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| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | |
| Pulmonary Arterial Hypertension (PAH) Agents | |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | |
| Pulmozyme | |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | |
| Pyrukynd (mitapivat) | |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | |
| Radicava (edaravone) | |

| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1096 |
|---|------|
| Ranolazine products | 1099 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1099 |
| Ravicti | 1102 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1102 |
| Rayos | 1105 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1105 |
| Reblozyl (luspatercept-aamt) | 1108 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1108 |
| Recorlev (levoketoconazole) | 1112 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1112 |
| Rectiv | 1115 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1115 |
| Regranex | |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1117 |
| Relistor | 1119 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1119 |
| Repatha | 1123 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1123 |
| Retinal Vascular Disease Agents | 1132 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1132 |
| Revcovi - AZ | 1138 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1138 |
| Revlimid (lenalidomide) | 1141 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1141 |
| Reyvow - Arizona | 1152 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1152 |
| Rezurock (belumosudil) | 1156 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1156 |
| Rhofade | 1159 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1159 |
| Rinvoq (upadacitinib) | 1161 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1161 |

| Ruconest-Arizona | 1175 |
|---|------|
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1175 |
| Samsca | 1178 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1178 |
| Sandostatin | 1180 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1180 |
| Sedative Hypnotics - AZM | 1189 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1189 |
| Serevent Diskus - Arizona | 1192 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1192 |
| SGLT-2 Inhibitors - AZM | 1195 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1195 |
| Short-Acting Opioid Products - AZM | 1198 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1198 |
| Signifor | 1210 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1210 |
| Siliq- Arizona | 1212 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1212 |
| Simponi- Arizona | 1216 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1216 |
| Sivextro | 1227 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | |
| Skyrizi (risankizumab-rzaa) | 1231 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1231 |
| Soliris- AZ | 1238 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1238 |
| Somatuline Depot (lanreotide) | 1249 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1249 |
| Somavert | 1254 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1254 |
| Soriatane | 1257 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1257 |
| Spinraza- Arizona | 1260 |

| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | |
|--|------|
| Spiriva Respimat | 1267 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1267 |
| Spravato - AZ | 1268 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1268 |
| Sprycel (dasatinib) | 1274 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1274 |
| Stelara (ustekinumab) | |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | |
| Strensiq | 1291 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1291 |
| Sublingual Immunotherapy (SLIT) | 1296 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1296 |
| Sublocade - Arizona | 1304 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1304 |
| Suboxone - AZM | 1308 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1308 |
| Sucraid | 1312 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1312 |
| Sunosi | 1315 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1315 |
| Sutent | 1320 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1320 |
| Symdeko | 1328 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1328 |
| Symlin | 1331 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1331 |
| Synagis | 1333 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1333 |
| Systane, Refresh, Gonak, Genteal, Tears Naturale | 1347 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1347 |
| Takhzyro | 1349 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1349 |

| Talicia and Mycobutin | |
|---|--|
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | |
| Taltz - Arizona | |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | |
| Tarceva | |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | |
| Targretin | |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | |
| Tarpeyo (budesonide) | |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | |
| Tasigna | |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | |
| Tavalisse - ARIZONA | |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | |
| Tegsedi | |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | |
| Temodar (temozolomide) | |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | |
| Test Strips | |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | |
| Testosterone - AZM | |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | |
| Tezspire (tezepelumab-ekko) | |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | |
| Thalomid | |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | |
| Tobramycin Inhalation - ARIZONA | |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | |
| Topical NSAIDs | |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | |
| Toujeo Solostar, Toujeo Max Solostar, Semglee, Basaglar | |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | |
| Trelegy Ellipta - ARIZONA | |

| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1440 |
|---|------|
| Tremfya - AZ | 1442 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1442 |
| Tretinoin Capsules - ARIZONA | 1449 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1449 |
| Tretinoin Topical | 1452 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1452 |
| Trikafta | 1454 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1454 |
| Triptans - AZ | 1457 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1457 |
| Twyneo (tretinoin-benzoyl peroxide 0.1-3% cream) | 1467 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1467 |
| Tykerb | 1469 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1469 |
| Tymlos - Arizona | 1475 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1475 |
| Uloric | 1479 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1479 |
| Ultomiris (ravulizumab-cwvz) | 1481 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1481 |
| Valchlor | 1486 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1486 |
| Valsartan oral solution | 1489 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1489 |
| Vancomycin - AZ | 1491 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1491 |
| Vecamyl | 1494 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1494 |
| Velphoro (sucroferric oxyhydroxide), Auryxia (ferric citrate) | 1496 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1496 |
| Veltassa | 1498 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | |

| Vemlidy | 1501 |
|--|------|
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1501 |
| Ventolin, Proventil, generic albuterol | 1503 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1503 |
| Verkazia (cyclosporine ophthalmic emulsion 0.1%) | 1505 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1505 |
| Vijoice (alpelisib) | 1508 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1508 |
| Vitamin B-12 | 1511 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1511 |
| Vitamin C | 1513 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1513 |
| Vitamin D | 1515 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1515 |
| Vivjoa (oteseconazole) | 1517 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1517 |
| Vonjo (pacritinib) | 1519 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1519 |
| Voquezna Triple Pak (vonoprazan, amoxicillin, clarithromycin), Voquezna Dual Pak (von amoxicillin) | • |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | |
| Votrient | |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | |
| Voxzogo (vosoritide) | |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | |
| Vtama (tapinarof) | |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | |
| Vyndagel and Vyndamax | |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | |
| Vyvgart (efgartigimod alfa-fcab) | |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | |
| Wakix | |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | |

| Xalkori | 1548 |
|---|------|
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1548 |
| Xeljanz, Xeljanz XR (tofacitinib) | 1553 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1553 |
| Xenazine | 1563 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1563 |
| Xenleta | 1567 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1567 |
| Xermelo | 1570 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1570 |
| Xofluza (baloxavir) | 1572 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1572 |
| Xolair | 1574 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1574 |
| Xopenex Respules | 1581 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1581 |
| Xuriden | 1583 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1583 |
| Xyrem, Xywav | |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1585 |
| Yonsa | 1590 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1590 |
| Zelboraf | 1594 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1594 |
| Zeposia (ozanimod) | |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | |
| Zimhi (naloxone) | 1608 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | |
| Zolgensma (onasemnogene abeparvovec-xioi) | |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | |
| Zolinza | |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | |
| Zontivity | |

| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1617 |
|---|------|
| Zortress - ARIZONA | 1619 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | 1619 |
| Ztalmy (ganaxolone) | 1621 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1621 |
| Zytiga | 1624 |
| Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) | 1624 |
| Ζγνοχ | 1628 |
| Medicaid - Arizona (AZM, AZMREF, AZMDDD) | |

Acetaminophen (Dose > 4 gm)

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99555 Acetaminophen (Dose > 4 gm)

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Acetaminophen (Dose > 4gm) | |
|--|----------------|
| Guideline Type | Administrative |
| | |
| Approval Criteria | |
| A Desurants for existencia on the descence are start then 1000mm new day, should be deviad | |

1 - Requests for acetaminophen dosages greater than 4000mg per day should be denied. The total dose of acetaminophen (cumulative total daily dose of 4000mg) is not supported by the Food and Drug Administration (FDA).

2. Revision History

| Date | Notes |
|-----------|--------------------|
| 6/28/2021 | 7/1 Implementation |

Actemra - AZM

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-105269 Actemra - AZM

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 4/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Actemra Actpen, Actemra SQ | |
|--|-----------------------|
| Diagnosis | Rheumatoid Arthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting ALL of the following:

1.1 Diagnosis of moderately to severely active Rheumatoid Arthritis (RA)

AND

1.2 History of failure to a 3 month trial of ONE non-biologic disease modifying anti-rheumatic drug (DMARD) [e.g., methotrexate, leflunomide, sulfasalazine, hydroxychloroquine] at maximally indicated doses within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*

AND

1.3 Patient is not receiving Actemra in combination with ANY of the following:

- Biologic DMARD [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

1.4 History of failure, contraindication, or intolerance to BOTH of the following:

- Humira (adalimumab)**
- Enbrel (etanercept)**

AND

1.5 Prescribed by, or in consultation with, a rheumatologist

OR

2 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting ALL of the following:

2.1 Patient is currently on Actemra therapy as documented by claims history or medical records (document drug, date, and duration of therapy)

AND

2.2 Diagnosis of moderately to severely active RA

AND

2.3 Patient is not receiving Actemra in combination with ANY of the following:

- Biologic DMARD [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

2.4 Prescribed by, or in consultation with, a rheumatologist

| Notes | *Claims history may be used in conjunction as documentation of drug, |
|-------|--|
| | date, and duration of trial. **Drug may require PA |

| Product Name: Actemra Actpen, Actemra SQ | |
|--|--|
| Diagnosis | Polyarticular Juvenile Idiopathic Arthritis (PJIA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting a diagnosis of moderately to severely active polyarticular juvenile idiopathic arthritis

AND

2 - One of the following:

2.1 History of failure, contraindication, or intolerance to both of the following:

- Humira (adalimumab)*
- Enbrel (etanercept)*

OR

2.2 Patient is currently on Actemra therapy as documented by claims history or medical records (document drug, date, and duration of therapy)

AND

3 - Patient is not receiving Actemra in combination with ANY of the following:

- Biologic disease modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

4 - Prescribed by, or in consultation with, a rheumatologist

Notes *May require PA

| Product Name: Actemra Actpen, Actemra SQ | |
|--|---|
| Diagnosis | Systemic Juvenile Idiopathic Arthritis (SJIA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting a diagnosis of active systemic juvenile idiopathic arthritis

AND

2 - Patient is not receiving Actemra in combination with ANY of the following:

- Biologic disease modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

3 - Prescribed by, or in consultation with, a rheumatologist

| Product Name: Actemra Actpen, Actemra SQ | |
|--|--|
| Diagnosis | Rheumatoid Arthritis, Polyarticular Juvenile Idiopathic Arthritis (PJIA), Systemic Juvenile Idiopathic Arthritis (SJIA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting positive clinical response to Actemra therapy

AND

2 - Patient is not receiving Actemra in combination with ANY of the following:

- Biologic disease modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

3 - Prescribed by, or in consultation with, a rheumatologist

| Product Name: Actemra Actpen, Actemra SQ, Actemra IV | |
|--|-----------------------|
| Diagnosis | Giant Cell Arteritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting a diagnosis of giant cell arteritis

AND

2 - One of the following:

2.1 History of failure, contraindication, or intolerance to ONE glucocorticoid (e.g., prednisone)

OR

2.2 Patient is currently on Actemra therapy as documented by claims history or medical records (document drug, date, and duration of therapy)

AND

3 - Patient is not receiving Actemra in combination with ANY of the following:

- Biologic disease modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

4 - Prescribed by or in consultation with a rheumatologist

| Product Name: Actemra Actpen, Actemra SQ, Actemra IV | |
|--|----------------------|
| Diagnosis | Giant Cell Arteritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting positive clinical response to Actemra therapy

AND

2 - Patient is not receiving Actemra in combination with ANY of the following:

- Biologic disease modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

3 - Prescribed by, or in consultation with, a rheumatologist

| Product Name: Actemra Actpen, Actemra SQ | |
|--|---|
| Diagnosis | Systemic Sclerosis-Associated Interstitial Lung Disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting a diagnosis of active systemic sclerosis-associated interstitial lung disease (SSc-ILD) as documented by ALL of the following:

1.1 ONE of the following:

1.1.1 Skin thickening of the fingers of both hands extending proximal to the metacarpophalangeal joints

OR

1.1.2 TWO of the following:

- Skin thickening of the fingers (e.g., puffy fingers, sclerodactyly of the fingers)
- Fingertip lesions (e.g., digital tip ulcers, fingertip pitting scars)
- Telangiectasia
- Abnormal nailfold capillaries
- Pulmonary arterial hypertension
- Raynaud's phenomenon
- SSc-related autoantibodies (e.g., anticentromere, anti-topoisomerase I, anti-RNA polymerase III)

AND

1.2 Presence of interstitial lung disease as determined by finding evidence of pulmonary fibrosis on HRCT (high-resolution computed tomography), involving at least 10% of the lungs

AND

2 - Patient is not receiving Actemra in combination with ANY of the following:

- Biologic disease modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

3 - Prescribed by, or in consultation with, a pulmonologist

| Product Name: Actemra Actpen, Actemra SQ | |
|--|---|
| Diagnosis | Systemic Sclerosis-Associated Interstitial Lung Disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting positive clinical response to Actemra therapy

AND

2 - Patient is not receiving Actemra in combination with ANY of the following:

- Biologic disease modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

3 - Prescribed by, or in consultation with, a pulmonologist

2. Revision History

| Date | Notes |
|-----------|---|
| 3/28/2022 | Added Actemra IV formulation to GCA criteria. Updated all criteria re garding concomitant use to indicate no options are allowed. Added S ubmission of Records to all criteria boxes. |

Acthar Gel, Cortrophin Gel

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-102899 | Acthar Gel, Cortrophin Gel |
|-----------|---|
| Formulary | Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) |

Formulary Note

Guideline Note:

| Effective Date: | 2/4/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Acthar Gel | | |
|--|--|--|
| Diagnosis | Infantile spasm (i.e., West Syndrome)* | |
| Approval Length | 4 Week(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of infantile spasms (i.e., West Syndrome)* | | |

AND

2 - Patient is less than 2 years old

AND

3 - Both of following:

3.1 Initial dose: 75 units per meters squared intramuscular (IM) twice daily for 2 weeks

AND

3.2 After 2 weeks, dose should be tapered according to the following schedule: 30 units per meters squared IM in the morning for 3 days; 15 units per meters squared IM in the morning for 3 days; 10 units per meters squared IM in the morning for 3 days; 10 units per meters squared IM every other morning for 6 days (3 doses)

| Notes | *Note: Acthar Gel is not medically necessary for treatment of acute ex |
|-------|--|
| | acerbations of multiple sclerosis. |

| Product Name: Acthar Gel, Cortrophin | |
|--------------------------------------|---|
| Diagnosis | Opsoclonus-myoclonus syndrome (i.e., OMS, Kinsbourne Syndrome)* |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of Opsoclonus-myoclonus syndrome (i.e., OMS, Kinsbourne Syndrome)*

AND

2 - For Cortrophin requests ONLY: Trial and failure or intolerance to Acthar Gel (verified via paid pharmacy claims or submission of medical records/chart notes)

| Notes | *Note: Acthar Gel is not medically necessary for treatment of acute ex |
|-------|--|
| | acerbations of multiple sclerosis. |

| Date | Notes |
|----------|---|
| 2/3/2022 | Added step through Acthar to get Cortrophin [for OMS Syndrome crit eria (not indicated for infantile spasms)] |

Actimmune

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99673 Actimmune

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Actimmune | |
|-------------------------|-------------------------------------|
| Diagnosis | Chronic Granulomatous Disease (CGD) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of chronic granulomatous disease

| Product Name: Actimmune | |
|-------------------------|-------------------------------------|
| Diagnosis | Chronic Granulomatous Disease (CGD) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

1 - Patient does not show evidence of progressive disease while on Actimmune

| Product Name: Actimmune | |
|-------------------------|---------------------------------|
| Diagnosis | Severe, Malignant Osteopetrosis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of severe, malignant osteopetrosis

| Product Name: Actimmune | |
|-------------------------|---------------------------------|
| Diagnosis | Severe, Malignant Osteopetrosis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Actimmune

| Product Name: Actimmune | |
|-------------------------|-----------------------------|
| Diagnosis | Primary Cutaneous Lymphomas |
| Approval Length | 12 month(s) |

| Therapy Stage | Initial Authorization |
|----------------|-----------------------|
| Guideline Type | Prior Authorization |
| | |
| | |

- 1 Patient has ONE of the following diagnoses:
 - Mycosis fungoides (MF)
 - Sézary syndrome (SS)

| Product Name: Actimmune | |
|-------------------------|-----------------------------|
| Diagnosis | Primary Cutaneous Lymphomas |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Actimmune

| Product Name: Actimmune | |
|-------------------------|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Actimmune will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Actimmune | |
|-------------------------|---------------------------|
| Diagnosis | NCCN Recommended Regimens |

| Approval Length | 12 month(s) |
|-----------------|---------------------|
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | • |

1 - Documentation of positive clinical response to Actimmune therapy

| Date | Notes |
|----------|--------------------|
| 6/7/2021 | 7.1 Implementation |

Adakveo

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99677 Adakveo

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Adakveo | | |
|-----------------------|---------------------|--|
| Diagnosis | Sickle cell disease | |
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |

1 - Diagnosis of sickle cell disease, identified by any genotype

| AND | | | | |
|--|--|--|--|--|
| 2 - ONE of the following: | | | | |
| 2.1 BOTH of the following: | | | | |
| Age 16 to 20 years Prescriber attests the service is medically necessary to correct or ameliorate a defect, a condition, or a physical or mental illness in an eligible patient | | | | |
| OR | | | | |
| 2.2 Age greater than or equal to 21 years | | | | |
| AND | | | | |
| 3 - Patient has experienced at least two vaso-occlusive crises within the past 12 months | | | | |

| Date | Notes |
|----------|--------------------|
| 4/8/2021 | 7/1 Implementation |

Adbry (tralokinumab-ldrm)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-104390 Adbry (tralokinumab-ldrm)

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 4/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Indications

Drug Name: Adbry (tralokinumab-ldrm)

Atopic Dermatitis Indicated for the treatment of moderate-to-severe atopic dermatitis in adult patients whose disease is not adequately controlled with topical prescription therapies or when those therapies are not advisable. May be used with or without topical corticosteroids.

