Geisinger Medicare 2025 Part B Step Therapy List & Criteria

TABLE OF CONTENTS

AVASTIN	3
BEOVU	4
BYOOVIZ	5
CIMERLI	6
EYLEA	7
EYLEA HD	8
GEL-ONE	9-10
HERCEPTIN	11
HYMOVIS	12-13
INFLIXIMAB (REMICADE OR UNBRANDED INFLIXIMAB)	14-15
KHAPZORY	16
LUCENTIS	17
MONOVISC	
RENFLEXIS	
RITUXAN	
SUSVIMO	25-26
SYNOJOYNT	
TRILURON	
TRIVISC	
VABYSMO	
ZILRETTA	35

AVASTIN (BEVACIZUMAB)

Affected Drugs

AVASTIN INTRAVENOUS SOLUTION 100MG/4ML VIAL AVASTIN INTAVENOUS SOLUTION 400MG/16ML VIAL

Step Therapy Criteria

 Medical record documentation of a therapeutic failure of, intolerance to, or contraindication to all of the following: Alymsys (bevacizumab-maly), Mvasi (bevacizumab-awwb), Vegzelma (bevacizumab-adcd), Zirabev (bevacizumab-bvzr).

AUTHORIZATION DURATION:

For adjuvant treatment of Stage III or IV Epithelial Ovarian, Fallopian Tube or Primary Peritoneal Cancer following initial surgical resection:

Authorization will be for one (1) 21 month approval. Authorization of Avastin for adjuvant treatment should not exceed the FDA-approved treatment duration of 21 months (28 cycles). For requests exceeding the above limit, medical record documentation of the following is required:

 Peer-reviewed literature citing well-designed clinical trials to indicate that the member's healthcare outcome will be improved by dosing beyond the FDA-approved treatment duration

For all other indications: Authorization will be open-ended

Step 1 Drugs

Alymsys (bevacizumab-maly) Mvasi (bevacizumab-awwb) Vegzelma (bevacizumab-adcd) Zirabev (bevacizumab-bvzr)

BEOVU (BROLUCIZUMAB)

Affected Drugs

BEOVU INTRAVITREAL SOLUTION 6MG/0.05ML VIAL BEOVU INTRAVITREAL SOLUTION 6MG/0.05ML PREFILLED SYRINGE

Step Therapy Criteria

- Medical record documentation of a diagnosis of neovascular age-related macular degeneration AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to intravitreal bevacizumab (Avastin)

OR

- Medical record documentation of a diagnosis of diabetic macular edema AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to intravitreal bevacizumab (Avastin)

AUTHORIZATION DURATION: Approvals will be given for a lifetime duration.

QUANTITY LIMIT: 12mg per 25 days (6mg per eye per 25 days)

Step 1 Drugs

Intravitreal Avastin (bevacizumab)

BYOOVIZ (RANIBIZUMAB-NUNA)

Affected Drugs

BYOOVIZ INTRAVITREAL SOLUTION 0.5MG/0.05ML VIAL

Step Therapy Criteria

- Medical record documentation of a diagnosis of neovascular age-related macular degeneration AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to intravitreal bevacizumab (Avastin)

OR

- Medical record documentation of a diagnosis of diabetic retinopathy with or without macular edema AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to intravitreal bevacizumab (Avastin)

OR

- Medical record documentation of a diagnosis of macular edema following retinal vein occlusion OR myopic choroidal neovascularization AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to intravitreal bevacizumab (Avastin)

AUTHORIZATION DURATION: Approvals will be given for a lifetime duration.

QUANTITY LIMIT: 0.1mL (1mg) per 28 days (0.5mg per eye per 28 days)

Step 1 Drugs

Intravitreal Avastin (bevacizumab)

CIMERLI (RANIBIZUMAB-EQRN)

Affected Drugs

CIMERLI INTRAVITREAL SOLUTION 0.5MG/0.05ML VIAL CIMERLI INTRAVITREAL SOLUTION 0.3MG/0.05ML VIAL

Step Therapy Criteria

- Medical record documentation of a diagnosis of neovascular age-related macular degeneration AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to intravitreal bevacizumab (Avastin)

OR

- Medical record documentation of a diagnosis of diabetic retinopathy with or without macular edema AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to intravitreal bevacizumab (Avastin)

OR

- Medical record documentation of a diagnosis of macular edema following retinal vein occlusion OR myopic choroidal neovascularization AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to intravitreal bevacizumab (Avastin)

AUTHORIZATION DURATION: Approvals will be given for a lifetime duration.

QUANTITY LIMIT: 0.1mL (1mg) per 28 days (0.5mg per eye per 28 days)

Step 1 Drugs

Intravitreal Avastin (bevacizumab)

EYLEA (AFLIBERCEPT)

Affected Drugs

EYLEA INTRAVITREAL SOLUTION 2MG/0.05ML VIAL EYLEA INTRAVITREAL SOLUTION 2MG/0.05ML PREFILLED SYRINGE

Step Therapy Criteria

- Medical record documentation of a diagnosis of neovascular age-related macular degeneration AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to intravitreal bevacizumab (Avastin)

OR

- Medical record documentation of a diagnosis of diabetic retinopathy with or without macular edema AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to Avastin OR medical record documentation of baseline best-corrected visual acuity 20/50 or worse

OR

- Medical record documentation of a diagnosis of macular edema following retinal vein occlusion AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to intravitreal bevacizumab (Avastin)

OR

• Medical record documentation of a diagnosis of retinopathy of prematurity (ROP)

AUTHORIZATION DURATION: Approvals will be given for a lifetime duration.

QUANTITY LIMIT: 0.1mL (4mg) per 25 days (2mg per eye per 25 days)

Step 1 Drugs

Intravitreal Avastin (bevacizumab)

EYLEA HD (AFLIBERCEPT)

Affected Drugs

EYLEA HD INTRAVITREAL SOLUTION 8MG/0.07ML VIAL

Step Therapy Criteria

- Medical record documentation of a diagnosis of neovascular age-related macular degeneration AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to one (1) of the following: Eylea*, Beovu*, Lucentis*, Byooviz*, or Cimerli* AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to Vabysmo*

OR

- Medical record documentation of a diagnosis of diabetic retinopathy with or without macular edema AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to one (1) of the following: Eylea*, Beovu*, Lucentis*, Byooviz*, or Cimerli* AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to Vabysmo*

*Prior authorization required

AUTHORIZATION DURATION: Approvals will be given for a lifetime duration.

