance Status 0 or 1 and has tried a gemcitabine-based regimen" and "according to prescriber patient is Eastern Cooperative Oncology Group Performance Status 0 or 1 and has tried a flouropyrimidine-based regimen but without prior irinotecan" as options for approval.</p> <p><b>Ampullary adenocarcinoma:</b> Added ampullary adenocarcinoma as a new condition for approval.</p>	9/1/2025
Selected Revision	<p><b>Alunbrig.</b> <b>Removed</b> criteria for Alunbrig for Employer Plans</p> <p><b>Bosulif.</b> <b>Removed</b> "Philadelphia Chromosome Positive" <b>Updated from</b> "Tasigna" <b>to</b> "nilotinib" for Employer Plans for Chronic Myeloid Leukemia (CML); <b>Added</b> "Tasigna" to prior use counting</p> <p><b>Danziten.</b> <b>Removed</b> "Philadelphia Chromosome Positive" <b>Added</b> "nilotinib" as an alternative for Chronic Myeloid Leukemia (CML); <b>Added</b> "Tasigna" to prior use counting for Employer Plans</p> <p><b>Fruzaqla.</b> <b>Removed</b> criteria for Fruzaqla.</p> <p><b>Iclusig.</b> <b>Removed</b> "Philadelphia Chromosome Positive" <b>Updated from</b> "Tasigna" <b>to</b> "nilotinib" for Employer Plans for Chronic Myeloid Leukemia (CML); <b>Added</b> "Tasigna" to prior use counting</p>	10/1/2025

	<p><b>Added</b> for Acute Lymphoblastic Leukemia “Patient is ≥ 18 years of age, has newly diagnosed disease, and is taking the requested medication with chemotherapy; Patient has T315I-positive mutation”</p> <p><b>Ivra.</b> <b>Added</b> criteria for Ivra.</p> <p><b>Nilotinib.</b> <b>Added</b> criteria for Nilotinib for Individual and Family Plans</p> <p><b>Provence.</b> <b>Removed</b> criteria for Provence.</p> <p><b>Scemblix.</b> <b>Removed</b> “Philadelphia Chromosome Positive”</p> <p><b>Tasigna.</b> <b>Removed</b> “Philadelphia Chromosome Positive” <b>Updated</b> criteria to “Documented trial of <b>nilotinib</b> (the bioequivalent generic product) [may require prior authorization] AND cannot take due to a formulation difference in the inactive ingredient(s) which would result in a significant allergy or serious adverse reaction”</p> <p><b>Tepylute.</b> <b>Added</b> criteria for Tepylute (Effective 11/1/2025)</p> <p><b>Zykadia.</b> <b>Added</b> “Or Alunbrig” to the Zykadia step for Employer Plans</p>	
Selected Revision	<p><b>Akeega.</b> <b>Added</b> criteria for Akeega for Individual and Family Plan (Effective 1/1/2026)</p> <p><b>Avgemsi.</b> <b>Added</b> criteria for Avgemsi (Effective 1/15/2026)</p> <p><b>Bosulif.</b> <b>Chronic Myeloid Leukemia, Employer Plans</b> <b>Updated from</b> “Trial of, contraindication, or significant intolerance to ONE” <b>to</b> “ Trial of, contraindication, or significant intolerance to TWO” <b>Removed</b> “Patient meets <b>ONE</b> of the following: Patient has a history of a serious, chronic lung disease or has had or is at risk of pleural effusion; OR Patient is at risk of bleeding; OR Patient has a prolonged QT interval or is at risk of developing QT interval prolongation <b>Chronic Myeloid Leukemia, Individual and Family Plans</b></p>	12/1/2025