2. Criteria

| Product Name: Adbry | | |
|---------------------|-----------------------|--|
| Diagnosis | Atopic Dermatitis | |
| Approval Length | 6 Months* | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

| Approval Criteria | | | | |
|---|--|--|--|--|
| 1 - Diagnosis of moderate to severe atopic dermatitis | | | | |
| AND | | | | |
| | | | | |
| 2 - Submission of documentation (e.g., chart notes) demonstrating one of the following: | | | | |
| Involvement of at least 10% body surface area (BSA) SCORing Atopic Dermatitis (SCORAD) index value of at least 25 [A] | | | | |
| AND | | | | |
| 3 - Patient is 18 years of age or older | | | | |
| AND | | | | |
| 4 - Prescribed by or in consultation with one of the following: | | | | |
| Dermatologist Allergist/Immunologist | | | | |
| AND | | | | |
| 5 - Trial and failure of a minimum 30-day supply (14-day supply for topical corticosteroids), contraindication, or intolerance to at least TWO of the following (verified via paid claims or submission of records): | | | | |
| Medium or higher potency topical corticosteroid Pimecrolimus cream^ Tacrolimus ointment Eucrisa (crisaborole) ointment^ | | | | |
| Notes*QL Override (For new starts only): Enter 2 PAs as follows: First PA: Approve 6 syringes per 28 days for one month; Second PA: Approve 4 syringes per 28 days (no overrides needed) for the remaining 11 months. (Adbry is hard-coded with a quantity of 4 syringes per 28 days); ^Product may require step therapy | | | | |

| Product Name: Adbry | | |
|---------------------|---------------------|--|
| Diagnosis | Atopic Dermatitis | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |

1 - Submission of documentation (e.g., chart notes) demonstrating positive clinical response to therapy as evidenced by at least ONE of the following:

- Reduction in body surface area involvement from baseline
- Reduction in SCORing Atopic Dermatitis (SCORAD) index value from baseline [A]

3. Background

| Clinical Practice Guidelines | | | | |
|------------------------------|--------------------------------------|-------------------------|-----------------|--|
| Class | Drug | Dosage Form | Strength (%) | |
| Very high potency | Augmented betamethasone dipropionate | Ointment, gel | 0.05 | |
| | Clobetasol propionate | Cream, foam, ointment | 0.05 | |
| | Diflorasone diacetate | Ointment | 0.05 | |
| | Halobetasol propionate | Cream, ointment | 0.05 | |
| High Potency | Amcinonide | Cream, lotion, ointment | 0.1 | |
| | Augmented betamethasone dipropionate | Cream, lotion | 0.05 | |

| | Betamethasone dipropionate | Cream, foam, ointment, solution | 0.05 |
|------------------------|----------------------------|---------------------------------|-------|
| | Desoximetasone | Cream, ointment | 0.25 |
| | Desoximetasone | Gel | 0.05 |
| | Diflorasone diacetate | Cream | 0.05 |
| | Fluocinonide | Cream, gel, ointment, solution | 0.05 |
| | Halcinonide | Cream, ointment | 0.1 |
| | Mometasone furoate | Ointment | 0.1 |
| | Triamcinolone acetonide | Cream, ointment | 0.5 |
| Medium | Betamethasone valerate | Cream, foam, lotion, ointment | 0.1 |
| potency Clocortolone p | Clocortolone pivalate | Cream | 0.1 |
| | Desoximetasone | Cream | 0.05 |
| | Fluocinolone acetonide | Cream, ointment | 0.025 |
| | Flurandrenolide | Cream, ointment, lotion | 0.05 |
| | Fluticasone propionate | Cream | 0.05 |
| | Fluticasone propionate | Ointment | 0.005 |
| | Mometasone furoate | Cream, lotion | 0.1 |
| | Triamcinolone acetonide | Cream, ointment, lotion | 0.1 |
| Lower- | Hydrocortisone butyrate | Cream, ointment, solution | 0.1 |
| medium potency | Hydrocortisone probutate | Cream | 0.1 |
| | Hydrocortisone valerate | Cream, ointment | 0.2 |
| | Prednicarbate | Cream | 0.1 |
| Low | Alclometasone dipropionate | Cream, ointment | 0.05 |
| potency | Desonide | Cream, gel, foam, ointment | 0.05 |
| | Fluocinolone acetonide | Cream, solution | 0.01 |
| | Dexamethasone | Cream | 0.1 |

| Lowest potency | Hydrocortisone | Cream, lotion, ointment, solution | 0.25, 0.5, 1 |
|-------------------|------------------------|-----------------------------------|--------------|
| | Hydrocortisone acetate | Cream, ointment | 0.5-1 |

| Date | Notes |
|----------|-------------|
| 3/4/2022 | New Program |

ADHD Agents - AZM

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-109904 ADHD Agents - AZM

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 8/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

Product Name: Brand Adderall, generic amphetamine/dextroamphetamine tablets, brand Adderall XR, brand Concerta ER, Daytrana, generic dexmethylphenidate tablets, brand Focalin XR, brand Methylin solution, generic methylphenidate tablets, brand Ritalin LA, generic methylphenidate ER (CD) capsules, Vyvanse capsules, generic atomoxetine, generic clonidine ER, generic guanfacine ER, generic dextroamphetamine tablets

| Diagnosis | ADHD Medications for Use in Children Under 6 Years Old |
|-------------------|--|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - The requesting clinician has documented that the child has a diagnosis of attention deficit hyperactivity disorder (ADHD)

AND

2 - The requesting clinician has documented that psychosocial issues have been evaluated before request for ADHD medications

AND

3 - The requesting clinician has documented non-medication alternatives that have been attempted before request for ADHD medications

AND

4 - The requested dose does NOT exceed the Food and Drug Administration (FDA) recommended maximum daily dosage unless the provider has submitted clinical justification for the dose exceeding the FDA maximum

Product Name: Non-Preferred Drugs: Brand Adhansia XR, Brand Adzenys XR-ODT, generic amphetamine ER, generic amphetamine, generic amphetamine/dextroamphetamine capsules, Brand Aptensio XR, Brand Azstarys, Brand Cotempla XR-ODT, Brand Desoxyn, Brand Dexedrine, generic dexmethylphenidate ER, generic dexmethylphenidate capsules, generic dexmethylphenidate ER, generic dextroamphetamine capsules, generic dextroamphetamine ER, Brand Dyanavel XR (oral suspension and chewable tablets), Brand Evekeo ODT, Brand Focalin, Brand Intuniv, Brand Jornay PM, generic methamphetamine, generic methylphenidate capsule, generic methylphenidate ER tablets, generic methylphenidate ER (LA), Brand Mydayis, Brand Procentra, Brand Qelbree, Brand Quillichew ER, Brand Quillivant XR, generic relexxii, Brand Ritalin, Brand Strattera, Brand Vyvanse Chewables, Brand Zenzedi

| Approval Length | 12 month(s) |
|-----------------|---------------------|
| Guideline Type | Prior Authorization |

Approval Criteria

1 - The patient has a history of failure, contraindication, or intolerance to a trial to FOUR of the following preferred products*:

| Brand Adderall Brand Concerta Daytrana generic dexmeta Brand Focalin 2 Brand Methylin generic methylt Brand Ritalin La generic methylt Vyvanse capsu generic atomoxic generic guanfa | tamine/dextroamphetamine tablets XR a ER thylphenidate tablets XR solution ohenidate tablets A ohenidate ER (CD) capsules iles tetine he ER |
|---|--|
| Notes | *Alternatives may require prior authorization |

| Date | Notes |
|-----------|---|
| 7/28/2022 | Added Dyanavel XR chewable tablets as target to NP section. |

Aduhelm (aducanumab-avwa)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-107262 Aduhelm (aducanumab-avwa)

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 5/17/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Aduhelm | |
|--|--|
| Diagnosis | Alzheimer's Disease - MEDICARE PART B* |
| Approval Length | 6 month(s) |
| Guideline Type | Medicare Part B |
| | |
| Approval Criteria | |
| 1 - Requested medication is billed through Medicare Part B | |

AND

2 - Submission of documentation confirming patient is enrolled in a CMS approved prospective comparative study

| Notes | *Note: THIS SECTION SHOULD ONLY BE USED FOR DUAL ELIGIB LE MEMBERS (WILL HAVE AZMDUAL PLAN CODE) COVERED UN |
|-------|--|
| | DER MEDICARE PART B THAT ARE REQUESTING SECONDARY COVERAGE. |

| Product Name: Aduhelm | |
|-----------------------|--|
| Diagnosis | Alzheimer's Disease - MEDICARE PART D* |
| Approval Length | None |
| Guideline Type | Prior Authorization requests from providers from Medicare Part D for Dual Eligible Members |

Approval Criteria

1 - Requested medication is billed through Medicare Part D

AND

2 - Requests for coverage of Aduhelm (aducanumab) are not authorized and will not be approved under Part D

| *Note: THIS SECTION SHOULD ONLY BE USED FOR DUAL ELIGIB LE MEMBERS (WILL HAVE AZMDUAL PLAN CODE). APPROVAL L ENGTH: NONE - REQUESTS FOR ADUHELM ARE NOT COVERED UNDER MEDICARE PART D AND SHALL BE DENIED AS A BENEF IT EXCLUSION. |
|--|
| T EAGEOSION. |

| Product Name: Aduhelm | |
|-----------------------|---------------------------------------|
| Diagnosis | Alzheimer's Disease - FEE-FOR-SERVICE |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

1 - Diagnosis of one of the following:

- Mild cognitive impairment (MCI) due to Alzheimer's Disease (AD)
- Mild dementia due to Alzheimer's Disease (AD)

AND

2 - Submission of medical records (e.g., chart notes, laboratory values, examination histories) documenting the basis for diagnosis, including all of the following:

2.1 Documentation of a comprehensive history and neurological examination, inclusive of a description of the nature and duration of cognitive symptoms within the previous 3 months

AND

2.2 Medical records documenting baseline (within the previous three months) cognitive function based on ONE of the following objective assessments:

- Mini-Mental State Examination (MMSE) score ≥ 24
- Montreal Cognitive Assessment (MoCA) score ≥ 15

AND

2.3 Medical records documenting confirmed evidence of clinically significant AD neuropathology based on ONE of the following:

- Cerebral Spinal Fluid (CSF) biomarkers
- Amyloid positron emission tomography (PET)

AND

3 - Patient has received recent (within the previous 3 months) baseline brain magnetic resonance imaging (MRI) prior to initiating treatment

AND

4 - Patient does not have significant cerebrovascular disease as established by brain MRI showing any of the following:

- Acute or sub-acute hemorrhage
- Prior macro-hemorrhage or prior subarachnoid hemorrhage (unless finding is not due to an underlying structural or vascular hemorrhage)
- 4 or more brain microhemorrhages
- Cortical infarct
- More than 1 lacunar infarct
- Superficial siderosis
- History of diffuse white matter disease

AND

- **5** Patient does not have any of the following non-AD neurodegenerative disorders:
 - Probable dementia with Lewy bodies by consensus criteria
 - Suspected frontotemporal degeneration
 - Dementia in down syndrome

AND

6 - Patient does not have any of the following exclusionary neurological or psychiatric conditions:

- Uncontrolled seizure disorder
- Uncontrolled mood disorder, anxiety disorder, or psychosis
- Substance use disorder active in the past 2 years

AND

7 - Patient does not have any of the following cardiovascular conditions:

- Uncontrolled hypertension
- Coronary artery disease (including unstable angina and myocardial infarction)
- Heart failure
- Arrhythmia
- Clinically significant carotid atherosclerosis and/or peripheral arterial disease

8 - Both of the following: Patient is not currently taking an anticoagulant or antiplatelet agent (unless aspirin 325 • mg/day or less) Patient has no history of transient ischemic attack (TIA), stroke, or unexplained loss of consciousness within previous year prior to initiating treatment AND 9 - Patient does not have any uncontrolled clinically significant chronic medical condition (e.g., liver disease, kidney disease, pulmonary disease, autoimmune disease requiring chronic immunosuppression, malignant neoplasm, active chronic infection [HIV, HCV], poorly controlled diabetes mellitus) AND 10 - Prescribed dosing is in accordance with the United States Food and Drug Administration approved labeling AND 11 - Prescribed by or in consultation with one of the following: Neurologist Geriatrics specialist AND **12** - Prescriber attests that the patient and/or authorized representative (e.g., power of attorney, invoked health care proxy) has shared in decision-making and has been informed on the known and potential risks and lack of established clinical benefit associated with Aduhelm (aducanumab-avwa) treatment AND

13 - Therapy should be discontinued permanently and the request should be denied if one or more of the following apply:

- If the patient has had ≥ 10 new incident microhemorrhages, regardless of clinical severity (including asymptomatic)
- If the patient had a serious event [Serious events include concern for immediate risk of death (a life-threatening event); inpatient hospitalization or prolongation of existing hospitalization due to symptoms; new persistent or significant disability/incapacity]
- If the patient has had ≥ 3 new incident areas of superficial siderosis, regardless of clinical severity (including asymptomatic) therapy should be discontinued permanently and the request should be denied

| Notes | *NOTE: If the patient has had ≥10 new incident microhemorrhages, re gardless of clinical severity (including asymptomatic) therapy should b e discontinued permanently and the request should be denied. *NOTE : If the patient had a serious event, therapy should be discontinued. † *NOTE: If the patient has had ≥3 new incident areas of superficial side rosis, regardless of clinical severity (including asymptomatic) therapy should be discontinued permanently and the request should be denie d. †Serious events include concern for immediate risk of death (a life-t hreatening event); inpatient hospitalization or prolongation of existing hospitalization due to symptoms; new persistent or significant disabilit y/incapacity. ‡Requests should be evaluated case-by-case with clinic al review and MD advisor. |
|-------|--|

| Product Name: Aduhelm | | |
|---|------------|--|
| Diagnosis Alzheimer's Disease - FEE-FOR-SERVICE | | |
| Approval Length | 6 month(s) | |
| Therapy Stage Reauthorization | | |
| Guideline Type Prior Authorization | | |