QUANTITY LIMIT: 0.14mL (16mg) per 21 days (8mg per eye per 21 days)

Step 1 Drugs

Neovascular age-related macular degeneration:

Intravitreal Eylea (aflibercept) Intravitreal Lucentis (ranibizumab) Intravitreal Beovu (brolucizumab) Intravitreal Byooviz (ranibizumab-nuna) Intravitreal Cimerli (ranibizumab-eqrn)

Diabetic Retinopathy/Diabetic Macular Edema:

Intravitreal Eylea (aflibercept) Intravitreal Lucentis (ranibizumab) Intravitreal Byooviz (ranibizumab-nuna) Intravitreal Cimerli (ranibizumab-eqrn)

Step 2 Drugs Neovascular age-related macular degeneration/Diabetic Retinopathy/Diabetic Macular Edema: Vabysmo (faricimab)

GEL-ONE (CROSS-LINKED HYALURONATE)

Affected Drugs

GEL-ONE INTRA-ARTICULAR 30MG/3ML PREFILLED SYRINGE

Step Therapy Criteria

- Physician documented symptomatic osteoarthritis of the knee, defined as knee pain associated with radiographic evidence of osteophytes in the knee joint provided the clinical presentation is not that of "bone-on-bone", morning stiffness of less than or equal to 30 minutes in duration, crepitus on range of motion; **AND**
- Physician documented knee joint pain sufficient to interfere with ambulatory functional activities; **AND**
- Physician documentation of non-pharmacologic modalities, e.g., weight loss, quadriceps muscle strengthening, other physical therapy modalities, or exercises that have not promoted satisfactory symptomatic relief; **AND**
- Physician documentation that there has been no significant improvement following pharmacologic therapy with a full-dose nonsteroidal anti-inflammatory drug (NSAID) regimen, with or without supplemental acetaminophen, over a 10-12 week period of time or if NSAID's are contraindicated, a failure of daily acetaminophen regimen over a 4 to 6 week period; AND
- Physician documentation that there has been no significant improvement following standard dose intra-articular corticosteroid injection(s) e.g., a satisfactory clinical response of greater than or equal to 3 months; this requirement does not apply if the use of corticosteroids might increase the risk of local or systemic bacterial infection, e.g., diabetes mellitus; **AND**
- Physician documentation of failure on, intolerance to or contraindication to two (2) of the following: Durolane, Euflexxa, Gelsyn-3, GenVisc, Hyalgan, Orthovisc, Supartz, Synvisc/Synvisc-One, and Visco-3

AUTHORIZATION DURATION: Initial approval will be for six (6) months. Subsequent approvals will be for six (6) months when members meet the following criteria:

- Medical record documentation of significant improvement in pain and function following the previous injection; AND
- Documented reduction of the doses of nonsteroidals or analgesics during the six-month period following the last injection in the previous series as well as no need for accompanying intra-articular steroid injections; **AND**
- Six months or longer have elapsed since the last injection in the previous series.

QUANTITY LIMIT: One (1) treatment course to the affected knee(s) (bilateral injections may be allowed if both knees meet the required coverage criteria)

LIMITATIONS:

• Gel-One treatment course is limited to 1 injection in a 6-month period.

CONTRAINDICATIONS:

- The use of these products for injection into any joint other than the knee.
- Injection of these products for indications other than the diagnosis of osteoarthritis
- Documented allergy to chickens or eggs.
- Knee joint infection, skin disease or infection around the area where the injection will be given.
- The insured individual has known sensitivity or contraindication to the use of either sodium hyaluronate or hylan G-F 20, e.g., crystal synovitis or hypersensitivity to hyaluronan preparations

Step 1 Drugs

Durolane, Euflexxa, Gelsyn-3, GenVisc, Hyalgan, Orthovisc, Supartz, Synvisc/Synvisc-One, Visco-3

HERCEPTIN (TRASTUZUMAB)

Affected Drugs

HERCEPTIN INTRAVENOUS SOLUTION 150MG VIAL

Step Therapy Criteria

 Medical record documentation of a therapeutic failure of, intolerance to, or contraindication to at least two (2) of the following: trastuzumab-anns (Kanjinti), trastuzumab-dkst (Ogivri), trastuzumab-dttb (Ontruzant), trastuzumab-qyyp (Trazimera), and/or trastuzumab-pkrb (Herzuma)

AUTHORIZATION DURATION:

For adjuvant treatment:

Authorization will be for one (1) 12-month approval. Authorization of Herceptin for adjuvant treatment should not exceed the FDA-approved treatment duration of 1 year (12 months). For requests exceeding the above limit, medical record documentation of the following is required:

 Peer-reviewed literature citing well-designed clinical trials to indicate that the member's healthcare outcome will be improved by dosing beyond the FDA-approved treatment duration

For all other indications: Authorization will be open-ended

Step 1 Drugs

Herzuma (trastuzumab-pkrb), Kanjinti (trastuzumab-anns), Ogivri (trastuzumab-dkst), Ontruzant (trastuzumab-dttb), Trazimera (trastuzumab-qyyp)

HYMOVIS (HIGH MOLECULAR WEIGHT VISCOELASTIC HYALURONAN)

Affected Drugs

HYMOVIS INTRA-ARTICULAR SOLUTION 24MG/3ML PREFILLED SYRINGE

Step Therapy Criteria

- Physician documented symptomatic osteoarthritis of the knee, defined as knee pain associated with radiographic evidence of osteophytes in the knee joint provided the clinical presentation is not that of "bone-on-bone", morning stiffness of less than or equal to 30 minutes in duration, crepitus on range of motion; **AND**
- Physician documented knee joint pain sufficient to interfere with ambulatory functional activities; **AND**
- Physician documentation of non-pharmacologic modalities, e.g., weight loss, quadriceps muscle strengthening, other physical therapy modalities, or exercises that have not promoted satisfactory symptomatic relief; **AND**
- Physician documentation that there has been no significant improvement following pharmacologic therapy with a full-dose nonsteroidal anti-inflammatory drug (NSAID) regimen, with or without supplemental acetaminophen, over a 10-12 week period of time or if NSAID's are contraindicated, a failure of daily acetaminophen regimen over a 4 to 6 week period; AND
- Physician documentation that there has been no significant improvement following standard dose intra-articular corticosteroid injection(s) e.g., a satisfactory clinical response of greater than or equal to 3 months; this requirement does not apply if the use of corticosteroids might increase the risk of local or systemic bacterial infection, e.g., diabetes mellitus; **AND**
- Physician documentation of failure on, intolerance to or contraindication to two (2) of the following: Durolane, Euflexxa, Gelsyn-3, GenVisc, Hyalgan, Orthovisc, Supartz, Synvisc/Synvisc-One, and Visco-3

AUTHORIZATION DURATION: Initial approval will be for six (6) months. Subsequent approvals will be for six (6) months when members meet the following criteria:

- Medical record documentation of significant improvement in pain and function following the previous injection; **AND**
- Documented reduction of the doses of nonsteroidals or analgesics during the six-month period following the last injection in the previous series as well as no need for accompanying intra-articular steroid injections; **AND**
- Six months or longer have elapsed since the last injection in the previous series.

