

STEP THERAPY CRITERIA FOR MEDICARE PART B DRUGS

This list is current as of 4/1/2025 and pertains to the following Independent Health Medicare Advantage Plans for 2025:

Independent Health's Encompass 65[®] Basic (HMO)
Independent Health's Encompass 65[®] Core (HMO)
Independent Health's Encompass 65[®] Direct (HMO)
Independent Health's Medicare Passport[®] Access (PPO)
Independent Health's Medicare Passport[®] Connect (PPO)
Independent Health's Medicare Family Choice[®] (HMO I-SNP)
Independent Health's Assure Advantage (HMO C-SNP)
Independent Health's Medicare Advantage Employer Group Waiver Plans (EGWP)

In some cases, we require that you first try certain drugs to treat your medical condition before we will cover another drug for that condition. For example, if Drug X and Drug Y both treat your medical condition, we may not cover Drug Y unless you try Drug X first. If Drug X does not work for you, we will then cover Drug Y.

The therapy outlined in this document may also involve a combination of Part B and Part D drugs. For example, we may not cover a Part B drug unless you try a Part D drug first. Or, we may not cover a Part D drug unless you try a Part B drug first. This is dependent on the therapy described to treat your medical condition. This document contains the Step Therapy protocols for Medicare Part B drugs that are associated with the Independent Health Medicare Advantage Plans mentioned above.

If you have any questions, please contact our Medicare Member Services Department at 1-800-665-1502 or, for TTY users 711, October 1st – March 31st: Monday through Sunday from 8 a.m. to 8 p.m., April 1st – September 30th: Monday through Friday from 8 a.m. to 8 p.m.

The formulary may change at any time. You will receive notice when necessary.

Amvuttra

Products Affected

- AMVUTTRA SOLUTION PREFILLED SYRINGE
25 MG/0.5ML SUBCUTANEOUS

Details

Criteria	For approval, the patient must have tried and failed to have an adequate response to either Onpattro or Tegsedi. This specific requirement applies to new starts only.
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Aphexda

Products Affected

- APHEXDA SOLUTION RECONSTITUTED 62 MG SUBCUTANEOUS

Details

Criteria	For approval, the patient must have tried and failed to have an adequate response to plerixafor. This specific requirement applies to new starts only.
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4/1/2025

Apretude

Products Affected

- APRETUDE SUSPENSION EXTENDED RELEASE
600 MG/3ML INTRAMUSCULAR

Details

Criteria	For approval, patient must have tried and had an intolerance to or has a contraindication to emtricitabine/tenofovir disoproxil fumarate.
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4/1/2025

Asceniv

Products Affected

- ASCENIV SOLUTION 5 GM/50ML
INTRAVENOUS

Details

Criteria	For approval, patient must have tried and failed to have a response to another intravenous immunoglobulin (IVIG) product. This specific requirement applies to new starts only and does not apply to patients using Asceniv for any indication not shared with preferred IVIG products.
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Bendamustine

Products Affected

- TREANDA SOLUTION RECONSTITUTED 100 MG INTRAVENOUS
- TREANDA SOLUTION RECONSTITUTED 25 MG INTRAVENOUS

Details

Criteria	For approval of Treanda, the patient must have tried and failed to have an adequate response to Belrapzo, bendamustine (Apotex/Baxter) or Bendeka. This specific requirement applies to new starts only and does not apply to patients using Treanda for any indication not shared with preferred agents.
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Beovu

Products Affected

- BEOVU SOLUTION PREFILLED SYRINGE 6
MG/0.05ML INTRAVITREAL

Details

Criteria	For approval for a diagnosis of neovascular (wet) age-related macular degeneration or diabetic macular edema, the patient must have tried and failed to have an adequate response to bevacizumab (Avastin). This specific requirement applies to new starts only and does not apply to patients using Beovu for any off-label indication not shared with bevacizumab.
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4/1/2025