1 - Prescribed dosing is in accordance with the United States Food and Drug Administration approved labeling

AND

2 - Follow-up MRIs have been conducted at the following timeframes:

- Week 14 (after 4th infusion, prior to first 6 mg/kg dose)
- Week 22 (after 6th infusion, prior to first 10 mg/kg dose)
- Week 30 (after 8th infusion, prior to third 10 mg/kg dose)
- Week 42 (after 11th infusion, prior to sixth 10 mg/kg dose)

• Every 6 months thereafter

AND

3 - Patient's diagnosis continues to be mild cognitive impairment or mild dementia stage due to Alzheimer's disease as established by one of the following examination scales:

3.1 One of the following:

- Mini Mental State Exam (MMSE) score ≥ 24
- Montreal Cognitive Assessment (MoCA) score ≥ 15

OR

3.2 Both of the following:

- MMSE <24 or MoCA <15
- Rate of decline was slower than expected (<2 points/year)

AND

4 - ONE of the following (ARIA-H, microhemorrhages):

- Patient has had no new incident microhemorrhage
- Patient has had 1 to 4 new incident microhemorrhage(s) AND microhemorrhages are asymptomatic (no clinical symptoms)
- Patient has had 5 to 9 new incident microhemorrhages AND microhemorrhages are asymptomatic (no clinical symptoms) AND the microhemorrhages have been stabilized
- Patient has had 1 to 9 new incident microhemorrhages AND microhemorrhages resulted in mild, moderate or severe clinical symptoms AND the microhemorrhages have been stabilized

AND

5 - ONE of the following (ARIA-H, superficial siderosis)

- Patient has had no new incident areas of superficial siderosis
- Patient has had 1 new incident area of superficial siderosis AND superficial siderosis is asymptomatic (no clinical symptoms)

- Patient has had 2 new incident areas of superficial siderosis AND superficial siderosis is asymptomatic (no clinical symptoms) AND the superficial siderosis has been stabilized
- Patient has had 1 to 2 new incident areas of superficial siderosis AND superficial siderosis resulted in mild, moderate or severe clinical symptoms AND the superficial siderosis has been stabilized

AND

- **6** ONE of the following (ARIA-E)
 - Patient has had no new ARIA-E
 - Patient has mild ARIA-E on MRI AND ARIA-E is asymptomatic (no clinical symptoms)
 - Patient has had moderate or severe ARIA-E on MRI AND ARIA-E is asymptomatic (no clinical symptoms) AND the ARIA-E is stable
 - Patient has had mild, moderate or severe ARIA-E on MRI AND ARIA-E resulted in mild, moderate or severe clinical symptoms AND the ARIA-E is stable

AND

- 7 One of the following:
- 7.1 Patient does not meet ANY of the following:
 - Initiation of anticoagulation
 - Development of active immune-mediated/autoimmune conditions (e.g., Crohn's disease, SLE, aplastic anemia, myasthenia gravis, meningitis/encephalitis)
 - Initiation of immunomodulatory medications (e.g., cancer immunotherapies, rituximab, azathioprine)
 - Development of other neurologic conditions (e.g., intracerebral bleeds, TBI, stroke)

OR

7.2 BOTH of the following:

- Patient does meet one of the above
- Prescriber documents clinical rationale for continued use of aducanumab‡

AND

8 - Prescribed by or in consultation with one of the following:

- Neurologist
- Geriatric specialist

AND

9 - Therapy should be discontinued permanently and the request should be denied if one or more of the following apply:

- If the patient has had ≥ 10 new incident microhemorrhages, regardless of clinical severity (including asymptomatic)
- If the patient had a serious event [Serious events include concern for immediate risk of death (a life-threatening event); inpatient hospitalization or prolongation of existing hospitalization due to symptoms; new persistent or significant disability/incapacity]
- If the patient has had ≥ 3 new incident areas of superficial siderosis, regardless of clinical severity (including asymptomatic) therapy should be discontinued permanently and the request should be denied

| Notes | *NOTE: If the patient has had ≥10 new incident microhemorrhages, re gardless of clinical severity (including asymptomatic) therapy should b e discontinued permanently and the request should be denied. *NOTE : If the patient had a serious event, therapy should be discontinued. † *NOTE: If the patient has had ≥3 new incident areas of superficial side rosis, regardless of clinical severity (including asymptomatic) therapy should be discontinued permanently and the request should be denie d. †Serious events include concern for immediate risk of death (a life-t hreatening event); inpatient hospitalization or prolongation of existing hospitalization due to symptoms; new persistent or significant disabilit y/incapacity. ‡Requests should be evaluated case-by-case with clinic al review and MD advisor. |
|-------|--|

2. Background

| Clinical Pr | ractice Guideline | S | | |
|-------------|-------------------|--------------|-----------------------|----|
| Appendix | | | | |
| | ARIA - H (Micro | hemorrhages) | | |
| | | New Inc | cident Microhemorrhag | es |
| | | Ra | adiographic Severity | |

| | | Mild (1 to 4) | Moderate (5 to 9) | Severe (≥10) |
|-------------------------------------|--------------------|---|--|---------------------|
| Clinical Sympto m Severity | Asymptomatic | Continue treatment; MRI q4w until stable | Suspend treatment; MRI q4w until stable; Restart once stable | Stop Permanently |
| Seventy | Mild | Suspend treatment; MRI q4w until stable | | Stop Permanently |
| | Moderate Severe | Restart once stable an resolved | nd clinical symptoms | |
| | Serious | | Stop Permanently | |
| | | | | |
| | ARIA - H (Supe | rficial Siderosis) | | |
| | | | of Superficial Siderosi | s (Central Read) |
| | | Ra | adiographic Severity | |
| | | Mild (1) | Moderate (2) | Severe (≥3) |
| Clinical Sympto m | Asymptomatic | Continue treatment; MRI q4w until stable | Suspend treatment; MRI q4w until stable; Restart once stable | Stop Permanently |
| Severity | Mild | Suspend treatment; MRI q4w until stable | | Stop Permanently |
| | Moderate | Restart once stable ar resolved | nd clinical symptoms | 1 official official |
| | Severe | Tesolveu | | |
| | Serious | | Stop Permanently | |
| | | | | |
| | <u>ARIA - E</u> | | | - |
| | | | everity on MRI (Central | Read) |
| | | Ra | adiographic Severity | |
| | | Mild | Moderate | Severe |
| Clinical Sympto m | Asymptomatic | Continue treatment; MRI q4w until stable | Suspend treatment; MRI q4w until stable; Restart once stable | |
| Severity | Mild | | | |

| | Moderate Severe | Suspend treatment; MRI q4w until stable Restart once stable and clinical symptoms resolved |
|---|--------------------|--|
| : | Serious | Stop Permanently |

| Date | Notes |
|-----------|--|
| 5/17/2022 | Updated Medicare sections for clarification. |

Aemcolo

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99426 Aemcolo

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Aemcolo | | | |
|--------------------------------------|---------------------|--|--|
| Approval Length | 1 month(s) | | |
| Guideline Type | Prior Authorization | | |
| | | | |
| Approval Criteria | Approval Criteria | | |
| 1 - Diagnosis of travelers' diarrhea | | | |
| | | | |
| | AND | | |
| | | | |

2 - History of failure, contraindication, or intolerance to ONE of the following:

- Azithromycin (generic Zithromax)
 Ciprofloxacin (generic Cipro)
 Levofloxacin (generic Levaquin)
 Ofloxacin (generic Floxin)

| Date | Notes |
|-----------|---|
| 3/10/2021 | Bulk Copy C&S Arizona to Arizona Standard |

Afinitor

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99709 Afinitor

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | | |
|--|--------------------------------|--|
| Diagnosis | iagnosis Neuroendocrine tumors | |
| Approval Length | 12 month(s) | |
| Therapy Stage Initial Authorization | | |
| Guideline Type Prior Authorization | | |

Approval Criteria

- **1** Diagnosis of one of the following:
 - Neuroendocrine tumors of pancreatic origin

- Neuroendocrine tumors of gastrointestinal origin Neuroendocrine tumors of lung origin Neuroendocrine tumors of thymic origin •
- ٠
- •

| , , | 0 |
|---|-----|
| | AND |
| 2 - Disease is progressive | |
| | AND |
| 3 - One of the following: | |
| Disease is unresectable Disease is locally advanced Disease is metastatic | |

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|-----------------------|
| Diagnosis | Neuroendocrine Tumors |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

1 - Patient does not show evidence of progressive disease while on Afinitor therapy

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|-----------------------|
| Diagnosis | Renal cell cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

| 1 - Diagnosis of renal cell cancer |
|--|
| AND |
| 2 - One of the following: |
| 2.1 Disease has relapsed |
| OR |
| 2.2 BOTH of the following |
| Medically or surgically unresectable tumor Diagnosis of Stage IV disease |
| AND |
| 3 - One of the following: |
| 3.1 Patient with non-clear cell histology |
| OR |
| 3.2 Both of the following: |
| 3.2.1 Patient with predominantly clear cell histology |
| AND |
| 3.2.2 History of failure, contraindication, or intolerance to at least one prior systemic therapy [e.g., Nexavar (sorafenib), Sutent (sunitinib), Opdivo (nivolumab), Cabometyx (cabozantinib)] |

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|-------------------|
| Diagnosis | Renal cell cancer |
| Approval Length | 12 month(s) |

| Therapy Stage | Reauthorization |
|----------------|---------------------|
| Guideline Type | Prior Authorization |
| | |

1 - Patient does not show evidence of progressive disease while on Afinitor therapy

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|--|
| Diagnosis | Renal Angiomyolipoma with Tuberous Sclerosis Complex |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of renal angiomyolipoma and tuberous sclerosis complex (TSC), not requiring immediate surgery

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|--|
| Renal Angiomyolipoma with Tuberous Sclerosis Complex | |
| 12 month(s) | |
| Reauthorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Afinitor therapy

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|--|
| Diagnosis | Subependymal Giant Cell Astrocytoma Associated with Tuberous Sclerosis Complex |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

1 - Diagnosis of subependymal giant cell astrocytoma (SEGA) associated with tuberous sclerosis (TS)

AND

2 - Patient is not a candidate for curative surgical resection

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|--|
| Diagnosis | Subependymal Giant Cell Astrocytoma Associated with Tuberous Sclerosis Complex |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Afinitor therapy

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|--|
| Waldenströms Macroglobulinemia or Lymphoplasmacytic Lymphoma | |
| 12 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |

Approval Criteria

- **1** Diagnosis of one of the following:
 - Waldenströms macroglobulinemia
 - Lymphoplasmacytic lymphoma

AND

- 2 One of the following:
 - Disease is non-responsive to primary treatment Disease is progressive Disease has relapsed •
 - •
 - •

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|--|
| Diagnosis | Waldenströms Macroglobulinemia or Lymphoplasmacytic Lymphoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Afinitor therapy

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | | |
|--|-----------------------|--|
| Diagnosis | Breast Cancer | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Diagnosis of breast cancer

AND

- 2 One of the following:
 - 2.1 Disease is recurrent

2.2 Disease is metastatic

AND

OR

3 - One of the following:

3.1 Disease is hormone receptor positive (HR+) [i.e., estrogen-receptor-positive (ER+) or progesterone-receptor-positive (PR+)]

OR

3.2 BOTH of the following:

- Disease is hormone receptor negative (HR-)
- Disease has clinical characteristics that predict a HR+ tumor

AND

4 - Disease is human epidermal growth factor receptor 2 (HER2)-negative

AND

5 - One of the following:

5.1 Patient is a postmenopausal woman

OR

5.2 BOTH of the following:

- Patient is a premenopausal woman
- Patient is being treated with ovarian ablation/suppression

OR

5.3 Patient is male

AND

6 - One of the following:

6.1 Both of the following:

6.1.1 Used in combination with Aromasin (exemestane)

AND

6.1.2 One of the following:

6.1.2.1 Disease progressed while on or within 12 months of non-steroidal aromatase inhibitor [e.g., Arimidex (anastrozole), Femara (letrozole)] therapy

OR

6.1.2.2 Patient was treated with tamoxifen at any time

OR

6.2 Used in combination with ONE of the following:

• Fulvestrant

• Tamoxifen

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|-----------------|
| Diagnosis | Breast Cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |

| Guideline Type | Prior Authorization |
|----------------|---------------------|
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Afinitor therapy

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|-----------------------|
| Diagnosis | Hodgkin Lymphoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of classical Hodgkin lymphoma

AND

- **2** ONE of the following:
 - •
 - Disease is refractory Disease has relapsed

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|---------------------|
| Diagnosis | Hodgkin Lymphoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Afinitor therapy

| Product Name: Brand | Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|---|--|
| Diagnosis | PEComa (perivascular epitheliod cell tumor), recurrent angiomyolipoma, lymphangioleiomyomatosis, or gastrointestinal stromal tumor (GIST) | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| Approval Criteria 1 - Diagnosis of PECor | na (perivascular epitheliod cell tumor) | |
| | OR | |
| 2 - Diagnosis of recurre | 2 - Diagnosis of recurrent angiomyolipoma | |
| | OR | |
| 3 - Diagnosis of lympha | angioleiomyomatosis | |
| | OR | |
| 4 - All of the following: | | |
| 4.1 Diagnosis of Gastrointestinal Stromal Tumor (GIST) | | |
| | AND | |
| 4.2 Disease has prog | ressed after single agent therapy with ONE of the following: | |
| Gleevec (imatinib) Sutent (sunitinib) Stivarga (regorafenib) | | |
| AND | | |

4.3 Used in combination with ONE of the following:

- Gleevec (imatinib)
- Sutent (sunitinib)
- Stivarga (regorafenib)

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|---|
| Diagnosis | PEComa (perivascular epitheliod cell tumor), recurrent angiomyolipoma, lymphangioleiomyomatosis, or gastrointestinal stromal tumor (GIST) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Afinitor therapy

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|-----------------------------|
| Diagnosis | Thymic Carcinoma or Thymoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- 1 One of the following:
 - Diagnosis of thymic carcinoma
 - Diagnosis of thymoma

AND

2 - ONE of the following:

2.1 History of failure, contraindication, or intolerance to at least one prior first-line chemotherapy regimen

OR

2.2 Patient has extrathoracic metastatic disease

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|-----------------------------|
| Diagnosis | Thymic Carcinoma or Thymoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Afinitor therapy

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|--|
| Diagnosis | Follicular carcinoma, Hürthle cell carcinoma, or papillary carcinoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- **1** Diagnosis of ONE of the following:
 - Follicular carcinoma
 - Hürthle cell carcinoma
 - Papillary carcinoma

AND

2 - ONE of the following:

- Unresectable locoregional recurrent disease •
- Persistent disease
- Metastatic disease

- 3 ONE of the following:
 - Patient has symptomatic disease Patient has progressive disease ٠
 - •

AND

4 - Disease is refractory to radioactive iodine treatment

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|--|
| Diagnosis | Follicular carcinoma, Hürthle cell carcinoma, or papillary carcinoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Afinitor therapy

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|--|
| Meningioma | |
| 12 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Diagnosis of meningioma

2 - Disease is recurrent or progressive

AND

3 - Surgery and/or radiation is not possible

AND

4 - Used in combination with bevacizumab (e.g., Avastin, Myasi)

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|---------------------|
| Diagnosis | Meningioma |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Afinitor therapy

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|-----------------------|
| Diagnosis | Endometrial Carcinoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| · · · | |

Approval Criteria

1 - Diagnosis of endometrial carcinoma

2 - Used in combination with letrozole

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|-----------------------|
| Diagnosis | Endometrial Carcinoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Afinitor therapy

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|--|
| Diagnosis | Tuberous Sclerosis Complex associated Partial-Onset Seizures |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of tuberous sclerosis complex associated partial-onset seizures