	<p><b>Added</b> criteria for dasatanib as a required alternative</p> <p><b>Acute Lymphoblastic Leukemia.</b>  <b>Added</b> Criteria for Acute Lymphoblastic Leukemia for Employer Plans and Individual and Family Plans</p> <p><b>Braftovi.</b>  <b>Added</b> criteria for Individual and Family Plan (Effective 1/1/2026)</p> <p><b>Danziten.</b>  <b>Chronic Myeloid Leukemia, Employer Plans</b>  <b>Removed</b> " Patient meets <b>BOTH</b> of the following: Patient meets <b>ONE</b> of the following: Patient has intermediate- to high-risk chronic phase CML, Patient has accelerated phase CML or blast phase CML; Patient meets <b>ONE</b> of the following: Patient has a history of serious, chronic lung disease or has had or is at risk of pleural effusion, Patient is at risk of bleeding</p> <p><b>Chronic Myeloid Leukemia, Individual and Family Plan</b>  <b>Removed</b> Sprycel as an alternative</p> <p><b>Fruzaqla.</b>  <b>Added</b> criteria for Fruzaqla for Individual and Family Plans (Effective 1/1/2026)</p> <p><b>Iclusig.</b>  <b>Chronic Myeloid Leukemia, Individual and Family Plan</b>  <b>Removed</b> Sprycel as an alternative  <b>Acute Lymphoblastic Leukemia, Individual and Family Plan</b>  <b>Removed</b> Sprycel as an alternative</p> <p><b>Mektovi.</b>  <b>Added</b> criteria for Individual and Family Plans (Effective 1/1/2026)</p> <p><b>Nilotinib.</b>  <b>Chronic Myeloid Leukemia, Individual and Family Plan</b>  <b>Removed</b> Sprycel as an alternative</p> <p><b>Nilotinib d-tartrate.</b>  <b>Added</b> criteria for nilotinib d-tartrate</p> <p><b>Onivyde.</b>  <b>Pancreatic Adenocarcinoma</b>  <b>Added</b> "According to the prescriber, the patient has baseline neuropathy"  <b>Ampullary Adenocarcinoma</b></p>	
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	<p><b>Added</b> "According to the prescriber, the patient has baseline neuropathy"</p> <p><b>Phyrago.</b> <b>Added</b> criteria for Phyrago for Individual and Family Plans</p> <p><b>Revlimid.</b> <b>Added</b> criteria for Revlimid for Employer Plans (Effective 2/1/2026)</p> <p><b>Scemblix</b> <b>Chronic Myeloid Leukemia, Individual and Family Plan</b> <b>Removed</b> Sprycel as an alternative</p> <p><b>Sprycel.</b> <b>Added</b> criteria for Sprycel for Individual and Family Plans</p> <p><b>Talzenna.</b> <b>Added</b> criteria for Talzenna for Individual and Family Plans (Effective 1/1/2026)</p> <p><b>Tasigna.</b> <b>Chronic Myeloid Leukemia, Individual and Family Plan</b> <b>Removed</b> Sprycel as an alternative</p>	
Selected Revision	<p><b>Bosulif.</b> <b>Added</b> capsules</p> <p><b>Docivyx.</b> <b>Added</b> "Patients with hypersensitivities to polysorbate 80"</p> <p><b>Vabrinty.</b> <b>Added</b> criteria for Vabrinty.</p>	1/1/2026
Selected Revision	<p><b>Jemperli.</b> <b>Removed</b> documentation requirements</p> <p><b>Kyxata.</b> <b>Added</b> criteria for Kyxata</p> <p><b>Vectibix.</b> <b>Updated from</b> "Patient has been started on Vectibix" <b>to</b> "Patient has been started on Vectibix or has already been started on therapy with Lumakras"</p>	2/15/2026
Selected Revision	<p><b>Allymsys, Avastin, Vegzelma</b> <b>Added</b> examples of formulation difference in the inactive ingredient(s) "[e.g., differences in stabilizing agent, buffering agent, and/or surfactant]"</p>	3/1/2026

	<p><b>Jobevne.</b> <b>Added</b> criteria for Jobevne</p>	
Selected Revision	<p><b>Abraxane intravenous infusion.</b> <b>Added</b> "Patient has tried paclitaxel intravenous infusion" for Breast Cancer, Cervical Cancer, Endometrial Cancer, Melanoma, Non-Small Cell Lung Cancer, and Ovarian Cancer.</p> <p><b>Akeega.</b> <b>Added</b> "Patient has BRCA2-mutated metastatic castration-sensitive prostate cancer</p> <p><b>Augtyro.</b> <b>Updated from</b> "If Augtyro has not been tried previously, approve if the patient has symptomatic systemic progression (multiple lesions) on Rozlytrek (entrectinib capsules and pellet packet), Xalkori (crizotinib capsules), Zykadia" <b>to</b> "If Augtyro has not been tried previously, approve if the patient has symptomatic systemic progression (multiple lesions) on Rozlytrek (entrectinib capsules and pellet packet), Xalkori (crizotinib capsules), or Ibtrozi (taletrectinib capsules)"</p> <p><b>Paclitaxel albumin-bound intravenous infusion.</b> <b>Added</b> "Patient has tried paclitaxel intravenous infusion" for Breast Cancer, Cervical Cancer, Endometrial Cancer, Melanoma, Non-Small Cell Lung Cancer, and Ovarian Cancer.</p> <p><b>Phyrago.</b> <b>Added</b> "Employer Group Plans" criteria <b>Added</b> "According to the prescriber, the patient requires co-administration with a histamine-2 (H2) antagonist or proton pump inhibitor (PPI). <u>Note:</u> Examples of H2 antagonists include famotidine, cimetidine, nizatidine. Examples of PPIs include omeprazole, esomeprazole, pantoprazole, rabeprazole"</p> <p><b>Ziihera.</b> <b>Removed</b> criteria for Ziihera.</p>	3/15/2026
Selected Revision	<p><b>Added</b> [documentation required] instructions</p> <p><b>Beizray.</b> <b>Added</b> criteria for Beizray</p> <p><b>Keytruda IV (pembrolizumab intravenous infusion).</b> <b>Updated from</b> "Patient has been started on Keytruda; Patient meets <b>ALL</b> of the following:</p>	4/1/2026