Bevacizumab

Products Affected

- ALYMSYS SOLUTION 100 MG/4ML
INTRAVENOUS
- ALYMSYS SOLUTION 400 MG/16ML
INTRAVENOUS
- AVASTIN SOLUTION 100 MG/4ML
INTRAVENOUS
- AVASTIN SOLUTION 400 MG/16ML
INTRAVENOUS
- VEGZELMA SOLUTION 100 MG/4ML
INTRAVENOUS
- VEGZELMA SOLUTION 400 MG/16ML
INTRAVENOUS

Details

Criteria	For approval of Alymsys, Avastin, or Vegzelma for oncology (cancer) indications, the patient must have tried and failed to have an adequate response to Mvasi or Zirabev. This specific requirement applies to new starts only. This requirement does not apply to patients using Avastin for ophthalmic (eye) indications or to patients using a bevacizumab agent for any indication not shared by Mvasi or Zirabev.
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Botulinum toxins

Products Affected

- DAXXIFY SOLUTION RECONSTITUTED 100 UNIT INTRAMUSCULAR
- DYSPORE SOLUTION RECONSTITUTED 300 UNIT INTRAMUSCULAR
- DYSPORE SOLUTION RECONSTITUTED 500 UNIT INTRAMUSCULAR
- MYOBLOC SOLUTION 10000 UNIT/2ML INTRAMUSCULAR
- MYOBLOC SOLUTION 2500 UNIT/0.5ML INTRAMUSCULAR
- MYOBLOC SOLUTION 5000 UNIT/ML INTRAMUSCULAR

Details

Criteria	For approval, the patient must have tried and failed to have an adequate response to Botox and Xeomin. This specific requirement applies to new starts only.
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Briumvi

Products Affected

- BRIUMVI SOLUTION 150 MG/6ML
INTRA VENOUS

Details

Criteria	For approval, the patient must have tried and failed to have an adequate response to Ocrevus. This specific requirement applies to new starts only.
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4/1/2025

Camcevi

Products Affected

- CAMCEVI PREFILLED SYRINGE 42 MG
SUBCUTANEOUS

Details

Criteria	For approval, the patient must have tried and failed to have an adequate response to either Eligard or Lupron. This specific requirement applies to new starts only.
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4/1/2025

Enjaymo

Products Affected

- ENJAYMO SOLUTION 1100 MG/22ML
INTRAVENOUS

Details

Criteria	For approval, the patient must have tried and failed to have an adequate response to Ruxience (or another rituximab product) or any other B-cell targeting therapy. This specific requirement applies to new starts only.
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4/1/2025

Erythropoietins

Products Affected

- EPOGEN SOLUTION 10000 UNIT/ML INJECTION
- EPOGEN SOLUTION 2000 UNIT/ML INJECTION
- EPOGEN SOLUTION 20000 UNIT/ML INJECTION
- EPOGEN SOLUTION 3000 UNIT/ML INJECTION
- EPOGEN SOLUTION 4000 UNIT/ML INJECTION
- PROCRIT SOLUTION 10000 UNIT/ML INJECTION
- PROCRIT SOLUTION 2000 UNIT/ML INJECTION
- PROCRIT SOLUTION 20000 UNIT/ML INJECTION
- PROCRIT SOLUTION 3000 UNIT/ML INJECTION
- PROCRIT SOLUTION 4000 UNIT/ML INJECTION
- PROCRIT SOLUTION 40000 UNIT/ML INJECTION

Details

Criteria	For approval, the patient must have tried and failed to have an adequate response to Retacrit. This specific requirement applies to new starts only and does not apply to patients using Epogen or Procrit for any indication not shared with Retacrit.
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Evkeeza

Products Affected

- EVKEEZA SOLUTION 1200 MG/8ML
INTRA VENOUS
- EVKEEZA SOLUTION 345 MG/2.3ML
INTRA VENOUS

Details

Criteria	For approval of Evkeeza, the patient must have tried and failed to have an adequate response to or have a contraindication to both a maximally-tolerated dose of a statin drug and a PCSK9 inhibitor. This specific requirement applies to new starts only.
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4/1/2025