AND

2 - Used as adjunctive therapy

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|--|
| Diagnosis | Tuberous Sclerosis Complex associated Partial-Onset Seizures |
| Approval Length | 12 month(s) |

| Therapy Stage | Reauthorization |
|----------------|---------------------|
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Afinitor therapy

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Use supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Brand Afinitor, generic everolimus, Afinitor Disperz | |
|--|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Afinitor therapy

| Date | Notes |
|-----------|-------------------------------------|
| 5/12/2021 | Arizona Medicaid 7.1 Implementation |

Afrezza

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99427 Afrezza

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Afrezza | |
|---------------------------|------------------------------------|
| Diagnosis | Type 1 or Type 2 diabetes mellitus |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - One of the following: | |

1.1 Diagnosis of type 1 diabetes mellitus and used in combination with a basal insulin or continuous insulin pump OR **1.2** Diagnosis of type 2 diabetes mellitus AND 2 - Patient is unable to self-inject medications (e.g. Humalog, Lantus, Levemir) due to ONE of the following: Physical impairment • Visual impairment ٠ Lipohypertrophy • Documented needle-phobia to the degree that the patient has previously refused any • injectable therapy or medical procedure (refer to DSM-5 for specific phobia diagnostic criteria) AND 3 - Forced Expiratory Volume (FEV1) within the last 60 days is greater than or equal to 70% of expected normal as determined by the physician AND 4 - Afrezza will not be approved in patients with ONE of the following: Who smoke cigarettes • Who recently quit smoking (within the past 6 months) • With chronic lung disease (e.g. asthma, chronic obstructive pulmonary disease) •

| Product Name: Afrezza | |
|-----------------------|------------------------------------|
| Diagnosis | Type 1 or Type 2 diabetes mellitus |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |

| Guideline Type | Prior Authorization | |
|--|---|--|
| | | |
| Approval Criteria | | |
| · | | |
| | 1 - Repeat pulmonary function test confirms that patient has NOT experienced a decline of 20% or more in Forced Expiratory Volume (FEV1) | |
| | | |
| | AND | |
| | | |
| 2 - Patient continues following: | to be unable to self-inject short-acting insulin due to ONE of the | |
| Physical impairment Visual impairment Lipohypertrophy Documented needle-phobia to the degree that the patient has previously refused any injectable therapy or medical procedure (refer to DSM-5 for specific phobia diagnostic criteria) | | |
| ontonay | | |
| AND | | |
| | | |
| | | |

3 - Patient continues to not smoke cigarettes

| Date | Notes |
|-----------|---|
| 3/10/2021 | Bulk Copy C&S Arizona to Arizona Standard |

Aldurazyme - Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99428 | Aldurazyme - Arizona |
|----------|----------------------|
|----------|----------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Aldurazyme | |
|---|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - One of the following: | |
| 1.1 Confirmed diagnosis of Hurler and Hurler-Scheie forms of Mucopolysaccharidosis I (MPS I) | |

| OR |
|---|
| 1.2 Both the following: |
| 1.2.1 Confirmed diagnosis of Scheie form of Mucopolysaccharidosis I (MPS I) |
| AND |
| 1.2.2 Have moderate to severe symptoms |

| Date | Notes |
|-----------|---|
| 3/10/2021 | Bulk Copy C&S Arizona to Arizona Standard |

Alecensa

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99674 Alecensa

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Alecensa | |
|---|------------------------------------|
| Diagnosis | Non-Small Cell Lung Cancer (NSCLC) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of non-small cell lung cancer (NSCLC) | |

2 - Disease is one of the following:

- Metastatic
- Recurrent

AND

3 - Tumor is anaplastic lymphoma kinase (ALK)-positive

| Product Name: Alecensa | |
|------------------------|------------------------------------|
| Diagnosis | Non-Small Cell Lung Cancer (NSCLC) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Alecensa therapy

| Product Name: Alecensa | |
|------------------------|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Alecensa will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Alecensa | |
|---|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Documentation of positive clinical response to Alecensa therapy | |

| Date | Notes |
|----------|--------------------|
| 6/3/2021 | 7/1 Implementation |

Alinia

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99429 Alinia

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Alinia, generic nitazoxanide | |
|--|------------------------------------|
| Diagnosis | Diarrhea caused by Giardia lamblia |
| Approval Length | 3 Day(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of giardiasis | |

2 - History of failure, contraindication, or intolerance to metronidazole

| Product Name: Brand Alinia, generic nitazoxanide | |
|--|--|
| Diarrhea caused by Cryptosporidium parvum | |
| 12 month(s) | |
| Prior Authorization | |
| | |
| | |

Approval Criteria

1 - Diagnosis of cryptosporidiosis

| Date | Notes |
|-----------|---|
| 3/10/2021 | Bulk Copy C&S Arizona to Arizona Standard |

Alpha Interferons - AZM

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-105169 A | Alpha Interferons - | AZM |
|-------------|---------------------|-----|
|-------------|---------------------|-----|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 4/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Intron A | | |
|---|---------------------|--|
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of hairy cell leukemia | | |
| OR | | |

2 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of condylomata acuminata (genital or perianal) OR 3 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of AIDS-related Kaposi's sarcoma OR 4 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of leptomeningeal metastases OR 5 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of meningiomas OR 6 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of kidney cancer OR 7 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting treatment of myeloproliferative neoplasms (MPNs) such as essential thrombocythemia (ET), polycythemia vera (PV), or primary myelofibrosis (PM) OR 8 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of follicular lymphoma

9 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of adult T-cell leukemia, lymphoma

OR

OR

10 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of mycosis fungoides, Sézary syndrome

OR

11 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of desmoid tumors/aggressive fibromatosis

OR

12 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of giant cell tumor of the bone

OR

13 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of malignant melanoma

| Product Name: Alferon N | |
|-------------------------|---------------------|
| Approval Length | 8 Week(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting treatment of refractory or recurring external condylomata acuminata (genital or venereal warts) due to the human papillomavirus (HPV) infection

| Date | Notes |
|-----------|--|
| 3/24/2022 | Removed Sylatron from guideline, Added Submission of Medical Rec ords |

Alzheimer's Agents - AZM

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-109871 Alzheimer's Agents - AZM

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 7/27/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

Product Name: Brand Aricept, generic donepezil, Brand Namenda/Namenda XR, generic
memantine/memantine XR, Brand Razadyne, generic galantamine hydrobromide, Brand
Razadyne ER, generic galantamine ERApproval Length12 month(s)

| pp: e . eg | |
|----------------|---------------------|
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of dementia of the Alzheimer's type

Product Name: Brand Exelon, generic rivastigmine

| Approval Length | 12 month(s) | | |
|---|---|--|--|
| Guideline Type | Prior Authorization | | |
| | | | |
| Approval Criteria | Approval Criteria | | |
| 1 - Diagnosis of dementia of the Alzheimer's type | | | |
| | | | |
| OR | | | |
| | | | |
| 2 - Diagnosis of deme | entia associated with Parkinson's disease | | |

| Product Name: Adlarity | / |
|--|--|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of demen | itia of the Alzheimer's type |
| | AND |
| | |
| 2 - One of the following | j: |
| 2.1 History of failure, of (verified via paid pharm | contraindication or intolerance to ALL of the following preferred drugs* nacy claims): |
| generic donepezil | |
| generic galantamine IR/ER generic memantine | |
| generic oral riva | |
| | |
| | OR |
| | |
| 2.2 Both of the followi | ng: |
| 2.2.1 History of failure (verified via paid pharm | e, contraindication or intolerance to generic rivastigmine patch* nacy claims) |

2.2.2 Patient is unable to swallow oral formulations or has documented swallowing difficulties

| Notes *PA may be required |
|---------------------------|
|---------------------------|

| Date | Notes |
|-----------|---|
| 7/27/2022 | Added XR formulations of Namenda/memantine to product name sec tion. No change to criteria. |

Ampyra

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99666 Ampyra

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Ampyra, generic dalfampridine ER | | |
|--|-----------------------|--|
| Diagnosis | Multiple Sclerosis | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| | | |

Approval Criteria

1 - Diagnosis of multiple sclerosis

2 - Physician confirmation that patient has difficulty walking (e.g., timed 25-foot walk test)

| Product Name: Brand Ampyra, generic dalfampridine ER | |
|--|---------------------|
| Diagnosis | Multiple Sclerosis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Physician confirmation that the patient's walking improved with Ampyra therapy

| Date | Notes |
|-----------|---|
| 3/18/2021 | Bulk Copy C&S Arizona SP to Medicaid Arizona SP for 7/1 eff |

Amvuttra (vutrisiran)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-114478 | Amvuttra | (vutrisiran) |
|-----------|----------|--------------|
|-----------|----------|--------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Amvuttra | |
|------------------------|--|
| 12 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Submission of documentation (e.g., chart notes) confirming diagnosis of hereditary transthyretin-mediated amyloidosis (hATTR amyloidosis) with polyneuropathy

| AND | |
|--|--|
| 2 - Patient has a transthyretin (TTR) mutation (e.g., V30M) | |
| AND | |
| 3 - Two of the following: | |
| Patient has a baseline polyneuropathy disability (PND) score less than or equal to IIII Patient has a baseline familial amyloidotic polyneuropathy (FAP) stage of 1 or 2 Patient has a baseline neuropathy impairment score (NIS) greater than or equal to 5 and less than or equal to 130 Patient has a baseline Karnofsky Performance Status score greater than or equal to 60% | |
| AND | |
| 4 - Presence of clinical signs and symptoms of the disease (e.g., peripheral/autonomic neuropathy, walking ability, quality of life) | |
| AND | |
| 5 - Patient has not had a liver transplant | |
| AND | |
| 6 - Prescribed by or in consultation with a neurologist | |

| Product Name: Amvuttra | |
|------------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| | |

Approval Criteria

1 - Submission of documentation (e.g., chart notes) confirming positive clinical response to therapy as evidenced by an improvement in clinical signs and symptoms from baseline (e.g., neuropathy, quality of life, gait speed, nutritional status, decrease in serum TTR level)

AND

2 - Two of the following:

- Patient continues to have a polyneuropathy disability (PND) score less than or equal to IIIb
- Patient continues to have a familial amyloidotic polyneuropathy (FAP) stage of 1 or 2
- Patient continues to have a neuropathy impairment score (NIS) greater than or equal to 5 and less than or equal to 130
- Patient continues to have a Karnofsky Performance Status score greater than or equal to 60%

| Date | Notes |
|-----------|-------------|
| 9/26/2022 | New Program |

Anthelmintics - Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99431 | Anthelmintics - Arizona |
|----------|-------------------------|
|----------|-------------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Albenza, generic albendazole | | |
|--|---------------------|--|
| Diagnosis | See Note section* | |
| Approval Length | 1 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of Enterobius vermicularis (pinworm) | | |

| OR |
|--|
| 2 - Diagnosis of Hydatid Disease [Echinococcosis (Tapeworm)] |
| OR |
| 3 - Diagnosis of Ancylostoma/Necatoriasis (Hookworm) |
| OR |
| 4 - Diagnosis of Ascariasis (Roundworm) |
| OR |
| 5 - Diagnosis of Mansonella perstans (Filariasis) |
| OR |
| 6 - Diagnosis of Toxocariasis (Roundworm) |
| OR |
| 7 - Diagnosis of Trichinellosis |
| OR |
| 8 - Diagnosis of Trichuriasis (Whipworm) |
| OR |
| 9 - Diagnosis of Capillariasis |

| Notes | * Enterobius vermicularis (pinworm), Hydatid Disease [Echinococcosis (Tapeworm)] Ancylostoma/Necatoriasis (Hookworm), Ascariasis (Rou |
|-------|---|
| | ndworm), Mansonella perstans (Filariasis), Toxocariasis (Roundworm) , Trichinellosis, Trichuriasis (Whipworm), Capillariasis |

| Product Name: Brand Albenza, generic albendazole | |
|--|--|
| Neurocysticercosis | |
| 6 month(s) | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Diagnosis of neurocysticercosis

| Product Name: Brand Stromectol, generic ivermectin | |
|--|---------------------|
| Approval Length | 1 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of intestinal strongyloidiasis due to the nematode parasite Strongyloides stercoralis

OR

2 - Diagnosis of onchocerciasis due to the nematode parasite Onchocerca volvulus

| Date | Notes |
|-----------|---|
| 3/10/2021 | Bulk Copy C&S Arizona to Arizona Standard |

Anticonvulsants - AZM

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-107448 Antice | onvulsants - AZM |
|------------------|------------------|
|------------------|------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 6/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Aptiom, Briviact, Brand Vimpat, generic lacosamide, Xcopri | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting ONE of the following:

1.1 All of the following:

1.1.1 Diagnosis of partial-onset seizures

1.1.2 History of greater than or equal to 8 week trial of at least TWO of the following (any release formulation qualifies):

- Carbamazepine*
- Divalproex*
- Gabapentin*
- Lamotrigine*
- Levetiracetam*
- Oxcarbazepine*
- Phenytoin*
- Pregabalin*
- Topiramate*
- Valproic acid*
- Zonisamide*

AND

1.1.3 One of the following:

1.1.3.3 Trial and failure, contraindication, or intolerance to generic lacosamide (APPLIES TO BRAND VIMPAT ONLY)

AND

1.1.3.1 Both of the following:

- Documented history of persisting seizures after titration to the highest tolerated dose with each medication trial
- Lack of compliance as a reason for treatment failure has been ruled out

OR

1.1.3.2 Both of the following:

- Documentation of failure due to intolerable side effects.
- Reasonable efforts were made to minimize the side effect (e.g. change timing of dosing, divide dose out for more frequent but smaller doses, etc.)

| OR | |
|---|----------------------|
| 1.2 For continuation of prior therapy for a seizure disorder | |
| Notes | *Drug may require PA |

| Product Name: Fycompa | |
|-----------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting ONE of the following:

1.1 All of the following:

1.1.1 Diagnosis of partial-onset or primary generalized tonic-clonic seizures

AND

1.1.2 History of greater than or equal to 8 week trial of at least TWO of the following (any release formulation qualifies):

- Carbamazepine*
- Divalproex*
- Gabapentin*
- Lamotrigine*
- Levetiracetam*
- Oxcarbazepine*
- Phenytoin*
- Pregabalin*
- Topiramate*
- Valproic acid*
- Zonisamide*

AND

1.1.3 One of the following:

| 1.1.3.1 Both of the fo | bllowing: |
|-------------------------|---|
| with each medie | story of persisting seizures after titration to the highest tolerated dose cation trial Ince as a reason for treatment failure has been ruled out |
| | OR |
| 1.1.3.2 Both of the fo | bllowing: |
| Reasonable eff | of failure due to intolerable side effects. orts were made to minimize the side effect (e.g. change timing of lose out for more frequent but smaller doses, etc.) |
| | OR |
| 1.2 For continuation of | f prior therapy for a seizure disorder |
| Notes | *Drug may require PA |

| Product Name: Epidiolex | |
|-------------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting ONE of the following:

1.1 Diagnosis of seizures associated with Dravet syndrome or tuberous sclerosis complex

OR

1.2 All of the following:

1.2.1 Diagnosis of seizures associated with Lennox-Gastaut syndrome

1.2.2 History of greater than or equal to 8 week trial, contraindication or intolerance of at least TWO of the following (any release formulation qualifies):

- Banzel (rufinamide)*
- Clobazam*
- Divalproex*
- Felbamate*
- Lamotrigine*
- Topiramate*
- Valproic acid*

AND

1.2.3 One of the following:

1.2.3.1 Both of the following:

- Documented history of persisting seizures after titration to the highest tolerated dose with each medication trial
- Lack of compliance as a reason for treatment failure has been ruled out

OR

1.2.3.2 Both of the following:

- Documentation of failure due to intolerable side effects.
- Reasonable efforts were made to minimize the side effect (e.g. change timing of dosing, divide dose out for more frequent but smaller doses, etc.)