	<p>Patient has recurrent, unresectable, oligometastatic, or metastatic disease; The medication is used in combination with cisplatin and gemcitabine; According to the prescriber, the patient has inadequate efficacy, contraindication, or significant intolerance to <b>Loqtorzi (toripalimab intravenous infusion)</b> [may require prior authorization]; Patient meets <b>ALL</b> of the following: Patient has recurrent, unresectable, oligometastatic, or metastatic disease; Tumor is tumor mutational burden-high (TMB-H) [<math>\geq 10</math> mutations/megabase]; Medication is used for subsequent therapy; Patient meets <b>ALL</b> of the following: Patient has recurrent or metastatic disease; AND Tumor is programmed death-ligand 1 positive (combined positive score [CPS] <math>\geq 1</math>)” to “According to the prescriber, the patient has tried, and has had inadequate efficacy or significant intolerance or patient has a contraindication to Loqtorzi;; OR Patient has been started on Keytruda IV or Keytruda Qlex; OR Patient has a diagnosis of head and neck squamous cell carcinoma other than nasopharyngeal carcinoma”</p> <p><b>Keytruda Qlex.</b>  <b>Added</b> criteria for Keytruda Qlex “<b>Keytruda Qlex is considered medically necessary when BOTH of the following are met:</b> When the Oncology Medications criteria above the table are met <b>ONE</b> of the following: Patient has tried and cannot take Keytruda intravenous (IV) [may require prior authorization]; Patient is unable to obtain IV access”</p> <p><b>Revlimid.</b>  <b>Added</b> documentation requirements.</p> <p><b>Zykadia.</b>  <b>Removed</b> criteria for Zykadia for Individual and Family Plans</p> <p><b>Effective 4/15/2026</b>  <b>Gleostine.</b>  <b>Added</b> criteria for Gleostine for Employer Plans</p>	
Selected Revision	<p><b>Augtyro.</b>  <b>Updated from</b> “If Augtyro has not been tried previously, approve if the patient has symptomatic systemic progression (multiple lesions) on Rozlytrek (entrectinib capsules and pellet packet), Xalkori (crizotinib capsules), or Ibtrozi (taletrectinib capsules)” <b>to</b> “If Augtyro has not been tried previously, approve if the patient has progression on Rozlytrek (entrectinib capsules and pellet</p>	5/1/2026

	<p>packet), Xalkori (crizotinib capsules), or Ibtrozi (taletrectinib capsules)”</p> <p><b>Bosulif, Danziten, Iclusig, Nilotinib hcl, Scemblix, Tasigna.</b>  <b>Removed</b> “may require prior authorization” for imatinib and dasatinib for Individual and Family Plans</p> <p><b>Danziten.</b>  <b>Removed</b> “may require prior authorization” for nilotinib hcl for Employer Plans</p> <p><b>Gleevec.</b>  <b>Removed</b> “may require prior authorization” for imatinib</p> <p><b>Imkeldi.</b>  <b>Removed</b> “may require prior authorization” for imatinib for Individual and Family Plans</p> <p><b>Nexavar.</b>  <b>Removed</b> “may require prior authorization” for sorafenib.</p> <p><b>Pazopanib 400 mg.</b>  <b>Added</b> criteria for Pazopanib 400 mg</p> <p><b>Sprycel</b>  <b>Removed</b> “may require prior authorization” for dasatinib</p> <p><b>Sutent.</b>  <b>Removed</b> “may require prior authorization” for sunitinib</p> <p><b>Temodar.</b>  <b>Removed</b> “may require prior authorization” for temozolomide</p> <p><b>Tykerb.</b>  <b>Removed</b> “may require prior authorization” for lapatinib.</p> <p><b>Xeloda.</b>  <b>Removed</b> “may require prior authorization” for capecitabine</p> <p><b>Zytiga.</b>  <b>Removed</b> “may require prior authorization” for abiraterone</p>	
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The policy effective date is in force until updated or retired.

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