Eylea

Products Affected

- EYLEA HD SOLUTION 8 MG/0.07ML INTRAVITREAL
- EYLEA SOLUTION 2 MG/0.05ML INTRAVITREAL
- EYLEA SOLUTION PREFILLED SYRINGE 2 MG/0.05ML INTRAVITREAL

Details

Criteria	For approval for a diagnosis of neovascular (wet) age-related macular degeneration, diabetic macular edema, or diabetic retinopathy in patients with diabetic macular edema, the patient must have tried and failed to have an adequate response to bevacizumab (Avastin). This specific requirement applies to new starts only.
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Feiba

Products Affected

- FEIBA SOLUTION RECONSTITUTED 1000 UNIT INTRAVENOUS
- FEIBA SOLUTION RECONSTITUTED 2500 UNIT INTRAVENOUS

Details

Criteria	For approval of Feiba, the patient must have tried and failed to have an adequate response to Hemlibra. This specific requirement applies to new starts only. This requirement does not apply to treatment of hemophilia B.
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Filgrastim

Products Affected

- GRANIX SOLUTION 300 MCG/ML SUBCUTANEOUS
- GRANIX SOLUTION 480 MCG/1.6ML SUBCUTANEOUS
- GRANIX SOLUTION PREFILLED SYRINGE 300 MCG/0.5ML SUBCUTANEOUS
- GRANIX SOLUTION PREFILLED SYRINGE 480 MCG/0.8ML SUBCUTANEOUS
- NEUPOGEN SOLUTION 300 MCG/ML INJECTION
- NEUPOGEN SOLUTION 480 MCG/1.6ML INJECTION
- NEUPOGEN SOLUTION PREFILLED SYRINGE 300 MCG/0.5ML INJECTION
- NEUPOGEN SOLUTION PREFILLED SYRINGE 480 MCG/0.8ML INJECTION
- NIVESTYM SOLUTION 300 MCG/ML INJECTION
- NIVESTYM SOLUTION 480 MCG/1.6ML INJECTION
- NIVESTYM SOLUTION PREFILLED SYRINGE 300 MCG/0.5ML INJECTION
- NIVESTYM SOLUTION PREFILLED SYRINGE 480 MCG/0.8ML INJECTION
- *releuko solution prefilled syringe 300 mcg/0.5ml subcutaneous*
- *releuko solution prefilled syringe 480 mcg/0.8ml subcutaneous*

Details

Criteria	For approval of Neupogen for any indication other than Hematopoietic Syndrome of Acute Radiation Syndrome, Granix, Nivestym, or Releuko, the patient must have tried and failed to have an adequate response to Zarxio. This specific requirement applies to new starts only and does not apply to patients using a non-preferred agent for any indication not shared with Zarxio.
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Growth hormone

