

CRITERIA FOR PRIOR AUTHORIZATION

Oncology - Auxiliary Treatment Agents

BILLING CODE TYPE For drug coverage and provider type information, see the [KMAP Reference Codes webpage](#).

MANUAL GUIDELINES Prior authorization will be required for all current and future dose forms available. All medication-specific criteria will be reviewed according to the criteria below.

Anti-thymocyte globulin (Atgam®)
Aprepitant (Cinvanti®, Emend® oral)
Darbepoetin alfa (Aranesp®)
Denosumab (Prolia®, Xgeva®)
(chemotherapy diluent) (Elliotts B® solution)
Epoetin alfa (Epogen®, Procrit®, Retacrit®)
Filgrastim (Neupogen®, Nivestym®, Zarxio®)
Fosaprepitant (Emend® IV)
Fosnetupitant/palonosetron (Akinzeo® IV)
Tbo-filgrastim (Granix®)
Glucarpidase (Voraxaze®)
Luspatercept (Reblozyl®)
Mesna (Mesnex®)
Netupitant/palonosetron (Akinzeo® oral)
Octreotide (Bynfezia™, Sandostatin®, Sandostatin® LAR)
Pegfilgrastim (Neulasta®, Neulasta Onpro®, Fulphila®, Nyvepria™, Udenyca®, Ziextenzo™)
[Piflufolastat F 18 \(Pylarify®\)](#)
Plerixafor (Mozobil®)
Rasburicase (Elitek®)
Rolapitant (Varubi®)
Sargramostim (Leukine®)

CRITERIA FOR INITIAL APPROVAL FOR ALL PRODUCTS (MUST MEET ALL OF THE FOLLOWING):

- Medication requested must be prescribed according to the FDA-approved indication, age, dose, and pre-requisite treatments located in the package insert.
- For all agents listed, the preferred PDL drug, if applicable, which treats the PA indication, is required unless the patient meets the non-preferred PDL PA criteria.

CRITERIA FOR RENEWAL FOR ALL PRODUCTS:

- Prescriber must attest that the patient has experienced a positive clinical response from continuous treatment with the requested medication and is able to tolerate therapy.
- Patient must continue to meet the criteria required for initial approval.

LENGTH OF APPROVAL: 12 MONTHS

FOR DRUGS THAT HAVE A CURRENT PA REQUIREMENT, BUT NOT FOR THE NEWLY APPROVED INDICATIONS, FOR OTHER FDA-APPROVED INDICATIONS, AND FOR CHANGES TO AGE REQUIREMENTS NOT LISTED WITHIN THE PA CRITERIA:

- **THE PA REQUEST WILL BE REVIEWED BASED UPON THE FOLLOWING PACKAGE INSERT INFORMATION: INDICATION, AGE, DOSE, AND ANY PRE-REQUISITE TREATMENT REQUIREMENTS FOR THAT INDICATION.**