QUANTITY LIMIT: One (1) treatment course to the affected knee(s) (bilateral injections may be allowed if both knees meet the required coverage criteria)

LIMITATIONS:

• Hymovis treatment course is limited to 2 injections in a 6-month period.

CONTRAINDICATIONS:

- The use of these products for injection into any joint other than the knee.
- Injection of these products for indications other than the diagnosis of osteoarthritis
- Documented allergy to chickens or eggs.
- Knee joint infection, skin disease or infection around the area where the injection will be given.
- The insured individual has known sensitivity or contraindication to the use of either sodium hyaluronate or hylan G-F 20, e.g., crystal synovitis or hypersensitivity to hyaluronan preparations

Step 1 Drugs

Durolane, Euflexxa, Gelsyn-3, GenVisc, Hyalgan, Orthovisc, Supartz, Synvisc/Synvisc-One, Visco-3

INFLIXIMAB (REMICADE OR UNBRANDED INFLIXIMAB)

Affected Drugs

REMICADE INTRAVENOUS SOLUTION RECONSTITUTED 100 MG VIAL INFLIXIMAB INTRAVENOUS SOLUTION RECONSTITUDED 100MG VIAL

Step Therapy Criteria

For Treatment of Rheumatoid Arthritis:

- Must be 18 years of age or greater AND
- Requesting provider must be a rheumatologist AND
- Diagnosis of moderate to severe rheumatoid arthritis according the American College of Rheumatology Criteria for the Classification and Diagnosis of Rheumatoid Arthritis AND
- Medical record documentation that the infliximab product is <u>not</u> being used concurrently with a TNF blocker or other biologic agent AND
- Continuation of effective dose of methotrexate during infliximab therapy AND
- Medical record documentation of a therapeutic failure on, intolerance to, or contraindication to two of the following: infliximab-dyyb (Inflectra) and infliximab-axxq (Avsola)

For Treatment of Crohn's Disease, Pediatric Crohn's Disease, and/or Fistulizing Crohn's Disease:

- Must be 6 years of age or older; AND
- Prescription is written by a gastroenterologist AND
- Medical record documentation of a diagnosis of moderate to severe Crohn's disease AND
- Medical record documentation that the infliximab product is <u>not</u> being used concurrently with a TNF blocker or other biologic agent AND
- Medical record documentation of a therapeutic failure on, intolerance to, or contraindication to two of the following: infliximab-dyyb (Inflectra) and infliximab-axxq (Avsola)

For Treatment of Ulcerative Colitis:

- Must be at least 6 years of age; AND
- Must be prescribed by a gastroenterologist; AND
- Physician provided documentation of a diagnosis of moderate to severe ulcerative colitis AND
- Medical record documentation that the infliximab product is <u>not</u> being used concurrently with a TNF blocker or other biologic agent **AND**
- Medical record documentation of a therapeutic failure on, intolerance to, or contraindication to two of the following: infliximab-dyyb (Inflectra) and infliximab-axxq (Avsola)

For Treatment of Ankylosing Spondylitis:

- Physician documentation of a diagnosis of ankylosing spondylitis AND
- Prescribing physician must be a rheumatologist AND
- Must be at least 18 years of age AND
- Medical record documentation that the infliximab product is <u>not</u> being used concurrently with a TNF blocker or other biologic agent AND

 Medical record documentation of a therapeutic failure on, intolerance to, or contraindication to two of the following: infliximab-dyyb (Inflectra) and infliximab-axxq (Avsola)

For the treatment of Plaque Psoriasis:

- Prescribed by a dermatologist AND
- Insured individual must be at least 18 years of age AND
- Physician provided documentation of a diagnosis of moderate to severe plaque psoriasis characterized by greater than or equal to 5% body surface area involved or disease affecting crucial body areas such as the hands, feet, face, or genitals **AND**
- Medical record documentation that the infliximab product is <u>not</u> being used concurrently with a TNF blocker or other biologic agent **AND**
- Medical record documentation of a therapeutic failure on, intolerance to, or contraindication to two of the following: infliximab-dyyb (Inflectra) and infliximab-axxq (Avsola)

For the treatment of Psoriatic Arthritis:

- Physician provided documentation of a diagnosis of moderately to severely active psoriatic arthritis which must include the following:
 - Documentation of either active psoriatic lesions or a documented history of psoriasis

AND

- Must be prescribed by a rheumatologist or dermatologist AND
- Must be at least 18 years of age AND
- Medical record documentation that the infliximab product is <u>not</u> being used concurrently with a TNF blocker or other biologic agent AND
- Medical record documentation of a therapeutic failure on, intolerance to, or contraindication to two of the following: infliximab-dyyb (Inflectra) and infliximab-axxq (Avsola)

AUTHORIZATION DURATION: Approval will be given for an initial duration of six (6) months. For continuation of coverage, medical record documentation of clinical improvement or lack of progression in the signs and symptoms of the treated indication at six (6) months of infliximab therapy is required.

After the initial six (6) month approval, subsequent approvals for coverage will be for a duration of one (1) year. Reevaluation of coverage will be every one (1) year requiring medical record documentation of continued or sustained improvement in the signs and symptoms of the treated indication while on infliximab therapy.

Step 1 Drugs

Avsola (infliximab-axxq), Inflectra (infliximab-dyyb)

KHAPZORY (LEVOLEUCOVORIN)

Affected Drugs

KHAPZORY SOLUTION RECONSTITUED 175 MG VIAL

Step Therapy Criteria

• Medical record documentation of intolerance to or contraindication to preferred levoleucovorin calcium products.