Products Affected

- GENOTROPIN CARTRIDGE 12 MG SUBCUTANEOUS
- GENOTROPIN CARTRIDGE 5 MG SUBCUTANEOUS
- GENOTROPIN MINIQUICK PREFILLED SYRINGE 0.2 MG SUBCUTANEOUS
- GENOTROPIN MINIQUICK PREFILLED SYRINGE 0.4 MG SUBCUTANEOUS
- GENOTROPIN MINIQUICK PREFILLED SYRINGE 0.6 MG SUBCUTANEOUS
- GENOTROPIN MINIQUICK PREFILLED SYRINGE 0.8 MG SUBCUTANEOUS
- GENOTROPIN MINIQUICK PREFILLED SYRINGE 1 MG SUBCUTANEOUS
- GENOTROPIN MINIQUICK PREFILLED SYRINGE 1.2 MG SUBCUTANEOUS
- GENOTROPIN MINIQUICK PREFILLED SYRINGE 1.4 MG SUBCUTANEOUS
- GENOTROPIN MINIQUICK PREFILLED SYRINGE 1.6 MG SUBCUTANEOUS
- GENOTROPIN MINIQUICK PREFILLED SYRINGE 1.8 MG SUBCUTANEOUS
- GENOTROPIN MINIQUICK PREFILLED SYRINGE 2 MG SUBCUTANEOUS
- HUMATROPE CARTRIDGE 12 MG INJECTION
- HUMATROPE CARTRIDGE 24 MG INJECTION
- HUMATROPE CARTRIDGE 6 MG INJECTION
- NORDITROPIN FLEXPPO SOLUTION PEN-INJECTOR 10 MG/1.5ML SUBCUTANEOUS
- NORDITROPIN FLEXPPO SOLUTION PEN-INJECTOR 15 MG/1.5ML SUBCUTANEOUS
- NORDITROPIN FLEXPPO SOLUTION PEN-INJECTOR 30 MG/3ML SUBCUTANEOUS
- NORDITROPIN FLEXPPO SOLUTION PEN-INJECTOR 5 MG/1.5ML SUBCUTANEOUS
- NUTROPIN AQ NUSPIN 10 SOLUTION PEN-INJECTOR 10 MG/2ML SUBCUTANEOUS
- NUTROPIN AQ NUSPIN 20 SOLUTION PEN-INJECTOR 20 MG/2ML SUBCUTANEOUS
- NUTROPIN AQ NUSPIN 5 SOLUTION PEN-INJECTOR 5 MG/2ML SUBCUTANEOUS
- OMNITROPE SOLUTION CARTRIDGE 10 MG/1.5ML SUBCUTANEOUS
- OMNITROPE SOLUTION CARTRIDGE 5 MG/1.5ML SUBCUTANEOUS
- OMNITROPE SOLUTION RECONSTITUTED 5.8 MG SUBCUTANEOUS

Details

Criteria	<p>For Commercial and Essential plans, the patient must have tried and failed to have an adequate response to Genotropin.</p> <p>For Medicaid plans, the patient must have tried and failed to have an adequate response to Norditropin.</p>
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4/1/2025

Ilumya

Products Affected

- ILUMYA SOLUTION PREFILLED SYRINGE 100 MG/ML SUBCUTANEOUS

Details

Criteria	For approval, the patient must have tried and failed to have an adequate response to or had an intolerance to at least the formulary-preferred IL-17 inhibitor (Cosentyx) and one of the following: generic infliximab or Remicade or one other on-formulary biologic agent for the treatment of psoriasis (on-formulary adalimumab biosimilars, Enbrel, Humira, Skyrizi, on-formulary ustekinumab biosimilars).
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Imjudo

Products Affected

- IMJUDO SOLUTION 25 MG/1.25ML
INTRA VENOUS
- IMJUDO SOLUTION 300 MG/15ML
INTRA VENOUS

Details

Criteria	For approval for hepatocellular carcinoma, the patient must have tried and failed to have an adequate response to or had an intolerance or contraindication to Avastin or Avastin-containing regimens. For approval for non-small cell lung cancer, the patient must have tried and failed to have an adequate response to or had an intolerance or contraindication to regimens containing Keytruda or Tecentriq.
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Infliximab

Products Affected

- AVSOLA SOLUTION RECONSTITUTED 100 MG INTRAVENOUS
- INFLECTRA SOLUTION RECONSTITUTED 100 MG INTRAVENOUS
- RENFLEXIS SOLUTION RECONSTITUTED 100 MG INTRAVENOUS

Details

Criteria	For approval of Avsola, Inflectra, or Renflexis, the patient must have tried and failed to have an adequate response to generic infliximab or Remicade. This specific requirement applies to new starts only. This requirement does not apply to patients using Avsola, Inflectra, Renflexis, or another infliximab biosimilar agent for any indication not shared with generic infliximab or Remicade.
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Invega Hafyera

Products Affected

- INVEGA HAFYERA SUSPENSION PREFILLED SYRINGE 1092 MG/3.5ML INTRAMUSCULAR
- INVEGA HAFYERA SUSPENSION PREFILLED SYRINGE 1560 MG/5ML INTRAMUSCULAR