OR

1.3 For continuation of prior therapy for a seizure disorder

| Notes | *Drug may require PA |
|-------|----------------------|
|-------|----------------------|

Product Name: Brand Onfi, generic clobazam

| Approval Length | 12 month(s) |
|--|---|
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Submission of medi the following: | cal records (e.g., chart notes, lab work, imaging) documenting one of |
| 1.1 Diagnosis of seizu | ires associated with Lennox-Gastaut syndrome |
| | OR |
| 1.2 Both of the followi | ng: |
| Diagnosis of Dravet syndrome Patient is currently taking Diacomit | |
| | OR |
| 2 - For continuation of | orior therapy for a seizure disorder |

| Product Name: Brand Banzel, generic rufinamide | |
|---|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| Approval Criteria | |
| 1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of seizures associated with Lennox-Gastaut syndrome | |

OR

2 - For continuation of prior therapy for a seizure disorder

Product Name: Brand Gabitril, generic tiagabine

| Approval Length | 12 month(s) | |
|--|---|--|
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting ONE of the following: | | |
| 1.1 All of the following: | | |
| 1.1.1 Diagnosis of pa | rtial-onset seizures | |
| | | |
| | AND | |
| 1.1.2 Used as adjunctive therapy (defined as accessory treatment used in combination to enhance primary treatment) | | |
| | AND | |
| 1.1.3 Not used as pri | mary treatment | |
| AND | | |
| 1.1.4 History of great release formulation qua | er than or equal to 8 week trial of at least TWO of the following (any alifies): | |
| Carbamazepine Divalproex* Gabapentin* Lamotrigine* Levetiracetam* Oxcarbazepine* Phenytoin* Pregabalin* Topiramate* Valproic acid* Zonisamide* | | |

| 1.2 For continuation o | f prior therapy for a seizure disorder |
|------------------------|--|
| Notes | *Drug may require PA |

| Product Name: Sympazan | |
|------------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting ONE of the following:

1.1 ALL of the following:

1.1.1 Diagnosis of seizures associated with Lennox-Gastaut syndrome (LGS)

AND

1.1.2 BOTH of the following:

- Used as adjunctive therapy (defined as accessory treatment used in combination to enhance primary treatment.)
- Not used as primary treatment

AND

1.1.3 History of greater than or equal to 8 week trial, contraindication or intolerance of at least TWO of the following (any release formulation qualifies):

- Divalproex*
- Lamotrigine*
- Topiramate*
- Valproic acid*
- Felbamate*
- Banzel*

AND

1.1.4 Prescriber provides a reason or special circumstance the patient cannot use generic clobazam tablets or suspension

OR

1.2 ALL of the following:

1.2.1 Diagnosis of refractory partial onset seizures (four or more uncontrolled seizures per month after an adequate trial of at least two antiepileptic drugs)

AND

1.2.2 BOTH of the following:

- Used as adjunctive therapy (defined as accessory treatment used in combination to enhance primary treatment.)
- Not used as primary treatment

AND

1.2.3 History of greater than or equal to 8 week trial of at least TWO of the following (any release formulation qualifies):

- Carbamazepine*
- Divalproex^{*}
- Gabapentin*
- Lamotrigine*
- Levetiracetam*
- Oxcarbazepine*
- Phenytoin*
- Pregabalin*
- Topiramate*
- Valproic acid*
- Zonisamide*

AND

1.2.4 Prescriber provides a reason or special circumstance the patient cannot use generic clobazam tablets or suspension

OR

1.3 ALL of the following:

1.3.1 Diagnosis of Dravet syndrome

AND

1.3.2 Patient is currently taking Diacomit

AND

1.3.3 Prescriber provides a reason or special circumstance the patient cannot use generic clobazam tablets or suspension

OR

1.4 For continuation of prior therapy for a seizure disorder

| Notes | *Drug may require PA |
|-------|----------------------|
|-------|----------------------|

| Product Name: Brand Sabril Oral Solution, generic vigabatrin oral solution, generic vigadrone oral solution | |
|---|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of infantile spasms

| 2 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting all of th following: | |
|---|--|
| 2.1 Diagnosis of complex partial seizures | |
| | |
| AND | |
| 2.2 Used as adjunctive therapy (defined as accessory treatment used in combination to enhance primary treatment) | |
| AND | |
| | |
| 2.3 Not used as primary treatment | |
| | |
| AND | |
| 2.4 History of greater than or equal to 8 week trial of at least TWO of the following (any release formulation qualifies): Carbamazepine* Divalproex* Gabapentin* Lamotrigine* Levetiracetam* Oxcarbazepine* Phenytoin* Pregabalin* Topiramate* Valproic acid* Zonisamide* | |
| OR | |
| | |
| 3 - For continuation of prior therapy for a seizure disorder | |
| Notes *Drug may require PA | |
| | |

| Product Name: Brand Sabril Tablets, generic vigabatrin tablets | |
|--|-------------|
| Approval Length | 12 month(s) |

| <u> </u> | | |
|---|--|--|
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Submission of med the following: | ical records (e.g., chart notes, lab work, imaging) documenting ONE of | |
| 1.1 All of the following | j: | |
| 1.1.1 Diagnosis of co | omplex partial seizures | |
| | AND | |
| 1.1.2 Used as adjund enhance primary treatr | ctive therapy (defined as accessory treatment used in combination to nent) | |
| | AND | |
| 1.1.3 Not used as primary treatment | | |
| | AND | |
| 1.1.4 History of great release formulation quant | er than or equal to 8 week trial of at least TWO of the following (any alifies): | |
| Carbamazepine* Divalproex* Gabapentin* Lamotrigine* Levetiracetam* Oxcarbazepine* Phenytoin* Pregabalin* Topiramate* Valproic acid* Zonisamide* | | |
| | OR | |
| 1.2 For continuation c | of prior therapy for a seizure disorder | |

| Notes *Drug may require PA | Notes | *Drug may require PA |
|----------------------------|-------|----------------------|
|----------------------------|-------|----------------------|

| Product Name: Diacomit | |
|------------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of Dravet syndrome and currently taking clobazam

OR

2 - For continuation of prior therapy for a seizure disorder

| Product Name: Fintepla | |
|------------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting ALL of the following:

1.1 Diagnosis of seizures associated with Dravet syndrome

AND

1.2 History of greater than or equal to 8-week trial of at least TWO of the following (any release formulation qualifies):

- Divalproex (e.g., generic Depakote)
- Levetiracetam (e.g., generic Keppra)
- Topiramate (e.g., generic Topamax)
- Valproic acid (e.g., generic Depakene)
- Zonisamide (generic Zonegran)

1.3 ONE of the following:

1.3.1 BOTH of the following:

1.3.1.1 Documented history of persisting seizures after titration to the highest tolerated dose with each medication trial

AND

1.3.1.2 Lack of compliance as a reason for treatment failure has been ruled out

OR

1.3.2 BOTH of the following:

1.3.2.1 Documentation of failure due to intolerable side effects

AND

1.3.2.2 Reasonable efforts were made to minimize the side effect (e.g., change timing of dosing, divide dose out for more frequent but smaller doses, etc.)

OR

2 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting ALL of the following:

2.1 Diagnosis of seizures associated with Lennox-Gastaut syndrome

AND

2.2 History of greater than or equal to 8 week trial, contraindication or intolerance of at least TWO of the following (any release formulation qualifies):

• Banzel (rufinamide)*

| Clobazam* Divalproex* Felbamate* Lamotrigine* Topiramate* Valproic Acid* | |
|---|--|
| | AND |
| 2.3 ONE of the followi | ng: |
| 2.3.1 BOTH of the fol | lowing: |
| with each medic | story of persisting seizures after titration to the highest tolerated dose cation trial nce as a reason for treatment failure has been ruled out |
| | OR |
| 2.3.2 BOTH of the fol | lowing: |
| | of failure due to intolerable side effects nce as a reason for treatment failure has been ruled out |
| | OR |
| 3 - For continuation of p | prior therapy for a seizure disorder |
| Notes | *Drug may require PA |

2. Revision History

| Date | Notes |
|-----------|--|
| 5/24/2022 | Added generic lacosamide as target. Added criteria for Fintepla's ne w indication of Lennox-Gastaut Syndrome |

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-107448 Anticonvulsants - AZM

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 6/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Aptiom, Briviact, Brand Vimpat, generic lacosamide, Xcopri | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting ONE of the following:

1.1 All of the following:

1.1.1 Diagnosis of partial-onset seizures

1.1.2 History of greater than or equal to 8 week trial of at least TWO of the following (any release formulation qualifies):

- Carbamazepine*
- Divalproex*
- Gabapentin*
- Lamotrigine*
- Levetiracetam*
- Oxcarbazepine*
- Phenytoin*
- Pregabalin*
- Topiramate*
- Valproic acid*
- Zonisamide*

AND

1.1.3 One of the following:

1.1.3.3 Trial and failure, contraindication, or intolerance to generic lacosamide (APPLIES TO BRAND VIMPAT ONLY)

AND

1.1.3.1 Both of the following:

- Documented history of persisting seizures after titration to the highest tolerated dose with each medication trial
- Lack of compliance as a reason for treatment failure has been ruled out

OR

1.1.3.2 Both of the following:

- Documentation of failure due to intolerable side effects.
- Reasonable efforts were made to minimize the side effect (e.g. change timing of dosing, divide dose out for more frequent but smaller doses, etc.)

| | OR |
|---|----------------------|
| 1.2 For continuation of prior therapy for a seizure disorder | |
| Notes | *Drug may require PA |

| Product Name: Fycompa | |
|-----------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting ONE of the following:

1.1 All of the following:

1.1.1 Diagnosis of partial-onset or primary generalized tonic-clonic seizures

AND

1.1.2 History of greater than or equal to 8 week trial of at least TWO of the following (any release formulation qualifies):

- Carbamazepine*
- Divalproex*
- Gabapentin*
- Lamotrigine*
- Levetiracetam*
- Oxcarbazepine*
- Phenytoin*
- Pregabalin*
- Topiramate*
- Valproic acid*
- Zonisamide*

AND

1.1.3 One of the following:

| 1.1.3.1 Both of the fo | bllowing: |
|-------------------------|---|
| with each medie | story of persisting seizures after titration to the highest tolerated dose cation trial Ince as a reason for treatment failure has been ruled out |
| | OR |
| 1.1.3.2 Both of the fo | bllowing: |
| Reasonable eff | of failure due to intolerable side effects. orts were made to minimize the side effect (e.g. change timing of lose out for more frequent but smaller doses, etc.) |
| | OR |
| 1.2 For continuation of | f prior therapy for a seizure disorder |
| Notes | *Drug may require PA |

| Product Name: Epidiolex | |
|-------------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting ONE of the following:

1.1 Diagnosis of seizures associated with Dravet syndrome or tuberous sclerosis complex

OR

1.2 All of the following:

1.2.1 Diagnosis of seizures associated with Lennox-Gastaut syndrome

1.2.2 History of greater than or equal to 8 week trial, contraindication or intolerance of at least TWO of the following (any release formulation qualifies):

- Banzel (rufinamide)*
- Clobazam*
- Divalproex*
- Felbamate*
- Lamotrigine*
- Topiramate*
- Valproic acid*

AND

1.2.3 One of the following:

1.2.3.1 Both of the following:

- Documented history of persisting seizures after titration to the highest tolerated dose with each medication trial
- Lack of compliance as a reason for treatment failure has been ruled out

OR

1.2.3.2 Both of the following:

- Documentation of failure due to intolerable side effects.
- Reasonable efforts were made to minimize the side effect (e.g. change timing of dosing, divide dose out for more frequent but smaller doses, etc.)

OR

1.3 For continuation of prior therapy for a seizure disorder

| Notes | *Drug may require PA |
|-------|----------------------|
|-------|----------------------|

Product Name: Brand Onfi, generic clobazam

| Approval Length | 12 month(s) |
|---|---|
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Submission of medi the following: | cal records (e.g., chart notes, lab work, imaging) documenting one of |
| 1.1 Diagnosis of seizu | ires associated with Lennox-Gastaut syndrome |
| | OR |
| 1.2 Both of the following | ng: |
| Diagnosis of Dra Patient is currer | avet syndrome ntly taking Diacomit |
| | OR |
| 2 - For continuation of p | prior therapy for a seizure disorder |

| Product Name: Brand Banzel, generic rufinamide | |
|--|--|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | cal records (e.g., chart notes, lab work, imaging) documenting ssociated with Lennox-Gastaut syndrome |

OR

2 - For continuation of prior therapy for a seizure disorder

Product Name: Brand Gabitril, generic tiagabine

| Approval Length | 12 month(s) |
|--|--|
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Submission of medi the following: | cal records (e.g., chart notes, lab work, imaging) documenting ONE of |
| 1.1 All of the following | j: |
| 1.1.1 Diagnosis of pa | rtial-onset seizures |
| | AND |
| | |
| 1.1.2 Used as adjunc enhance primary treatm | ctive therapy (defined as accessory treatment used in combination to nent) |
| | AND |
| 1.1.3 Not used as pri | mary treatment |
| | AND |
| 1.1.4 History of great release formulation qua | er than or equal to 8 week trial of at least TWO of the following (any alifies): |
| Carbamazepine Divalproex* Gabapentin* Lamotrigine* Levetiracetam* Oxcarbazepine* Phenytoin* Pregabalin* Topiramate* Valproic acid* Zonisamide* | |

| 1.2 For continuation o | f prior therapy for a seizure disorder |
|------------------------|--|
| Notes | *Drug may require PA |

| Product Name: Sympazan | |
|------------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting ONE of the following:

1.1 ALL of the following:

1.1.1 Diagnosis of seizures associated with Lennox-Gastaut syndrome (LGS)

AND

1.1.2 BOTH of the following:

- Used as adjunctive therapy (defined as accessory treatment used in combination to enhance primary treatment.)
- Not used as primary treatment

AND

1.1.3 History of greater than or equal to 8 week trial, contraindication or intolerance of at least TWO of the following (any release formulation qualifies):

- Divalproex*
- Lamotrigine*
- Topiramate*
- Valproic acid*
- Felbamate*
- Banzel*

AND

1.1.4 Prescriber provides a reason or special circumstance the patient cannot use generic clobazam tablets or suspension

OR

1.2 ALL of the following:

1.2.1 Diagnosis of refractory partial onset seizures (four or more uncontrolled seizures per month after an adequate trial of at least two antiepileptic drugs)

AND

1.2.2 BOTH of the following:

- Used as adjunctive therapy (defined as accessory treatment used in combination to enhance primary treatment.)
- Not used as primary treatment

AND

1.2.3 History of greater than or equal to 8 week trial of at least TWO of the following (any release formulation qualifies):

- Carbamazepine*
- Divalproex^{*}
- Gabapentin*
- Lamotrigine*
- Levetiracetam*
- Oxcarbazepine*
- Phenytoin*
- Pregabalin*
- Topiramate*
- Valproic acid*
- Zonisamide*