Preferred products: levoleucovorin calcium vial, powder for reconstitution, levoleucovorin calcium vial, solution

AUTHORIZATION DURATION: Initial approval will be for 12 months or less if the reviewing provider feels it is medically appropriate. Subsequent approvals will be for an additional 12 months or less if the reviewing provider feels it is medically appropriate and will require medical record documentation of continued disease improvement or lack of disease progression. The medication will no longer be covered if patient experiences toxicity or worsening of disease.

Step 1 Drugs

levoleucovorin calcium vial, powder for reconstitution; levoleucovorin calcium vial, solution

LUCENTIS (RANIBIZUMAB)

Affected Drugs

LUCENTIS INTRAVITREAL SOLUTION 0.5MG/0.05ML PREFILLED SYRINGE LUCENTIS INTRAVITREAL SOLUTION 0.3MG/0.05ML PREFILLED SYRINGE

Step Therapy Criteria

- Medical record documentation of a diagnosis of neovascular age-related macular degeneration AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to intravitreal bevacizumab (Avastin)

OR

- Medical record documentation of a diagnosis of diabetic retinopathy with or without macular edema AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to intravitreal bevacizumab (Avastin)

OR

- Medical record documentation of a diagnosis of macular edema following retinal vein occlusion OR myopic choroidal neovascularization AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to intravitreal bevacizumab (Avastin)

AUTHORIZATION DURATION: Approvals will be given for a lifetime duration.

QUANTITY LIMIT: 0.1mL (1mg) per 28 days (0.5mg per eye per 28 days)

Step 1 Drugs

Intravitreal Avastin (bevacizumab)

MONOVISC (HIGH MOLECULAR WEIGHT HYALURONON)

Affected Drugs

MONOVISC INTRA-ARTICULAR SOLUTION 88MG/4ML PREFILLED SYRINGE

Step Therapy Criteria

- Physician documented symptomatic osteoarthritis of the knee, defined as knee pain associated with radiographic evidence of osteophytes in the knee joint provided the clinical presentation is not that of "bone-on-bone", morning stiffness of less than or equal to 30 minutes in duration, crepitus on range of motion; **AND**
- Physician documented knee joint pain sufficient to interfere with ambulatory functional activities; **AND**
- Physician documentation of non-pharmacologic modalities, e.g., weight loss, quadriceps muscle strengthening, other physical therapy modalities, or exercises that have not promoted satisfactory symptomatic relief; **AND**
- Physician documentation that there has been no significant improvement following pharmacologic therapy with a full-dose nonsteroidal anti-inflammatory drug (NSAID) regimen, with or without supplemental acetaminophen, over a 10-12 week period of time or if NSAID's are contraindicated, a failure of daily acetaminophen regimen over a 4 to 6 week period; AND
- Physician documentation that there has been no significant improvement following standard dose intra-articular corticosteroid injection(s) e.g., a satisfactory clinical response of greater than or equal to 3 months; this requirement does not apply if the use of corticosteroids might increase the risk of local or systemic bacterial infection, e.g., diabetes mellitus; **AND**
- Physician documentation of failure on, intolerance to or contraindication to two (2) of the following: Durolane, Euflexxa, Gelsyn-3, GenVisc, Hyalgan, Orthovisc, Supartz, Synvisc/Synvisc-One, and Visco-3

AUTHORIZATION DURATION: Initial approval will be for six (6) months. Subsequent approvals will be for six (6) months when members meet the following criteria:

- Medical record documentation of significant improvement in pain and function following the previous injection; **AND**
- Documented reduction of the doses of nonsteroidals or analgesics during the six-month period following the last injection in the previous series as well as no need for accompanying intra-articular steroid injections; **AND**
- Six months or longer have elapsed since the last injection in the previous series.

QUANTITY LIMIT: One (1) treatment course to the affected knee(s) (bilateral injections may be allowed if both knees meet the required coverage criteria)

LIMITATIONS:

Monovisc treatment course is limited to 1 injection in a 6-month period.

CONTRAINDICATIONS:

- The use of these products for injection into any joint other than the knee.
- Injection of these products for indications other than the diagnosis of osteoarthritis
- Documented allergy to chickens or eggs.
- Knee joint infection, skin disease or infection around the area where the injection will be given.
- The insured individual has known sensitivity or contraindication to the use of either sodium hyaluronate or hylan G-F 20, e.g., crystal synovitis or hypersensitivity to hyaluronan preparations

Step 1 Drugs

Durolane, Euflexxa, Gelsyn-3, GenVisc, Hyalgan, Orthovisc, Supartz, Synvisc/Synvisc-One, Visco-3

RENFLEXIS (INFLIXIMAB-ABDA)

Affected Drugs

RENFLEXIS INTRAVENOUS SOLUTION RECONSTITUTED 100 MG VIAL

Step Therapy Criteria

For Treatment of Rheumatoid Arthritis:

- Must be 18 years of age or greater AND
- Requesting provider must be a rheumatologist AND
- Diagnosis of moderate to severe rheumatoid arthritis according the American College of Rheumatology Criteria for the Classification and Diagnosis of Rheumatoid Arthritis AND
- Medical record documentation that the infliximab product is <u>not</u> being used concurrently with a TNF blocker or other biologic agent AND
- Continuation of effective dose of methotrexate during infliximab therapy AND
- Medical record documentation of a therapeutic failure on, intolerance to, or contraindication to two of the following: infliximab-dyyb (Inflectra) and infliximab-axxq (Avsola)

For Treatment of Crohn's Disease, Pediatric Crohn's Disease, and/or Fistulizing Crohn's Disease:

- Must be 6 years of age or older; **AND**
- Prescription is written by a gastroenterologist AND
- Medical record documentation of a diagnosis of moderate to severe Crohn's disease AND
- Medical record documentation that the infliximab product is <u>not</u> being used concurrently with a TNF blocker or other biologic agent AND
- Medical record documentation of a therapeutic failure on, intolerance to, or contraindication to two of the following: infliximab-dyyb (Inflectra) and infliximab-axxq (Avsola)

For Treatment of Ulcerative Colitis:

- Must be at least 6 years of age; AND
- Must be prescribed by a gastroenterologist; AND
- Physician provided documentation of a diagnosis of moderate to severe ulcerative colitis AND
- Medical record documentation that the infliximab product is <u>not</u> being used concurrently with a TNF blocker or other biologic agent AND
- Medical record documentation of a therapeutic failure on, intolerance to, or contraindication to two of the following: infliximab-dyyb (Inflectra) and infliximab-axxq (Avsola)

For Treatment of Ankylosing Spondylitis:

- Physician documentation of a diagnosis of ankylosing spondylitis AND
- Prescribing physician must be a rheumatologist AND
- Must be at least 18 years of age AND
- Medical record documentation that the infliximab product is <u>not</u> being used concurrently with a TNF blocker or other biologic agent **AND**

 Medical record documentation of a therapeutic failure on, intolerance to, or contraindication to two of the following: infliximab-dyyb (Inflectra) and infliximab-axxq (Avsola)

For the treatment of Plaque Psoriasis:

- Prescribed by a dermatologist AND
- Insured individual must be at least 18 years of age AND
- Physician provided documentation of a diagnosis of moderate to severe plaque psoriasis characterized by greater than or equal to 5% body surface area involved or disease affecting crucial body areas such as the hands, feet, face, or genitals **AND**
- Medical record documentation that the infliximab product is <u>not</u> being used concurrently with a TNF blocker or other biologic agent **AND**
- Medical record documentation of a therapeutic failure on, intolerance to, or contraindication to two of the following: infliximab-dyyb (Inflectra) and infliximab-axxq (Avsola)

For the treatment of Psoriatic Arthritis:

- Physician provided documentation of a diagnosis of moderately to severely active psoriatic arthritis which must include the following:
 - Documentation of either active psoriatic lesions or a documented history of psoriasis

AND

- Must be prescribed by a rheumatologist or dermatologist AND
- Must be at least 18 years of age AND
- Medical record documentation that the infliximab product is <u>not</u> being used concurrently with a TNF blocker or other biologic agent AND
- Medical record documentation of a therapeutic failure on, intolerance to, or contraindication to two of the following: infliximab-dyyb (Inflectra) and infliximab-axxq (Avsola)

AUTHORIZATION DURATION: Approval will be given for an initial duration of six (6) months. For continuation of coverage, medical record documentation of clinical improvement or lack of progression in the signs and symptoms of the treated indication at six (6) months of infliximab therapy is required.

After the initial six (6) month approval, subsequent approvals for coverage will be for a duration of one (1) year. Reevaluation of coverage will be every one (1) year requiring medical record documentation of continued or sustained improvement in the signs and symptoms of the treated indication while on infliximab therapy.

Step 1 Drugs

Avsola (infliximab-axxq), Inflectra (infliximab-dyyb)

RITUXAN (RITUXIMAB)

Affected Drugs

RITUXAN INTRAVENOUS SOLUTION 100 MG/10 ML VIAL RITUXAN INTRAVENOUS SOLUTION 500 MG/50 ML VIAL

Step Therapy Criteria

For Rheumatoid Arthritis:

- Physician documentation of a diagnosis of moderate to severe rheumatoid arthritis in accordance with the American College of Rheumatology Criteria for the Classification and Diagnosis of Rheumatoid Arthritis; **AND**
- At least 18 years of age or older; AND
- Prescription written by a rheumatologist; AND
- Medical record documentation that an effective dose of methotrexate will be continued during rituximab therapy; AND
- Medical record documentation that Rituxan is not being used concurrently with a TNF blocker AND
- For rituximab reference product requests (i.e. Rituxan), medical record documentation of a therapeutic failure on, intolerance to, or contraindication to two (2) of the following: rituximab-pvvr (Ruxience), rituximab-arrx (Riabni), or rituximab-abbs (Truxima).

For Chronic Immunothrombocytopenia (ITP):

- Diagnosis of primary chronic ITP AND
- Platelet count of < 30,000/mm3 with active bleeding; or platelet count < 30,000/mm3 and a documented history of significant bleeding; or < 20,000/mm3 AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to corticosteroids and/or IVIG* (*prior authorization required) AND
- For rituximab reference product requests (i.e. Rituxan), medical record documentation of a therapeutic failure on, intolerance to, or contraindication to two (2) of the following: rituximab-pvvr (Ruxience), rituximab-arrx (Riabni), or rituximab-abbs (Truxima).

For Acute Lymphoblastic Leukemia, Hairy Cell Leukemia, and Chronic Lymphoid Leukemia:

- Medical record documentation of a diagnosis of Chronic Lymphocytic Leukemia (CLL) AND
- For rituximab reference product requests (i.e. Rituxan), medical record documentation of a therapeutic failure on, intolerance to, or contraindication to two (2) of the following: rituximab-pvvr (Ruxience), rituximab-arrx (Riabni), or rituximab-abbs (Truxima).

For Microscopic Polyarteritis Nodosa (PAN)

- Medical record documentation of a diagnosis of microscopic polyarteritis nodosa used in combination with glucocorticoids AND
- For rituximab reference product requests (i.e. Rituxan), medical record documentation of a therapeutic failure on, intolerance to, or contraindication to two (2) of the following: rituximab-pvvr (Ruxience), rituximab-arrx (Riabni), or rituximab-abbs (Truxima).

For Granulomatosis with Polyangiitis (GPA) (Wegener's Granulomatosis) and Microscopic Polyangiitis (MPA)

- Medical record documentation of a diagnosis of Granulomatosis with Polyangiitis (GPA) (Wegener's granulomatosis) or Microscopic Polyangiitis (MPA) used in combination with glucocorticoids
 - AND
- For rituximab reference product requests (i.e. Rituxan), medical record documentation of a therapeutic failure on, intolerance to, or contraindication to two (2) of the following: rituximab-pvvr (Ruxience), rituximab-arrx (Riabni), or rituximab-abbs (Truxima).

For Non-Hodgkin Lymphoma

- Medical record documentation of a diagnosis of Non-Hodgkin Lymphoma AND
- For rituximab reference product requests (i.e. Rituxan), medical record documentation of a therapeutic failure on, intolerance to, or contraindication to two (2) of the following: rituximab-pvvr (Ruxience), rituximab-arrx (Riabni), or rituximab-abbs (Truxima).

For Hodgkin Lymphoma

- Medical record documentation of a diagnosis of Hodgkin Lymphoma AND
- For rituximab reference product requests (i.e. Rituxan), medical record documentation of a therapeutic failure on, intolerance to, or contraindication to two (2) of the following: rituximab-pvvr (Ruxience), rituximab-arrx (Riabni), or rituximab-abbs (Truxima).