Details

Criteria	For approval, documentation of at least 4 months' treatment with 1-month paliperidone palmitate extended-release injectable suspension or at least one 3-month injection of 3-month paliperidone palmitate extended-release injectable suspension. This specific requirement applies to new starts only.
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4/1/2025

Invega Trinza

Products Affected

- INVEGA TRINZA SUSPENSION PREFILLED SYRINGE 273 MG/0.88ML INTRAMUSCULAR
- INVEGA TRINZA SUSPENSION PREFILLED SYRINGE 410 MG/1.32ML INTRAMUSCULAR
- INVEGA TRINZA SUSPENSION PREFILLED SYRINGE 546 MG/1.75ML INTRAMUSCULAR
- INVEGA TRINZA SUSPENSION PREFILLED SYRINGE 819 MG/2.63ML INTRAMUSCULAR

Details

Criteria	For approval, documentation of at least 4 months' treatment with 1-month paliperidone palmitate extended-release injectable suspension. This specific requirement applies to new starts only.
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Leucovorins

Products Affected

- KHAPZORY SOLUTION RECONSTITUTED 175 MG INTRAVENOUS
- *leucovorin calcium solution reconstituted 50 mg injection*

Details

Criteria	For approval, the patient must have tried and failed to have an adequate response to generic leucovorin. This specific requirement applies to new starts only and does not apply to patients using levoleucovorin for any indication not shared with leucovorin.
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Long-acting hemophilia factors

Products Affected

- *adynovate solution reconstituted 1000 unit intravenous*
- *adynovate solution reconstituted 1500 unit intravenous*
- *adynovate solution reconstituted 2000 unit intravenous*
- *adynovate solution reconstituted 250 unit intravenous*
- *adynovate solution reconstituted 3000 unit intravenous*
- *adynovate solution reconstituted 500 unit intravenous*
- *adynovate solution reconstituted 750 unit intravenous*
- ELOCTATE SOLUTION RECONSTITUTED 1000 UNIT INTRAVENOUS
- ELOCTATE SOLUTION RECONSTITUTED 2000 UNIT INTRAVENOUS
- UNIT INTRAVENOUS
- ELOCTATE SOLUTION RECONSTITUTED 250 UNIT INTRAVENOUS
- ELOCTATE SOLUTION RECONSTITUTED 4000 UNIT INTRAVENOUS
- ELOCTATE SOLUTION RECONSTITUTED 6000 UNIT INTRAVENOUS
- ESPEROCT SOLUTION RECONSTITUTED 1000 UNIT INTRAVENOUS
- ESPEROCT SOLUTION RECONSTITUTED 1500 UNIT INTRAVENOUS
- ESPEROCT SOLUTION RECONSTITUTED 2000 UNIT INTRAVENOUS
- ESPEROCT SOLUTION RECONSTITUTED 3000 UNIT INTRAVENOUS
- ESPEROCT SOLUTION RECONSTITUTED 500 UNIT INTRAVENOUS

Details

Criteria	For approval of Adynovate, Eloctate, or Esperoct, the patient must have tried and failed to have an adequate response to Jivi. This specific requirement applies to new starts only.
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Opdualag

Products Affected

- OPDUALAG SOLUTION 240-80 MG/20ML
INTRA VENOUS

Details

Criteria	For approval, the patient must have tried and failed to have an adequate response to Opdivo plus Yervoy or has a contraindication to the use of Yervoy. This specific requirement applies to new starts only.
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4/1/2025

Paclitaxel

Products Affected

- ABRAXANE SUSPENSION RECONSTITUTED 100 MG INTRAVENOUS
- *paclitaxel protein-bound part suspension reconstituted 100 mg intravenous*

Details

Criteria	For approval of Abraxane for any indication other than pancreatic cancer or small bowel carcinoma, the patient must have tried and failed to have an adequate response to generic paclitaxel. This specific requirement applies to new starts only and does not apply to patients using Abraxane for any indication not shared with generic paclitaxel.
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Pegfilgrastim