AND

1.2.4 Prescriber provides a reason or special circumstance the patient cannot use generic clobazam tablets or suspension

OR

1.3 ALL of the following:

1.3.1 Diagnosis of Dravet syndrome

AND

1.3.2 Patient is currently taking Diacomit

AND

1.3.3 Prescriber provides a reason or special circumstance the patient cannot use generic clobazam tablets or suspension

OR

1.4 For continuation of prior therapy for a seizure disorder

| Notes | *Drug may require PA |
|-------|----------------------|
|-------|----------------------|

| Product Name: Brand Sabril Oral Solution, generic vigabatrin oral solution, generic vigadrone oral solution | |
|---|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of infantile spasms

| 2 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting all of the following: |
|---|
| 2.1 Diagnosis of complex partial seizures |
| |
| AND |
| 2.2 Used as adjunctive therapy (defined as accessory treatment used in combination to enhance primary treatment) |
| AND |
| |
| 2.3 Not used as primary treatment |
| |
| AND |
| 2.4 History of greater than or equal to 8 week trial of at least TWO of the following (any release formulation qualifies): Carbamazepine* Divalproex* Gabapentin* Lamotrigine* Levetiracetam* Oxcarbazepine* Phenytoin* Pregabalin* Topiramate* Valproic acid* Zonisamide* |
| OR |
| |
| 3 - For continuation of prior therapy for a seizure disorder |
| Notes *Drug may require PA |
| |

| Product Name: Brand Sabril Tablets, generic vigabatrin tablets | |
|--|-------------|
| Approval Length | 12 month(s) |

| <u> </u> | | |
|---|--|--|
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Submission of med the following: | ical records (e.g., chart notes, lab work, imaging) documenting ONE of | |
| 1.1 All of the following | j: | |
| 1.1.1 Diagnosis of co | omplex partial seizures | |
| | AND | |
| 1.1.2 Used as adjund enhance primary treatr | ctive therapy (defined as accessory treatment used in combination to nent) | |
| | AND | |
| 1.1.3 Not used as primary treatment | | |
| | AND | |
| 1.1.4 History of great release formulation quant | er than or equal to 8 week trial of at least TWO of the following (any alifies): | |
| Carbamazepine Divalproex* Gabapentin* Lamotrigine* Levetiracetam* Oxcarbazepine Phenytoin* Pregabalin* Topiramate* Valproic acid* Zonisamide* | | |
| | OR | |
| 1.2 For continuation c | of prior therapy for a seizure disorder | |

| Notes *Drug may require PA | Notes | *Drug may require PA |
|----------------------------|-------|----------------------|
|----------------------------|-------|----------------------|

| Product Name: Diacomit | |
|------------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of Dravet syndrome and currently taking clobazam

OR

2 - For continuation of prior therapy for a seizure disorder

| Product Name: Fintepla | |
|------------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting ALL of the following:

1.1 Diagnosis of seizures associated with Dravet syndrome

AND

1.2 History of greater than or equal to 8-week trial of at least TWO of the following (any release formulation qualifies):

- Divalproex (e.g., generic Depakote)
- Levetiracetam (e.g., generic Keppra)
- Topiramate (e.g., generic Topamax)
- Valproic acid (e.g., generic Depakene)
- Zonisamide (generic Zonegran)

1.3 ONE of the following:

1.3.1 BOTH of the following:

1.3.1.1 Documented history of persisting seizures after titration to the highest tolerated dose with each medication trial

AND

1.3.1.2 Lack of compliance as a reason for treatment failure has been ruled out

OR

1.3.2 BOTH of the following:

1.3.2.1 Documentation of failure due to intolerable side effects

AND

1.3.2.2 Reasonable efforts were made to minimize the side effect (e.g., change timing of dosing, divide dose out for more frequent but smaller doses, etc.)

OR

2 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting ALL of the following:

2.1 Diagnosis of seizures associated with Lennox-Gastaut syndrome

AND

2.2 History of greater than or equal to 8 week trial, contraindication or intolerance of at least TWO of the following (any release formulation qualifies):

• Banzel (rufinamide)*

| Clobazam* Divalproex* Felbamate* Lamotrigine* Topiramate* Valproic Acid* | | |
|---|--|--|
| | AND | |
| 2.3 ONE of the followi | ng: | |
| 2.3.1 BOTH of the fol | lowing: | |
| Documented history of persisting seizures after titration to the highest tolerated dose with each medication trial Lack of compliance as a reason for treatment failure has been ruled out | | |
| | OR | |
| 2.3.2 BOTH of the fol | lowing: | |
| | of failure due to intolerable side effects nce as a reason for treatment failure has been ruled out | |
| | OR | |
| 3 - For continuation of p | prior therapy for a seizure disorder | |
| Notes | *Drug may require PA | |

2. Revision History

| Date | Notes |
|-----------|--|
| 5/24/2022 | Added generic lacosamide as target. Added criteria for Fintepla's ne w indication of Lennox-Gastaut Syndrome |

Antidepressants - AZM

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-112734 Antidepressants - AZM

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 8/27/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: generic citalopram oral solution, generic fluoxetine oral solution, generic sertraline oral conc for solution | |
|---|--|
| Diagnosis | Requests for Patients greater than 12 years of age |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - The member is unable to swallow the oral tablet/capsule.

Product Name: Amitriptyline, amoxapine, bupropion tabs/SR tabs/XL tabs (150 and 300mg), citalopram tabs/oral soln, clomipramine, desipramine, doxepin caps/oral conc for solution, duloxetine capsules (20, 30, 60mg), escitalopram, fluoxetine caps/oral soln, fluvoxamine IR, generic mirtazapine tabs/ODT, imipramine tabs/caps, nortriptyline caps/oral soln, paroxetine tabs, protriptyline, sertraline tabs/oral soln, trazodone, trimipramine, venlafaxine tabs/ER capsules

| Diagnosis | PREFERRED DRUG Requests for patient 6 years of age or younger |
|-----------------|---|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - The patient is unresponsive to other treatment modalities, unless contraindicated (i.e. other medications or behavioral modification attempted)

AND

2 - The physician attests that the requested medication is medically necessary. (Document rationale for use)

| Notes | Drug may require PA |
|-------|---------------------|
|-------|---------------------|

Product Name: Aplenzin, Brand Anafranil, Brand Celexa, generic citalopram capsules, Brand Cymbalta, generic duloxetine 40mg caps, Drizalma, Brand Effexor XR, generic venlafaxine ER tabs, Emsam, Fetzima, fluvoxamine ER, Brand Lexapro, maprotiline, Marplan, Brand Nardil, generic phenelzine, nefazodone, Brand Norpramin, Brand Pamelor caps/oral soln, Brand Parnate, generic tranylcypromine, Brand Paxil, generic paroxetine capsules, Paxil susp, Brand Paxil CR, generic paroxetine ER, Pexeva, Brand Pristiq, generic desvenlafaxine ER, Brand Prozac, generic fluoxetine tablets, Brand Remeron SLTB, Brand Remeron, Trintellix, Viibryd, Brand Wellbutrin SR, Brand Wellbutrin XL/Forfivo, generic bupropion ER (XL) 450mg tabs, Brand Zoloft, generic sertraline capsules

| Diagnosis | Non-Preferred Drugs |
|-----------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - The patient is unresponsive to other treatment modalities, unless contraindicated (i.e. other medications or behavioral modification attempted)

2 - The physician attests that the requested medication is medically necessary. (Document rationale for use)

AND

3 - Patient has a history of failure, contraindication or intolerance to at least 3 preferred alternatives*

- Bupropion (Generic Wellbutrin)
- Bupropion SR (Generic Wellbutrin SR)
- Bupropion XL (Generic Wellbutrin XL)
- Citalopram (Generic Celexa)
- Escitalopram Tablets (Generic Lexapro)
- Esketamine (Spravato)
- Fluoxetine Capsules (Generic Prozac)
- Fluoxetine Solution (Generic Prozac)
- Fluvoxamine Tablets (Generic Luvox)
- Mirtazapine (Generic Remeron)
- Paroxetine (Generic Paxil)
- Sertraline (Generic Zoloft)
- Trazodone (Generic Desyrel)
- Venlafaxine (Generic Effexor)
- Venlafaxine ER Capsules (Generic Effexor ER)

| Notes | *Drug may require PA |
|-------|----------------------|
|-------|----------------------|

| Product Name: Brand Venlafaxine besylate ER | |
|---|---------------------|
| Diagnosis | Non-Preferred Drugs |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - The patient is unresponsive to other treatment modalities, unless contraindicated (i.e. other medications or behavioral modification attempted)

2 - The physician attests that the requested medication is medically necessary. (Document rationale for use)

AND

3 - Patient has history of failure or intolerance to preferred generic venlafaxine or venlafaxine ER

AND

4 - Patient has a history of failure, contraindication or intolerance to at least 2 preferred alternatives*

- Bupropion (Generic Wellbutrin)
- Bupropion SR (Generic Wellbutrin SR)
- Bupropion XL (Generic Wellbutrin XL)
- Citalopram (Generic Celexa)
- Escitalopram Tablets (Generic Lexapro)
- Esketamine (Spravato)
- Fluoxetine Capsules (Generic Prozac)
- Fluoxetine Solution (Generic Prozac)
- Fluvoxamine Tablets (Generic Luvox)
- Mirtazapine (Generic Remeron)
- Paroxetine (Generic Paxil)
- Sertraline (Generic Zoloft)
- Trazodone (Generic Desyrel)

| Notes | *Drug may require PA |
|-------|----------------------|
|-------|----------------------|

2. Revision History

| Date | Notes |
|-----------|--|
| 8/26/2022 | Removed Caplyta from guideline, drug-specific guideline created. |

Antiemetics - Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99432 Antiemetics - Arizona

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Anzemet, granisetron tablet, ondansetron 24mg tablet | |
|--|---|
| Diagnosis | Nausea and vomiting associated with cancer chemotherapy |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | • |

Approval Criteria

1 - Prevention or treatment of nausea and vomiting associated with cancer chemotherapy

Product Name: Anzemet, granisetron tablet, ondansetron 24mg tablet

| Diagnosis | Nausea and vomiting associated with radiotherapy |
|-----------------|--|
| Approval Length | 3 month(s) |
| Guideline Type | Prior Authorization |

1 - Prevention or treatment of nausea and vomiting associated with radiotherapy (total body irradiation, single high-dose fraction to the abdomen, or daily fractions to the abdomen)

| Product Name: Anzemet, granisetron tablet, ondansetron 24mg tablet | |
|--|--------------------------------------|
| Diagnosis | Postoperative nausea and/or vomiting |
| Approval Length | 1 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Prevention of postoperative nausea and/or vomiting (administration prior to induction of anesthesia)

2. Revision History

| Date | Notes |
|-----------|---|
| 3/10/2021 | Bulk Copy C&S Arizona to Arizona Standard |

Antiglaucoma Agents - Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99587 | Antiglaucoma Agents - Arizona |
|----------|-------------------------------|
|----------|-------------------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Zioptan | | |
|--|---------------------|--|
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| Approval Criteria 1 - Diagnosis of elevated intraocular pressure due to ocular hypertension or open angle | | |
| glaucoma | | |

2. Revision History

| Date | Notes |
|------------|--|
| 10/25/2021 | Removed Azopt, Brand/generic Travatan Z as targets |

Antipsoriatic Agents

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99551 Antipsoriatic Agents

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|--------------------|-----------|
| P&T Approval Date: | |
| P&T Revision Date: | |

1. Criteria

| Product Name: Brand Dovonex cream, generic calcipotriene cream, Brand Calcitrene ointment, generic calcipotriene ointment, Brand Vectical, generic calcitriol ointment | |
|--|---------------------|
| Diagnosis | Psoriasis |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of psoriasis

AND

2 - History of failure, contraindication, or intolerance to TWO medium to Very high potency corticosteroid topical treatments (see Table 1 in Background section)

2. Background

Benefit/Coverage/Program Information

| - | Dosage Form | Strength |
|---|-----------------------------------|----------|
| Super High Potency | 1 | I |
| Augmented betamethasone dipropionate (Diprolene) | Gel, Ointment | 0.05% |
| Clobetasol propionate (Temovate, Temovate E) | Cream, Solution | 0.05% |
| Halobetasol propionate (Ultravate) | Cream | 0.05% |
| | | |
| dipropionate (Diprolene, | Cream, Lotion | 0.05% |
| dipropionate (Diprolene, Diprolene AF) | Cream, Lotion Lotion, Ointment | 0.05% |
| Augmented betamethasone dipropionate (Diprolene, Diprolene AF) Betamethasone dipropionate Fluocinonide (Lidex, Lidex E) | | |

| Betamethasone valerate (Beta- Val) | Cream | 0.1% |
|---------------------------------------|------------------------------|--------|
| Fluocinolone acetonide (Synalar) | Cream, Ointment | 0.025% |
| Fluticasone propionate (Cutivate) | Cream, Lotion | 0.05% |
| | Ointment | |
| | | 0.005% |
| Hydrocortisone butyrate (Locoid) | Ointment, Solution | 0.1% |
| Mometasone furoate (Elocon) | Cream, Ointment, Solution | 0.1% |
| Prednicarbate (Dermatop) | Cream | 0.1% |
| Triamcinolone acetonide (Kenalog) | Cream, Lotion, Ointment | 0.1% |
| | Ointment | |
| | | 0.025% |

Antipsychotics - AZM

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-114856 Antipsychotics - AZM

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/3/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

Product Name: haloperidol concentrate, haloperidol tablets, loxapine, perphenazine, thioridazine, thiothixene, generic pimozide, fluphenazine (tablets, concentrate and elixir), trifluoperazine, chlorpromazine (tabs and inj), lithium carbonate (caps, tabs and oral solution), generic lithium carbonate ER, generic aripiprazole tabs, generic ziprasidone, Latuda, generic risperidone (tabs and oral solution), risperidone ODT tabs, generic quetiapine, generic olanzapine (tabs and ODT tabs)

| Diagnosis | Preferred Antipsychotics: Children Under 6 Years Old |
|-----------------|--|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| | |

Approval Criteria

1 - The patient has been diagnosed per current DSM (Diagnostic and Statistical Manual of Mental Disorders) criteria with one of the following disorders:

- Bipolar Spectrum Disorder
- Schizophrenic Spectrum Disorder
- Tourette's or other tic disorder
- Autism Spectrum Disorder

AND

2 - The requesting clinician has documented that psychosocial issues have been evaluated before request for antipsychotic medications

AND

3 - The requesting clinician has documented non-medication alternatives that have been attempted before request for antipsychotic medications

AND

4 - The above documentation includes information on the expected outcomes and an evaluation of potential adverse events

AND

5 - The patient does not have a known hypersensitivity to the requested agent

| Product Name: Invega Sustenna | |
|-------------------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Patient has a diagnosis of schizophrenia or schizoaffective disorder

2 - ONE of the following:

2.1 BOTH of the following:

- Patient is non-adherent with oral atypical antipsychotic dosage forms
- Patient has established tolerability with oral paliperidone or oral risperidone

OR

2.2 Patient is unable to take oral solid alternatives

AND

3 - If the patient is less than 18 years of age, the prescriber attests they are aware of FDA (Food and Drug Administration) labeling regarding use of long acting injectable antipsychotic products in patients less than 18 years of age and feels the treatment with the requested product is medically necessary (Document rationale for use)

| Product Name: Risperdal Consta | |
|--------------------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

- 1 Patient has ONE of the following diagnoses:
 - Schizophrenia or schizoaffective disorder
 - Bipolar disorder

AND

- 2 ONE of the following:
- **2.1** BOTH of the following:

- Patient is non-adherent with oral atypical antipsychotic dosage forms
- Patient has established tolerability with oral risperidone

OR

2.2 Patient is unable to take oral solid alternatives

AND

3 - If the patient is less than 18 years of age, the prescriber attests they are aware of FDA (Food and Drug Administration) labeling regarding use of long acting injectable antipsychotic products in patients less than 18 years of age and feels the treatment with the requested product is medically necessary (Document rationale for use)

| Product Name: Abilify Maintena | |
|--------------------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| | |

- 1 Patient has ONE of the following diagnoses:
 - Schizophrenia or schizoaffective disorder
 - Bipolar disorder

AND

2 - ONE of the following:

2.1 BOTH of the following:

- Patient is non-adherent with oral atypical antipsychotic dosage forms
- Patient has established tolerability with aripiprazole