For Multiple Sclerosis (MS)

- Medical record documentation of a diagnosis of Multiple Sclerosis AND
- For rituximab reference product requests (i.e. Rituxan), medical record documentation of a therapeutic failure on, intolerance to, or contraindication to two (2) of the following: rituximab-pvvr (Ruxience), rituximab-arrx (Riabni), or rituximab-abbs (Truxima).

For Refractory Chronic Debilitating Myasthenia Gravis

- Medical record documentation of refractory Chronic Debilitating Myasthenia Gravis AND
- Prescribed by or in consultation with a neuromuscular specialist AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to at least one corticosteroid **AND**
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to at least one cholinesterase inhibitor AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to at least one non-steroidal immunosuppressive therapy AND
- For rituximab reference product requests (i.e. Rituxan), medical record documentation of a therapeutic failure on, intolerance to, or contraindication to two (2) of the following: rituximab-pvvr (Ruxience), rituximab-arrx (Riabni), or rituximab-abbs (Truxima).

For Pemphigus Vulgaris (PV)

- Prescription written by a dermatologist AND
- Member is 18 years of age or older AND
- Medical record documentation of a diagnosis of moderate to severe pemphigus vulgaris AND

- Medical record documentation of use in combination with corticosteroids or a contraindication or intolerance to corticosteroids AND
- For rituximab reference product requests (i.e. Rituxan), medical record documentation of a therapeutic failure on, intolerance to, or contraindication to two (2) of the following: rituximab-pvvr (Ruxience), rituximab-arrx (Riabni), or rituximab-abbs (Truxima).

AUTHORIZATION DURATION:

<u>For Multiple Sclerosis:</u> Initial approval will be for 12 months or less if the reviewing provider feels it is medically appropriate. Subsequent approvals will be for an additional 12 months or less if the reviewing provider feels it is medically appropriate and will require medical record documentation of continued disease improvement or lack of disease progression. The medication will no longer be covered if patient experiences toxicity or worsening of disease.

For all other indications: Initial approval will be for 6 months or less if the reviewing provider feels it is medically appropriate. Subsequent approvals will be for an additional 12 months or less if the reviewing provider feels it is medically appropriate and will require medical record documentation of continued disease improvement or lack of disease progression. The medication will no longer be covered if patient experiences toxicity or worsening of disease.

Step 1 Drugs

Ruxience (rituximab-pvvr) Riabni (rituximab-arrx) Truxima (rituximab-abbs)

SUSVIMO (RANIBIZUMAB)

Affected Drugs

SUSVIMO (IMPLANT 1ST FILL) INTRAVITREAL SOLUTION 10MG/0.1ML VIAL SUSVIMO OCULAR IMPLANT INTRAVITREAL IMPLANT SUSVIMO (IMPLANT REFILL) INTRAVITREAL SOLUTION 10MG/0.1ML VIAL

Step Therapy Criteria

- Medical record documentation of a diagnosis of neovascular age-related macular degeneration AND
- Medical record documentation patient has previously responded to at least two (2) intravitreal doses of a Vascular Endothelial Growth Factor (VEGF) inhibitor medication AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to intravitreal bevacizumab (Avastin) AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to one (1) additional intravitreal VEGF inhibitor (e.g. Eylea, Beovu, or Lucentis)

AND

- Medical record documentation that Susvimo (ranibizumab) will not be given in combination with an intravitreal Vascular Endothelial Growth Factor (VGEF) inhibitor OR
- If the request is for use in combination with an intravitreal VEGF inhibitor, <u>all</u> of the following must be met:
 - Medical record documentation Susvimo (ranibizumab) will be given in combination with intravitreal ranibizumab injection (Lucentis) AND
 - Medical record documentation intravitreal ranibizumab injection will be administered on an as needed basis, as determined by the prescriber

AUTHORIZATION DURATION: Approval will be given for an **initial duration of two years (2) years** or less if the reviewing provider feels it is medically appropriate. After the initial two (2) year approval, subsequent approvals will be for a **lifetime duration** or less if the reviewing provider feels it is medically appropriate, and will require:

Medical record documentation that Susvimo (ranibizumab) will not be given in combination with an intravitreal Vascular Endothelial Growth (VGEF) inhibitor

QUANTITY LIMIT: 0.2mL (2 vials) per 24 weeks (to allow 2mg per 24 weeks per treated eye)

LIMITATION: Susvimo (ranibizumab) to be given in combination with intravitreal ranibizumab (Lucentis) injections after 92 weeks from the start of Susvimo therapy has not been studied in clinical trials and will require prior authorization.

Step 1 Drugs

Intravitreal Avastin (bevacizumab)

Step 2 Drugs

Intravitreal Beovu (brolucizumab) Intravitreal Byooviz (ranibizumab-nuna) Intravitreal Cimerli (ranibizumab-eqrn) Intravitreal Eylea (aflibercept) Intravitreal Lucentis (ranibizumab)

SYNOJOYNT (SODIUM HYALURONATE)

Affected Drugs

SYNOJOYNT INTRA-ARTICULAR SOLUTION 20MG/2ML PREFILLED SYRINGE

Step Therapy Criteria

- Physician documented symptomatic osteoarthritis of the knee, defined as knee pain associated with radiographic evidence of osteophytes in the knee joint provided the clinical presentation is not that of "bone-on-bone", morning stiffness of less than or equal to 30 minutes in duration, crepitus on range of motion; **AND**
- Physician documented knee joint pain sufficient to interfere with ambulatory functional activities; **AND**
- Physician documentation of non-pharmacologic modalities, e.g., weight loss, quadriceps muscle strengthening, other physical therapy modalities, or exercises that have not promoted satisfactory symptomatic relief; **AND**
- Physician documentation that there has been no significant improvement following pharmacologic therapy with a full-dose nonsteroidal anti-inflammatory drug (NSAID) regimen, with or without supplemental acetaminophen, over a 10-12 week period of time or if NSAID's are contraindicated, a failure of daily acetaminophen regimen over a 4 to 6 week period; AND
- Physician documentation that there has been no significant improvement following standard dose intra-articular corticosteroid injection(s) e.g., a satisfactory clinical response of greater than or equal to 3 months; this requirement does not apply if the use of corticosteroids might increase the risk of local or systemic bacterial infection, e.g., diabetes mellitus; **AND**
- Physician documentation of failure on, intolerance to or contraindication to two (2) of the following: Durolane, Euflexxa, Gelsyn-3, GenVisc, Hyalgan, Orthovisc, Supartz, Synvisc/Synvisc-One, and Visco-3

AUTHORIZATION DURATION: Initial approval will be for six (6) months. Subsequent approvals will be for six (6) months when members meet the following criteria:

- Medical record documentation of significant improvement in pain and function following the previous injection; **AND**
- Documented reduction of the doses of nonsteroidals or analgesics during the six-month period following the last injection in the previous series as well as no need for accompanying intra-articular steroid injections; **AND**
- Six months or longer have elapsed since the last injection in the previous series.