Products Affected

- FULPHILA SOLUTION PREFILLED SYRINGE 6 MG/0.6ML SUBCUTANEOUS
- FYLNETRA SOLUTION PREFILLED SYRINGE 6 MG/0.6ML SUBCUTANEOUS
- NEULASTA ONPRO PREFILLED SYRINGE KIT 6 MG/0.6ML SUBCUTANEOUS
- NEULASTA SOLUTION PREFILLED SYRINGE 6 MG/0.6ML SUBCUTANEOUS
- NYVEPRIA SOLUTION PREFILLED SYRINGE 6 MG/0.6ML SUBCUTANEOUS
- ROLVEDON SOLUTION PREFILLED SYRINGE 13.2 MG/0.6ML SUBCUTANEOUS
- STIMUFEND SOLUTION PREFILLED SYRINGE 6 MG/0.6ML SUBCUTANEOUS
- ZIEXTENZO SOLUTION PREFILLED SYRINGE 6 MG/0.6ML SUBCUTANEOUS

Details

Criteria	
	For approval of Fulphila, Fylnetra, Neulasta, Nyvepria, Rolvedon, Stimufend, or Ziextenzo, the patient must have tried and failed to have an adequate response to Udenyca. This specific requirement applies to new starts only and does not apply to patients using a non-preferred pegfilgrastim product for any indication not shared with a preferred pegfilgrastim product.

Pemetrexed

Products Affected

- AXTLER SOLUTION RECONSTITUTED 100 MG INTRAVENOUS
- AXTLER SOLUTION RECONSTITUTED 500 MG INTRAVENOUS
- PEMFEXY SOLUTION 500 MG/20ML
- INTRAVENOUS
- PEMRYDI RTU SOLUTION 100 MG/10ML INTRAVENOUS
- PEMRYDI RTU SOLUTION 500 MG/50ML INTRAVENOUS

Details

Criteria	For approval of Axtle (J9292), Pemfexy (J9304), or Pemrydi RTU (J9324), the patient must have tried and failed to have an adequate response to one of the following drugs: pemetrexed (J9294, J9296, J9297, J9305, J9314, J9322, J9323) unless contraindicated or not tolerated. This specific requirement applies to new starts only and does not apply to patients using Axtle, Pemfexy, or Pemrydi for an indication not shared with preferred agents.
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Ranibizumab

Products Affected

- BYOOVIZ SOLUTION 0.5 MG/0.05ML INTRAVITREAL
- LUCENTIS SOLUTION PREFILLED SYRINGE 0.3 MG/0.05ML INTRAVITREAL
- LUCENTIS SOLUTION PREFILLED SYRINGE 0.5 MG/0.05ML INTRAVITREAL
- SUSVIMO (IMPLANT 1ST FILL) SOLUTION 10 MG/0.1ML INTRAVITREAL
- SUSVIMO (IMPLANT REFILL) SOLUTION 10 MG/0.1ML INTRAVITREAL

Details

Criteria	For approval of Byooviz, Lucentis, or Susvimo for shared indications, the patient must have tried and failed to have an adequate response to both (Step 1) bevacizumab (Avastin) and (Step 2) either aflibercept (Eylea/Eylea HD), brolucizumab (Beovu), or faricimab (Vabysmo). This specific requirement applies to new starts only.
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Rebyota

Products Affected

- REBYOTA SUSPENSION 150 ML RECTAL

Details

Criteria	For approval, the patient must have had prior therapy with bezlotoxumab or has a contraindication to its use.
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4/1/2025

Rituximab

Products Affected

- RITUXAN HYCELA SOLUTION 1400-23400 MG -UT/11.7ML SUBCUTANEOUS INTRAVENOUS
- RITUXAN HYCELA SOLUTION 1600-26800 MG -UT/13.4ML SUBCUTANEOUS INTRAVENOUS
- RITUXAN SOLUTION 500 MG/50ML INTRAVENOUS
- TRUXIMA SOLUTION 100 MG/10ML INTRAVENOUS
- TRUXIMA SOLUTION 500 MG/50ML INTRAVENOUS