2.2 Patient is unable to take oral solid alternatives

AND

3 - If the patient is less than 18 years of age, the prescriber attests they are aware of FDA (Food and Drug Administration) labeling regarding use of long acting injectable antipsychotic products in patients less than 18 years of age and feels the treatment with the requested product is medically necessary (Document rationale for use)

| Product Name: Invega Trinza | | |
|-----------------------------|---------------------|--|
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Patient has a diagnosis of schizophrenia or schizoaffective disorder

AND

2 - Patient has been treated with Invega Sustenna for at least 4 months

AND

3 - If the patient is less than 18 years of age, the prescriber attests they are aware of FDA (Food and Drug Administration) labeling regarding use of long acting injectable antipsychotic products in patients less than 18 years of age and feels the treatment with the requested product is medically necessary (Document rationale for use)

| Product Name: Aristada, Aristada Initio | |
|---|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Patient has a diagnosis of schizophrenia or schizoaffective disorder

AND

2 - ONE of the following:

2.1 BOTH of the following:

- Patient is non-adherent with oral atypical antipsychotic dosage forms
- Patient has established tolerability with oral aripiprazole

OR

2.2 Patient is unable to take oral solid alternatives

AND

3 - If the patient is less than 18 years of age, the prescriber attests they are aware of FDA (Food and Drug Administration) labeling regarding use of long acting injectable antipsychotic products in patients less than 18 years of age and feels the treatment with the requested product is medically necessary (Document rationale for use)

| Product Name: Perseris | |
|------------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient has a diagnosis of schizophrenia or schizoaffective disorder

AND

2 - ONE of the following:

2.1 BOTH of the following:

- Patient is non-adherent with oral atypical antipsychotic dosage forms
- Patient has established tolerability with oral risperidone

OR

2.2 Patient is unable to take oral solid alternatives

AND

3 - If the patient is less than 18 years of age, the prescriber attests they are aware of FDA (Food and Drug Administration) labeling regarding use of long acting injectable antipsychotic products in patients less than 18 years of age and feels the treatment with the requested product is medically necessary (Document rationale for use)

Product Name: Brand Abilify tabs, Brand Clozaril tabs, Brand Geodon, Brand Haldol decanoate inj, Brand Lithobid, Brand Orap, Brand Risperdal (tabs and oral soln), Brand Seroquel, Brand Zyprexa, Brand Zyprexa Zydis, perphenazine-amitriptyline, molindone, aripiprazole ODT, aripiprazole oral solution, Brand Invega, generic paliperidone, Fanapt, Brand Seroquel XR, generic quetiapine ER, Rexulti, Saphris, Secuado, Brand Symbyax, generic fluoxetine-olanzapine, Versacloz, Vraylar, Zyprexa Relprevv

| Diagnosis | Non-Preferred Drugs |
|-----------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - One of the following:

1.1 All of the following:

1.1.1 ONE of the following:

1.1.1.1 Patient has a history of failure, contraindication or intolerance to at least THREE preferred alternatives

- Aripiprazole (Abilify Maintena)
- Aripiprazole (Aristada Initio)
- Aripiprazole (Aristada)
- Aripiprazole (Generic Abilify)
- Clozapine (Generic Clozaril)

- Clozapine ODT (Generic Fazaclo ODT)
- Lurasidone (Latuda)
- Olanzapine (Generic Zyprexa)
- Olanzapine ODT (Generic Zyprexa Zydis)
- paliperidone (Invega Sustenna)
- paliperidone (Invega Trinza)
- Quetiapine (Generic Seroquel)
- Risperidone (Generic Risperdal)
- Risperidone (Risperdal Consta)
- Risperidone ODT (Generic Risperdal ODT)
- Ziprasidone (Generic Geodon)

OR

1.1.1.2 There are no preferred formulary alternatives for the requested drug

AND

1.1.2 If the request is for a multi-source brand medication (i.e., MSC O), ONE of the following:

1.1.2.1 BOTH of the following:

- The brand is being requested because of an adverse reaction, allergy or sensitivity to the generic and the prescriber must attest to submitting the FDA MedWatch Form for allergic reactions to the medications
- If there are generic product(s), the member has tried at least three (if available)

OR

1.1.2.2 ONE of the following:

- The brand is being requested due to a therapeutic failure with the generic (please provide reason for therapeutic failure)
- The brand is being requested because transition to the generic could result in destabilization of the patient (rationale must be provided)
- Special clinical circumstances exist that preclude the use of the generic equivalent of the multi-source brand medication for the patient (rationale must be provided)

1.1.3 ONE of the following:

1.1.3.1 The requested drug must be used for an FDA (Food and Drug Administration)-approved indication

OR

1.1.3.2 The use of this drug is supported by information from ONE of the following appropriate compendia of current literature:

- Food and Drug Administration (FDA) approved indications and limits
- Published practice guidelines and treatment protocols
- Comparative data evaluating the efficacy, type and frequency of side effects and potential drug interactions among alternative products as well as the risks, benefits and potential member outcomes
- Drug Facts and Comparisons
- American Hospital Formulary Service Drug Information
- United States Pharmacopeia Drug Information
- DRUGDEX Information System
- UpToDate
- MicroMedex
- Peer-reviewed medical literature, including randomized clinical trials, outcomes, research data and pharmacoeconomic studies
- Other drug reference resources

AND

1.1.4 ONE of the following:

1.1.4.1 The drug is being prescribed within the manufacturer's published dosing guidelines

OR

1.1.4.2 The drug falls within dosing guidelines found in ONE of the following compendia of current literature:

- Food and Drug Administration (FDA) approved indications and limits
- Published practice guidelines and treatment protocols
- Comparative data evaluating the efficacy, type and frequency of side effects and potential drug interactions among alternative products as well as the risks, benefits and potential member outcomes
- Drug Facts and Comparisons

- American Hospital Formulary Service Drug Information
- United States Pharmacopeia Drug Information
- DRUGDEX Information System
- UpToDate
- MicroMedex
- Peer-reviewed medical literature, including randomized clinical trials, outcomes, research data and pharmacoeconomic studies
- Other drug reference resources

1.1.5 The drug is being prescribed for a medically accepted indication that is recognized as a covered benefit by the applicable health plans' program*

OR

1.2 The requested medication is a behavioral health medication and ONE of the following:

1.2.1 The patient has been receiving treatment with the requested non-preferred behavioral health medication and is new to the plan (enrollment effective date within the past 90 days)

OR

1.2.2 The patient is currently receiving treatment with the requested non-preferred behavioral health medication in the hospital and must continue upon discharge

| al dysfunction purposes are NOT medically accepted indications and | Notes | *Note: Medications used solely for anti-obesity/weight loss, cosmetic (e.g., alopecia, actinic keratosis, vitiligo), erectile dysfunction, and sexu al dysfunction purposes are NOT medically accepted indications and are NOT recognized as a covered benefit. Erectile dysfunction drugs (Ciplic/Tadalafit) are covered for clinical diagnoses other than ED |
|--|-------|--|
|--|-------|--|

| Product Name: Abilify Mycite | |
|------------------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - One of the following:

1.1 All of the following:

- **1.1.1** Patient has ONE of the following:
 - Schizophrenia or schizoaffective disorder
 - Bipolar disorder
 - Autism
 - Major depressive disorder
 - Tourette's

AND

1.1.2 Submission of medical records or claims history documenting the patient is currently prescribed aripiprazole and tolerates the medication

AND

1.1.3 Submission of medical records or claims history documenting the patient's adherence to aripiprazole is less than 80 percent within the past 6 months (medication adherence percentage is defined as the number of pills absent in a given time period divided by the number of pills prescribed during that same time, multiplied by 100)

AND

1.1.4 ALL of the following strategies (if applicable to the patient) to improve patient adherence have been tried without success:

- Utilization of a pill box
- Utilization of a smart phone reminder (ex. alarm, application, or text reminder)
- Involving family members or friends to assist
- Coordinating timing of dose to coincide with dosing of another daily medication

AND

1.1.5 Submission of medical records or claims history documenting patient has experienced life-threatening or potentially life-threatening symptoms, or has experienced a severe worsening of symptoms leading to a hospitalization which was attributed to the lack of adherence to aripiprazole

1.1.6 Prescriber acknowledges that Abilify MyCite has not been shown to improve patient adherence and attests that Abilify MyCite is medically necessary for the patient to maintain compliance, avoid life-threatening worsening of symptoms, and reduce healthcare resources utilized due to lack of adherence

AND

1.1.7 Prescriber agrees to track and document adherence of Abilify MyCite through software provided by the manufacturer

AND

1.1.8 The patient has a history of failure, contraindication, or intolerance or reason or special circumstance they cannot use TWO of the following: (Drug may require PA)

- Abilify Maintena
- Invega Sustenna
- Risperdal Consta
- Aristada
- Perseris

OR

1.2 ONE of the following:

1.2.1 The patient has been receiving treatment with the requested non-preferred behavioral health medication and is new to the plan (enrollment effective date within the past 90 days)

OR

1.2.2 The patient is currently receiving treatment with the requested non-preferred behavioral health medication in the hospital and must continue upon discharge

Product Name: Abilify Mycite

| Approval Length | 12 month(s) | | |
|---|---------------------|--|--|
| Therapy Stage | Reauthorization | | |
| Guideline Type | Prior Authorization | | |
| | | | |
| Approval Criteria | Approval Criteria | | |
| 1 - Documentation that patient is clinically stable on Abilify MyCite | | | |
| | | | |
| | AND | | |
| 2 - Submission of medical records or claims history documenting that the use of Abilify MyCite has increased adherence to 80 percent or more | | | |
| AND | | | |
| 3 - Prescriber attests that the patient requires the continued use of Abilify MyCite to remain adherent | | | |

| Product Name: Lybalvi | |
|-----------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - One of the following:

1.1 All of the following:

1.1.1 Paid claims or submission of medical records (e.g., chart notes) (document drug, duration, dose and date of use) confirming BOTH of the following:

1.1.1.1 Patient has a history of failure, contraindication or intolerance to at least FOUR preferred alternatives:

- Aripiprazole (Abilify Maintena)
- Aripiprazole (Aristada Initio)
- Aripiprazole (Aristada)
- Aripiprazole (Generic Abilify)
- Clozapine (Generic Clozaril)

- Clozapine ODT (Generic Fazaclo ODT)
- Lurasidone (Latuda)
- Paliperidone (Invega Sustenna)
- Paliperidone (Invega Trinza)
- Quetiapine (Generic Seroquel)
- Risperidone (Generic Risperdal)
- Risperidone (Risperdal Consta)
- Risperidone ODT (Generic Risperdal ODT)
- Ziprasidone (Generic Geodon)

1.1.1.2 Failure to respond to generic olanzapine (Generic Zyprexa) given at maximum dosage

AND

1.1.2 ONE of the following:

1.1.2.1 The requested drug must be used for an FDA (Food and Drug Administration)-approved indication

OR

1.1.2.2 The use of this drug is supported by information from ONE of the following appropriate compendia of current literature:

- Food and Drug Administration (FDA) approved indications and limits
- Published practice guidelines and treatment protocols
- Comparative data evaluating the efficacy, type and frequency of side effects and potential drug interactions among alternative products as well as the risks, benefits and potential member outcomes
- Drug Facts and Comparisons
- American Hospital Formulary Service Drug Information
- United States Pharmacopeia Drug Information
- DRUGDEX Information System
- UpToDate
- MicroMedex
- Peer-reviewed medical literature, including randomized clinical trials, outcomes, research data and pharmacoeconomic studies
- Other drug reference resources

1.1.3 ONE of the following:

1.1.3.1 The drug is being prescribed within the manufacturer's published dosing guidelines

OR

1.1.3.2 The drug falls within dosing guidelines found in ONE of the following compendia of current literature:

- Food and Drug Administration (FDA) approved indications and limits
- Published practice guidelines and treatment protocols
- Comparative data evaluating the efficacy, type and frequency of side effects and potential drug interactions among alternative products as well as the risks, benefits and potential member outcomes
- Drug Facts and Comparisons
- American Hospital Formulary Service Drug Information
- United States Pharmacopeia Drug Information
- DRUGDEX Information System
- UpToDate
- MicroMedex
- Peer-reviewed medical literature, including randomized clinical trials, outcomes, research data and pharmacoeconomic studies
- Other drug reference resources

AND

1.1.4 The drug is being prescribed for a medically accepted indication that is recognized as a covered benefit by the applicable health plans' program*

OR

1.2 The requested medication is a behavioral health medication and ONE of the following:

1.2.1 The patient has been receiving treatment with the requested non-preferred behavioral health medication and is new to the plan (enrollment effective date within the past 90 days)

| 1.2.2 The patient is currently receiving treatment with the requested non-preferred behavioral health medication in the hospital and must continue upon discharge | |
|--|---|
| Notes | *Note: Medications used solely for anti-obesity/weight loss, cosmetic (e.g., alopecia, actinic keratosis, vitiligo), erectile dysfunction, and sexu al dysfunction purposes are NOT medically accepted indications and are NOT recognized as a covered benefit. Erectile dysfunction drugs (Cialis/Tadalafil) are covered for clinical diagnoses other than ED. |

| Product Name: generic haloperidol decanoate inj, fluphenazine decanoate, generic clozapine (tabs and ODT tabs) | |
|--|---------------------------|
| Diagnosis | Patients <18 years of age |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - ONE of the following:

1.1 BOTH of the following:

1.1.1 ONE of the following:

1.1.1.1 The requested medication must be used for an FDA (Food and Drug Administration) approved indication

OR

1.1.1.2 The use of the drug is supported by information in ONE of the following appropriate compendia of literature:

- Food and Drug Administration (FDA) approved indications and limits
- Published practice guidelines and treatment protocols
- Comparative data evaluating the efficacy, type and frequency of side effects and potential drug interactions among alternative products as well as the risks, benefits and potential member outcomes
- Drug Facts and Comparisons
- American Hospital Formulary Service Drug Information
- United States Pharmacopeia Drug Information
- DRUGDEX Information System
- UpToDate
- MicroMedex

- Peer-reviewed medical literature, including randomized clinical trials, outcomes, research data and pharmacoeconomic studies
- Other drug reference resources

1.1.2 The patient meets the FDA minimum age limit or the prescriber attests they are aware of FDA labeling regarding the use of the antipsychotic medication and feels the treatment with the requested medication is medically necessary (Document rationale for use)

OR

1.2 The patient is currently on the requested medication

| Date | Notes |
|-----------|--|
| 10/3/2022 | Removed NP criteria for quetiapine 150mg, now preferred. |

Anxiolytics- AZM

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-105536 Anxiolytics- AZM

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 4/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

Product Name: buspirone, Brand Xanax tabs, generic alprazolam tabs, alprazolam ODT, alprazolam conc, Brand Xanax XR, generic alprazolam ER, chlordiazepoxide, Brand Tranxene T, generic clorazepate dipotassium, Brand Valium tabs, generic diazepam tabs, diazepam conc, diazepam oral soln, Brand Ativan, Loreev XR, generic lorazepam, lorazepam conc, generic oxazepam, Brand Klonopin tabs, generic clonazepam tabs, clonazepam ODT

| Diagnosis | Requests for Patients less than 6 years of age |
|-------------------|--|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - The patient is unresponsive to other treatment modalities, unless contraindicated (i.e. other medications or behavioral modification attempted).