QUANTITY LIMIT: One (1) treatment course to the affected knee(s) (bilateral injections may be allowed if both knees meet the required coverage criteria)

LIMITATIONS:

Synojoynt treatment course is limited to 3 injections in a 6-month period.

CONTRAINDICATIONS:

- The use of these products for injection into any joint other than the knee.
- Injection of these products for indications other than the diagnosis of osteoarthritis
- Documented allergy to chickens or eggs.
- Knee joint infection, skin disease or infection around the area where the injection will be given.
- The insured individual has known sensitivity or contraindication to the use of either sodium hyaluronate or hylan G-F 20, e.g., crystal synovitis or hypersensitivity to hyaluronan preparations

Step 1 Drugs

Durolane, Euflexxa, Gelsyn-3, GenVisc, Hyalgan, Orthovisc, Supartz, Synvisc/Synvisc-One, Visco-3

TRILURON (SODIUM HYALURONATE)

Affected Drugs

TRILURON INTRA-ARTICULAR SOLUTION 20MG/2ML PREFILLED SYRINGE

Step Therapy Criteria

- Physician documented symptomatic osteoarthritis of the knee, defined as knee pain associated with radiographic evidence of osteophytes in the knee joint provided the clinical presentation is not that of "bone-on-bone", morning stiffness of less than or equal to 30 minutes in duration, crepitus on range of motion; **AND**
- Physician documented knee joint pain sufficient to interfere with ambulatory functional activities; AND
- Physician documentation of non-pharmacologic modalities, e.g., weight loss, quadriceps muscle strengthening, other physical therapy modalities, or exercises that have not promoted satisfactory symptomatic relief; **AND**
- Physician documentation that there has been no significant improvement following pharmacologic therapy with a full-dose nonsteroidal anti-inflammatory drug (NSAID) regimen, with or without supplemental acetaminophen, over a 10-12 week period of time or if NSAID's are contraindicated, a failure of daily acetaminophen regimen over a 4 to 6 week period; AND
- Physician documentation that there has been no significant improvement following standard dose intra-articular corticosteroid injection(s) e.g., a satisfactory clinical response of greater than or equal to 3 months; this requirement does not apply if the use of corticosteroids might increase the risk of local or systemic bacterial infection, e.g., diabetes mellitus; **AND**
- Physician documentation of failure on, intolerance to or contraindication to two (2) of the following: Durolane, Euflexxa, Gelsyn-3, GenVisc, Hyalgan, Orthovisc, Supartz, Synvisc/Synvisc-One, and Visco-3

AUTHORIZATION DURATION: Initial approval will be for six (6) months. Subsequent approvals will be for six (6) months when members meet the following criteria:

- Medical record documentation of significant improvement in pain and function following the previous injection; **AND**
- Documented reduction of the doses of nonsteroidals or analgesics during the six-month period following the last injection in the previous series as well as no need for accompanying intra-articular steroid injections; **AND**
- Six months or longer have elapsed since the last injection in the previous series.

QUANTITY LIMIT: One (1) treatment course to the affected knee(s) (bilateral injections may be allowed if both knees meet the required coverage criteria)

LIMITATIONS:

Triluron treatment course is limited to 3 injections in a 6-month period.

CONTRAINDICATIONS:

- The use of these products for injection into any joint other than the knee.
- Injection of these products for indications other than the diagnosis of osteoarthritis
- Documented allergy to chickens or eggs.
- Knee joint infection, skin disease or infection around the area where the injection will be given.
- The insured individual has known sensitivity or contraindication to the use of either sodium hyaluronate or hylan G-F 20, e.g., crystal synovitis or hypersensitivity to hyaluronan preparations

Step 1 Drugs

Durolane, Euflexxa, Gelsyn-3, GenVisc, Hyalgan, Orthovisc, Supartz, Synvisc/Synvisc-One, Visco-3

TRIVISC (SODIUM HYALURONATE)

Affected Drugs

TRIVISC INTRA-ARTICULAR SOLUTION 25MG/2.5ML PREFILLED SYRINGE

Step Therapy Criteria

- Physician documented symptomatic osteoarthritis of the knee, defined as knee pain associated with radiographic evidence of osteophytes in the knee joint provided the clinical presentation is not that of "bone-on-bone", morning stiffness of less than or equal to 30 minutes in duration, crepitus on range of motion; **AND**
- Physician documented knee joint pain sufficient to interfere with ambulatory functional activities; **AND**
- Physician documentation of non-pharmacologic modalities, e.g., weight loss, quadriceps muscle strengthening, other physical therapy modalities, or exercises that have not promoted satisfactory symptomatic relief; **AND**
- Physician documentation that there has been no significant improvement following pharmacologic therapy with a full-dose nonsteroidal anti-inflammatory drug (NSAID) regimen, with or without supplemental acetaminophen, over a 10-12 week period of time or if NSAID's are contraindicated, a failure of daily acetaminophen regimen over a 4 to 6 week period; AND
- Physician documentation that there has been no significant improvement following standard dose intra-articular corticosteroid injection(s) e.g., a satisfactory clinical response of greater than or equal to 3 months; this requirement does not apply if the use of corticosteroids might increase the risk of local or systemic bacterial infection, e.g., diabetes mellitus; **AND**
- Physician documentation of failure on, intolerance to or contraindication to two (2) of the following: Durolane, Euflexxa, Gelsyn-3, GenVisc, Hyalgan, Orthovisc, Supartz, Synvisc/Synvisc-One, and Visco-3

AUTHORIZATION DURATION: Initial approval will be for six (6) months. Subsequent approvals will be for six (6) months when members meet the following criteria:

- Medical record documentation of significant improvement in pain and function following the previous injection; **AND**
- Documented reduction of the doses of nonsteroidals or analgesics during the six-month period following the last injection in the previous series as well as no need for accompanying intra-articular steroid injections; **AND**
- Six months or longer have elapsed since the last injection in the previous series.