Details

Criteria	For approval of Rituxan for all indications except pemphigus vulgaris (PV), Rituxan Hycela, or Truxima, the patient must have tried and failed to have an adequate response to Riabni or Ruxience. This specific requirement applies to new starts only. This requirement does not apply to patients using a rituximab agent for any indication not shared by Riabni or Ruxience.
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Soliris

Products Affected

- SOLIRIS SOLUTION 300 MG/30ML
INTRA VENOUS

Details

Criteria	For a diagnosis of neuromyelitis optica spectrum disorder (NMOSD), the patient must have tried and failed to have an adequate response to or had an intolerance to or contraindication to Enspryng and either Uplizna or rituximab. For a diagnosis of generalized myasthenia gravis, the patient must have tried and failed to have an adequate response to or had an intolerance to or contraindication to Vyvgart. These requirements apply to new starts only.
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4/1/2025

Spinraza

Products Affected

- SPINRAZA SOLUTION 12 MG/5ML
INTRATHECAL

Details

Criteria	For approval, the patient must have tried and failed to have an adequate response to or had an intolerance to or contraindication to Evrysdi. This specific requirement applies to new starts only.
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4/1/2025

Tezspire

Products Affected

- TEZSPIRE SOLUTION PREFILLED SYRINGE 210 MG/1.91ML SUBCUTANEOUS

Details

Criteria	For approval, the patient must have (1) tried and failed, was intolerant to, or had a contraindication to dupilumab plus one other biologic for severe asthma, including either an IL-5 antagonist, an IL-5 receptor antagonist, or omalizumab or (2) an eosinophil count below that required to use these other medications. This specific requirement applies to new starts only.
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4/1/2025

Tocilizumab

Products Affected

- ACTEMRA SOLUTION 200 MG/10ML
INTRAVENOUS
- ACTEMRA SOLUTION 400 MG/20ML
INTRAVENOUS
- ACTEMRA SOLUTION 80 MG/4ML
INTRAVENOUS
- TOFIDENCE SOLUTION 200 MG/10ML
INTRAVENOUS
- TOFIDENCE SOLUTION 400 MG/20ML
INTRAVENOUS
- TOFIDENCE SOLUTION 80 MG/4ML
INTRAVENOUS

Details

Criteria	For approval of Actemra or Tofidence, the patient must have tried and failed to have an adequate response to Tyenne. This specific requirement applies to new starts only. This requirement does not apply to patients using Actemra, Tofidence, or another tocilizumab biosimilar agent for any indication not shared with Tyenne.
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Trastuzumab

Products Affected

- HERCEPTIN HYLECTA SOLUTION 600-10000 MG-UNT/5ML SUBCUTANEOUS
- HERCEPTIN SOLUTION RECONSTITUTED 150 MG INTRAVENOUS
- HERZUMA SOLUTION RECONSTITUTED 150 MG INTRAVENOUS
- HERZUMA SOLUTION RECONSTITUTED 420 MG INTRAVENOUS
- OGIVRI SOLUTION RECONSTITUTED 150 MG INTRAVENOUS
- OGIVRI SOLUTION RECONSTITUTED 420 MG INTRAVENOUS
- ONTRUZANT SOLUTION RECONSTITUTED 150 MG INTRAVENOUS
- ONTRUZANT SOLUTION RECONSTITUTED 420 MG INTRAVENOUS

Details

Criteria	For approval of Herceptin, Herceptin Hylecta, Herzuma, Ogivri, or Ontruzant, the patient must have tried and failed to have an adequate response to Kanjinti or Trazimera. This specific requirement applies to new starts only. This requirement does not apply to patients using a trastuzumab agent for any indication not shared by Kanjinti or Trazimera.
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Ultomiris

Products Affected

- ULTOMIRIS SOLUTION 1100 MG/11ML
INTRA VENOUS
- ULTOMIRIS SOLUTION 300 MG/3ML
INTRA VENOUS

Details

Criteria	For a diagnosis of generalized myasthenia gravis, the patient must have tried and failed to have an adequate response to or had an intolerance to or contraindication to Vyvgart. These requirements apply to new starts only.
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Vabysmo

Products Affected

- VABYSMO SOLUTION 6 MG/0.05ML
INTRAVITREAL

Details

Criteria	For approval for a diagnosis of neovascular (wet) age-related macular degeneration or diabetic macular edema, the patient must have tried and failed to have an adequate response to bevacizumab (Avastin). This specific requirement applies to new starts only.
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4/1/2025

Viscosupplements

Products Affected

- DUROLANE PREFILLED SYRINGE 60 MG/3ML INTRA-ARTICULAR
- GEL-ONE PREFILLED SYRINGE 30 MG/3ML INTRA-ARTICULAR
- GELSYN-3 SOLUTION PREFILLED SYRINGE 16.8 MG/2ML INTRA-ARTICULAR
- GENVISC 850 SOLUTION PREFILLED SYRINGE 25 MG/2.5ML INTRA-ARTICULAR
- HYALGAN SOLUTION 20 MG/2ML INTRA-ARTICULAR
- HYALGAN SOLUTION PREFILLED SYRINGE 20 MG/2ML INTRA-ARTICULAR
- HYMOVIS SOLUTION PREFILLED SYRINGE 24 MG/3ML INTRA-ARTICULAR
- MONOVISC SOLUTION PREFILLED SYRINGE 88 MG/4ML INTRA-ARTICULAR
- ORTHOVISC SOLUTION PREFILLED SYRINGE 30 MG/2ML INTRA-ARTICULAR
- SUPARTZ FX SOLUTION PREFILLED SYRINGE 25 MG/2.5ML INTRA-ARTICULAR
- SYNOJOYNT SOLUTION PREFILLED SYRINGE 20 MG/2ML INTRA-ARTICULAR
- SYNVISC SOLUTION PREFILLED SYRINGE 16 MG/2ML INTRA-ARTICULAR
- TRILURON SOLUTION PREFILLED SYRINGE 20 MG/2ML INTRA-ARTICULAR
- TRIVISC SOLUTION PREFILLED SYRINGE 25 MG/2.5ML INTRA-ARTICULAR

Details

Criteria	For approval of viscosupplements other than Euflexxa or Synvisc-One, the patient must have tried and failed to have an adequate response to or had an intolerance to both preferred agents (Euflexxa, Synvisc-One). All viscosupplements require medical prior authorization. Because all viscosupplements are considered medical devices and not drugs by the FDA, they can only be billed through Medicare Part B. This specific requirement applies to new starts only and does not apply to patients using a non-preferred viscosupplement for any indication not shared with Euflexxa or Synvisc-One.
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Vyepti

Products Affected

- VYEPTI SOLUTION 100 MG/ML
INTRA VENOUS

Details

Criteria	For approval of Vyepti, the patient must have tried and failed to have an adequate response to any two oral agents used to prevent/reduce frequency of migraines and have had prior use of one injectable CGRP antagonist/receptor antagonist. This specific requirement applies to new starts only.
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4/1/2025

Vyvgart

Products Affected

- VYVGART HYTRULO SOLUTION 180-2000 MG-UNIT/ML SUBCUTANEOUS
- VYVGART SOLUTION 400 MG/20ML INTRAVENOUS

Details

Criteria	For approval, the patient must have failed to respond to therapy with at least two of the following drug groups: acetylcholinesterase inhibitors, corticosteroids, nonsteroidal immunosuppressive therapies (e.g., methotrexate, azathioprine, mycophenolate, cyclosporine). This specific requirement applies to new starts only.
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4/1/2025

Zilretta

Products Affected

- ZILRETTA SUSPENSION RECONSTITUTED ER
32 MG INTRA-ARTICULAR

Details

Criteria	For approval, the patient must have tried and failed to have an adequate response to at least one other injectable corticosteroid. This specific requirement applies to new starts only.
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4/1/2025

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