QUANTITY LIMIT: One (1) treatment course to the affected knee(s) (bilateral injections may be allowed if both knees meet the required coverage criteria)

LIMITATIONS:

TriVisc treatment course is limited to 3 injections in a 6-month period..

CONTRAINDICATIONS:

- The use of these products for injection into any joint other than the knee.
- Injection of these products for indications other than the diagnosis of osteoarthritis
- Documented allergy to chickens or eggs.
- Knee joint infection, skin disease or infection around the area where the injection will be given.
- The insured individual has known sensitivity or contraindication to the use of either sodium hyaluronate or hylan G-F 20, e.g., crystal synovitis or hypersensitivity to hyaluronan preparations

Step 1 Drugs

Durolane, Euflexxa, Gelsyn-3, GenVisc, Hyalgan, Orthovisc, Supartz, Synvisc/Synvisc-One, Visco-3

VABYSMO (FARICIMAB)

Affected Drugs

VABYSMO INTRAVITREAL SOLUTION 6MG/0.05ML VIAL VABYSMO INTRAVITREAL SOLUTION 6MG/0.05ML PREFILLED SYRINGE

Step Therapy Criteria

- Medical record documentation of a diagnosis of neovascular age-related macular degeneration AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to intravitreal bevacizumab (Avastin) AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to one (1) additional intravitreal VEGF inhibitor (e.g. Eylea, Beovu, or Lucentis)

OR

- Medical record documentation of a diagnosis of diabetic macular edema AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to intravitreal bevacizumab (Avastin) **AND**
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to one (1) additional intravitreal VEGF inhibitor (e.g. Eylea or Lucentis)

OR

- Medical record documentation of a diagnosis of macular edema following retinal vein occlusion AND
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to intravitreal bevacizumab (Avastin) **AND**
- Medical record documentation of therapeutic failure on, intolerance to, or contraindication to one (1) additional intravitreal VEGF inhibitor (e.g. Eylea or Lucentis)

AUTHORIZATION DURATION: Approvals will be given for a lifetime duration.

QUANTITY LIMIT: 0.1mL (12mg) per 21 days (6mg per eye per 21 days)

Step 1 Drugs Neovascular age-related macular degeneration, Diabetic Macular Edema & Retinal Vein Occlusion:

Intravitreal Avastin (bevacizumab)

Step 2 Drugs

Neovascular age-related macular degeneration:

Intravitreal Beovu (brolucizumab)

Intravitreal Byooviz (ranibizumab-nuna)

Intravitreal Cimerli (ranibizumab-eqrn)

Intravitreal Eylea (aflibercept)

Intravitreal Lucentis (ranibizumab)

Diabetic Macular Edema:

Intravitreal Byooviz (ranibizumab-nuna) Intravitreal Cimerli (ranibizumab-eqrn) Intravitreal Eylea (aflibercept) Intravitreal Lucentis (ranibizumab)

Retinal Vein Occlusion:

Intravitreal Eylea (aflibercept) Intravitreal Lucentis (ranibizumab)

ZILRETTA (TRIAMCINOLONE ACETONIDE ER INJECTION)

Affected Drugs

ZILRETTA INTRA-ARTICULAR INJECTION ER SUSPENSION 32MG

Step Therapy Criteria

- Prescribed by a rheumatologist or orthopedic specialist AND
- Patient is 18 years of age or older AND
- Medical record documentation of a diagnosis of osteoarthritic pain of the knee AND
- Medical record documentation that patient has not received a previous administration of Zilretta to the requested knee AND
- Medical record documentation that non-pharmacologic modalities (e.g. Weight loss, aerobic/resistance land-based exercise or aquatic exercise, other physical therapy modalities or exercises) have not promoted satisfactory symptomatic relief **AND**
- Medical record documentation that there has been no significant improvement following a 10-12 week trial of full-dose nonsteroidal anti-inflammatory drug (NSAID) therapy, with or without supplemental acetaminophen OR if NSAIDs are contraindicated, a failure of daily acetaminophen regimen over a 4 to 6 week period AND
- Medical record documentation of a therapeutic failure on or intolerance to two different intra-articular steroid injections (e.g. triamcinolone, methylprednisolone, betamethasone, dexamethasone).

AUTHORIZATION DURATION: One injection per knee per lifetime (Facets RX count 32 per knee per lifetime)

NOTES:

The safety and efficacy of repeat administrations of Zilretta have not been studied. The safety and efficacy of Zilretta for management of osteoarthritis pain in joints other than the knee have not been studied.

Zilretta is for intra-articular use only and should not be administered by epidural, intrathecal, intravenous, intraocular, intramuscular, intradermal, or subcutaneous routes.

Step 1 Drugs

Injectable triamcinolone, methylprednisolone, betamethasone, dexamethasone

Devised: 9/15/20

Revised: 6/18/21 (Renflexis, Inflectra, Remicade), 1/18/22 (Rituxan), 3/15/22 (Susvimo, Vabysmo, Beovu), 5/17/22 (VEGf QL), 7/19/22 (VEGf Best-Corrected VA), 9/13/22 (RTX ITP duration/alts), 10/25/22 (VEGf QL, Beovu DME indication), 12/21/22 (Beovu affected drugs, clarified intravitreal bevacizumab (Avastin), Inflectra/Remicade delete, Renflexis alternatives edit), 1/17/23 (Byooviz, Cimerli), 6/23/23 (Eylea ROP), 7/18/23 (Herceptin), 8/25/23 (Gel-One, Hymovis, Monovisc, Synojoynt, Triluron, TriVisc), 11/21/23 (added infliximab/Remicade, Renflexis alternatives edit, Susvimo/Vabysmo step drugs), 4/19/24 (added Avastin [per 1/16/24 P&T], RTX indications), 7/16/24 (Vabysmo RVO, Eylea alt for RVO, Lucentis/Byooviz/Cimerli alt for RVO & MCN), 11/29/24 (added Khapzory [per 1/16/24 P&T], added Zilretta, Rituxan Hodgkin lymphoma alts)

Reviewed:

MA UM Committee approval: 12/31/23, 5/22/24, 8/30/24