AND

2 - The physician attests that the requested medication is medically necessary (Document rationale for use)

| Product Name: Loreev XR | |
|-------------------------|--|
| Diagnosis | Requests for Patients 6 years of age and older |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Trial and failure, or contraindication to generic lorazepam

AND

2 - The physician attests that the requested medication is medically necessary (Document rationale for use)

Product Name: buspirone, Brand Xanax tabs, generic alprazolam tabs, alprazolam ODT, alprazolam conc, Brand Xanax XR, generic alprazolam ER, chlordiazepoxide, Brand Tranxene T, generic clorazepate dipotassium, Brand Valium tabs, generic diazepam tabs, diazepam conc, diazepam oral soln, Brand Ativan, Loreev XR, generic lorazepam, lorazepam conc, generic oxazepam, Brand Klonopin tabs, generic clonazepam tabs, clonazepam ODT

| Reject 88: Drug Utilization Review: Greater than 1 Anxiolytic in 30 days |
|--|
| 12 month(s) |
| Prior Authorization |
| |

Approval Criteria

1 - The medication is being used to adjust the dose of the drug

OR

2 - The medication will be used in place of the previously prescribed drug, and not in addition to it

OR

3 - The medication dosage form will be used in place of the previously prescribed medication dosage form, and not in addition to it

OR

4 - The physician attests they are aware of the multiple anxiolytics prescribed to the patient and feels treatment with both medications is medically necessary (Document rationale for use)

| Date | Notes |
|-----------|------------------------------|
| 3/31/2022 | Added criteria for Loreev XR |

Apidra and Apidra Solostar

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99571 | Apidra and Apidra Solostar |
|----------|----------------------------|
|----------|----------------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Apidra and Apidra Solostar | | | |
|--|--|--|--|
| Guideline Type | Prior Authorization | | |
| | | | |
| Approval Criteria | | | |
| | 1 - Requests for Apidra, and Apidra Solostar should be denied. The plan's preferred products | | |
| Generic Novolo | are. • Generic Novolog • Generic Humalog • Novolog | | |

| Date | Notes |
|-----------|------------|
| 7/13/2021 | New policy |

Apomorphine products (Apokyn, Kynmobi)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-107440 Apomorphine products (Apokyn, Kynmobi)

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 6/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Brand Apokyn, generic apomorphine injection, Kynmobi | |
|--|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting all of the following:

1.1 Diagnosis of Parkinson's disease

1.2 Medication will be used as intermittent treatment for OFF episodes

AND

1.3 Patient is currently on a stable dose of a carbidopa/levodopa-containing medication and will continue receiving treatment with a carbidopa/levodopa-containing medication while on therapy

AND

1.4 Patient continues to experience greater than or equal to 2 hours of OFF time per day despite optimal management of carbidopa/levodopa therapy including BOTH of the following:

- Taking carbidopa/levodopa on an empty stomach or at least one half-hour or more before or one hour after a meal or avoidance of high protein diet
- Dose and dosing interval optimization

AND

1.5 History of failure, contraindication, or intolerance to TWO anti-Parkinson's disease therapies from the following adjunctive pharmacotherapy classes (trial must be from two different classes):

- Dopamine agonists (e.g., pramipexole, ropinirole)
- Catechol-O-methyl transferase (COMT) inhibitors (e.g., entacapone)
- Monoamine oxidase (MAO) B inhibitors (e.g., rasagiline, selegiline)

AND

2 - Prescribed by or in consultation with a neurologist or specialist in the treatment of Parkinson's disease

| Product Name: Brand Apokyn, generic apomorphine injection, Kynmobi | |
|--|-------------|
| Approval Length | 12 month(s) |

| Therapy Stage | Reauthorization | | |
|--|--|--|--|
| Guideline Type | Prior Authorization | | |
| | | | |
| Approval Criteria | Approval Criteria | | |
| 1 - Documentation of positive clinical response to therapy | | | |
| | | | |
| AND | | | |
| | | | |
| 2 - Patient will continue | e to receive treatment with a carbidopa/levodopa-containing medication | | |

| Date | Notes |
|-----------|--|
| 5/24/2022 | Added generic apomorphine injection and Kynmobi as targets |

Aquadeks

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99514 Aquadeks

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Aquadeks | |
|----------------------------------|---------------------|
| Diagnosis | Cystic Fibrosis |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of cystic fibrosis | |

| Date | Notes |
|-----------|--------------------|
| 4/10/2021 | 7/1 Implementation |

Aralast NP - Arizona

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-100647 Aralast NP - Arizona

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|--------------------|-----------|
| P&T Approval Date: | |
| P&T Revision Date: | |

1. Criteria

| Product Name: Aralast NP | | |
|--|---------------------|--|
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Patient has clinically evident emphysema | | |

2 - Patient has a diagnosis of severe congenital deficiency of Alpha1- proteinase inhibitor (alpha1 antitrypsin deficiency)

| Date | Notes |
|------------|---|
| 12/16/2021 | no changes to criteria, formulary name update |

Arcalyst

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-105172 Arcalyst

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 4/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Arcalyst | |
|------------------------|--|
| Diagnosis | Cryopyrin-Associated Periodic Syndromes (CAPS) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting a diagnosis of Cryopyrin-Associated Periodic Syndromes (CAPS) [including Familial Cold Auto-inflammatory Syndrome (FCAS), Muckle-Wells Syndrome (MWS), etc]

| Product Name: Arcalyst | |
|------------------------|--|
| Diagnosis | Cryopyrin-Associated Periodic Syndromes (CAPS) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting positive clinical response to Arcalyst therapy

| Date | Notes |
|-----------|---|
| 3/24/2022 | Updated diagnosis verbiage for clarification. Added Submission of M edical Records. |

Arikayce

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99710 Arikayce

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Arikayce | |
|------------------------|---|
| Diagnosis | Refractory Mycobacterium avium complex (MAC) lung disease |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | · |

Approval Criteria

1 - Diagnosis of refractory Mycobacterium avium complex (MAC) lung disease

2 - Submission of medical records (e.g., chart notes, laboratory values) or claims history documenting respiratory cultures positive for MAC within the previous 6 months

AND

3 - Submission of medical records (e.g., chart notes, laboratory values) or claims history documenting the patient has been receiving a multidrug background regimen containing at least TWO of the following agents for a minimum of 6 consecutive months within the past 12 months (prescription claims history may be used in conjunction as documentation of medication use, dose, and duration):

- Macrolide antibiotic* (e.g., azithromycin, clarithromycin)
- Ethambutol*
- Rifamycin antibiotic* (e.g., rifampin, rifabutin)

AND

4 - Patient will continue to receive a multidrug background regimen

AND

5 - Documentation that the patient has not achieved negative sputum cultures after receipt of a multidrug background regimen for a minimum of 6 consecutive months

AND

6 - In vitro susceptibility testing of recent (within 6 months) positive culture documents that the MAC isolate is susceptible to amikacin with a minimum inhibitory concentration (MIC) of less than or equal to 64 micrograms per milliliter (mcg/mL)

AND

7 - Prescribed by or in consultation with one of the following:

- Infectious disease specialist
- Pulmonologist

| Notes | *Drug may require PA) |
|-------|-----------------------|

| Product Name: Arikayce | |
|------------------------|---|
| Diagnosis | Refractory Mycobacterium avium complex (MAC) lung disease |
| Approval Length | 6 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - ONE of the following:

1.1 Documentation that the patient has achieved negative respiratory cultures

OR

1.2 ALL of the following:

1.2.1 Patient has not achieved negative respiratory cultures while on Arikayce

AND

1.2.2 Physician attestation that patient has demonstrated clinical benefit while on Arikayce

AND

1.2.3 In vitro susceptibility testing of most recent (within 6 months) positive culture with available susceptibility testing documents that the Mycobacterium avium complex (MAC) isolate is susceptible to amikacin with an minimum inhibitory concentration (MIC) of less than 64 micrograms per milliliter (mcg/mL)

| AND | | |
|---|---|--|
| | 1.2.4 Patient has NOT received greater than 12 months of Arikayce therapy with continued positive respiratory cultures | |
| | AND | |
| 2 - Submission of medical records (e.g., chart notes, laboratory values) or claims history documenting that the patient continues to receive a multidrug background regimen containing at least TWO of the following agents (prescription claims history may be used in conjunction as documentation of medication use, dose, and duration): | | |
| Ethambutol* | otic* (e.g., azithromycin, clarithromycin) otic* (e.g., rifampin, rifabutin) | |
| | AND | |
| 3 - Prescribed by or in consultation with one of the following: | | |
| Infectious disease specialistPulmonologist | | |
| Notes | *Drug may require PA | |

| Date | Notes |
|-----------|-------------------------------------|
| 5/12/2021 | Arizona Medicaid 7.1 Implementation |

Austedo (deutetrabenazine)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-114377 Austedo (deutetrabenazine)

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Austedo | |
|-----------------------|---|
| Diagnosis | Chorea Associated with Huntington Disease |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of chorea associated with Huntington's Disease

AND 2 - Prescribed by or in consultation with a neurologist AND 3 - Patient is 18 years of age or older AND 4 - Targeted mutation analysis demonstrates a cytosine-adenine-guanine (CAG) trinucleotide expansion of \geq 36 repeats in the huntingtin (HTT) gene AND 5 - Patient has a Unified Huntington Disease Rating Scale (UHDRS) score ranging from 1 to 4 on any one of UHDRS chorea items 1 through 7 AND 6 - Austedo is not prescribed concurrently with tetrabenazine or Ingrezza AND

7 - Dose does not exceed 48 mg per day

| Product Name: Austedo | |
|-----------------------|---------------------------------------|
| Diagnosis | Moderate to Severe Tardive dyskinesia |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

| Approval Criteria |
|--|
| 1 - Diagnosis of moderate to severe tardive dyskinesia (TD) secondary to treatment with a centrally acting dopamine receptor blocking agent (DRBA) |
| AND |
| 2 - Prescribed by or in consultation with a psychiatrist or neurologist |
| AND |
| 3 - Patient is 18 years of age or older |
| AND |
| 4 - Patient has an Abnormal Involuntary Movement Scale (AIMS) score of 3 or 4 on any one of the AIMS items 1 through 9 |
| AND |
| 5 - Austedo is not prescribed concurrently with tetrabenazine or Ingrezza |
| AND |
| 6 - Dose does not exceed 48 mg per day |

| Product Name: Austedo | |
|-----------------------|---|
| Diagnosis | Chorea Associated with Huntington Disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| | |

Approval Criteria

1 - Patient is responding positively to therapy as evidenced by a reduction in the baseline score of any one of the UHDRS chorea items 1 through 7

AND

2 - Austedo is not prescribed concurrently with tetrabenazine or Ingrezza

AND

3 - Dose does not exceed 48 mg per day

| Product Name: Austedo | |
|-----------------------|---------------------------------------|
| Diagnosis | Moderate to Severe Tardive dyskinesia |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient is responding positively to therapy as evidenced by a reduction in the baseline score of any one of the AIMS items 1 through 9

AND

2 - Austedo is not prescribed concurrently with tetrabenazine or Ingrezza

AND

3 - Dose does not exceed 48 mg per day

| Date | Notes |
|-----------|------------------------------------|
| 9/22/2022 | Removed step through tetrabenazine |

Azole Antifungals

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99585 Azole Antifungals

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Sporanox capsules, generic itraconazole capsules | | |
|--|----------------------------|--|
| Diagnosis | Systemic Fungal Infections | |
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - ONE of the following | | |
| 1.1 Diagnosis of ONE of the following: | | |
| Blastomycosis | | |

- Histoplasmosis
- Aspergillosis

OR

1.2 Both of the following:

1.2.1 Diagnosis of coccidioidomycosis

AND

1.2.2 Patient has a history of failure, contraindication, intolerance, or resistance to fluconazole (generic Diflucan) as evidenced by submission of medical records or claims history

| Product Name: Brand Sporanox capsules, generic itraconazole capsules | |
|--|---------------------------|
| Diagnosis | Onychomycosis Fingernails |
| Approval Length | 2 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of fingernail onychomycosis confirmed by ONE of the following:

- KOH (potassium hydroxide) test
- Fungal culture
- Nail biopsy

AND

2 - Patient has a history of at least a 6-week trial resulting in therapeutic failure, contraindication, intolerance, or resistance to Terbinafine as evidenced by submission of medical records or claims history

Product Name: Brand Sporanox capsules, generic itraconazole capsules

| Diagnosis | Onychomycosis Fingernails |
|-----------------|---------------------------|
| Approval Length | 2 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Both of the following:

1.1 Three months have elapsed since completion of initial therapy for fingernail onychomycosis

AND

1.2 Documentation of positive clinical response to therapy

| Product Name: Brand Sporanox capsules, generic itraconazole capsules | |
|--|------------------------|
| Diagnosis | Onychomycosis Toenails |
| Approval Length | 3 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of toenail onychomycosis confirmed by ONE of the following:

- KOH (potassium hydroxide) test
- Fungal culture
- Nail biopsy

AND

2 - Patient has a history of at least a 12-week trial resulting in therapeutic failure, contraindication, intolerance, or resistance to Terbinafine as evidenced by submission of medical records or claims history.

| Product Name: Brand Sporanox capsules, generic itraconazole capsules | |
|--|------------------------|
| Diagnosis | Onychomycosis Toenails |
| Approval Length | 3 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - BOTH of the following:

1.1 Nine months have elapsed since completion of initial therapy for toenail onychomycosis

AND

1.2 Documentation of positive clinical response to therapy

| Product Name: Brand Sporanox Oral Solution, generic itraconazole oral solution | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - ONE of the following diagnoses: | |
| Oropharyngeal candidiasisEsophageal candidiasis | |

| Product Name: Brand Vfend tablets, generic voriconazole tablets | |
|---|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - One of the following:

1.1 Diagnosis of invasive aspergillosis including Aspergillus fumigatus OR **1.2** ALL of the following: Diagnosis of Candidemia • Patient is non-neutropenic • Patient has a history of failure, contraindication, intolerance, or resistance to • fluconazole (generic Diflucan) as evidenced by submission of medical records or claims history OR **1.3** Both of the following: **1.3.1** ONE of the following diagnoses: Candida infection in the abdomen ٠ Candida infection in the kidney • Candida infection in the bladder wall • Candida infection in wounds • Disseminated Candida infections in skin • Esophageal candidiasis • AND **1.3.2** Patient has a history of failure, contraindication, intolerance, or resistance to fluconazole (generic Diflucan) as evidenced by submission of medical records or claims history OR 1.4 Diagnosis of Scedosporium apiospermum infection (asexual form of Pseudallescheria boydii)

1.5 Diagnosis of Fusarium spp. infection including Fusarium solani

OR

1.6 Diagnosis of Exserohilum species infection

| Product Name: Brand Vfend Powder for Oral Suspension, generoral suspension | ric voriconazole powder for |
|---|-----------------------------|
| Approval Length 12 month(s) | |
| Guideline Type Prior Authorization | |
| | |
| Approval Criteria | |
| 1 - Both of the following: | |
| 1.1 One of the following: | |
| 1.1.1 Diagnosis of invasive aspergillosis including Aspergillus | fumigatus |
| | |
| OR | |
| | |
| 1.1.2 ALL of the following: | |
| Diagnosis of Candidemia | |
| Patient is non-neutropenic Patient has a history of failure, contraindication, intolerar | oce or resistance to |
| fluconazole (generic Diflucan) as evidenced by submissi | |
| claims history | |
| OR | |
| | |
| 1.1.3 Both of the following: | |
| 1.1.3.1 ONE of the following diagnoses: | |
| Candida infection in the abdomen | |
| Candida infection in the kidneyCandida infection in the bladder wall | |

• Candida infection in wounds

- Disseminated Candida infections in skin
- Esophageal candidiasis

AND

1.1.3.2 Patient has a history of failure, contraindication, intolerance, or resistance to fluconazole (generic Diflucan) as evidenced by submission of medical records or claims history

OR

1.1.4 Diagnososis of Scedosporium apiospermum infection (asexual form of Pseudallescheria boydii)

OR

1.1.5 Diagnosis of Fusarium spp. infection including Fusarium solani

OR

1.1.6 Diagnosis of Exserohilum species infection

AND

1.2 Physician has provided rationale for the patient needing to use voriconazole oral suspension instead of voriconazole tablets.

| Product Name: Brand Noxafil tablets, generic posaconazole tablets | |
|---|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - BOTH of the following: | |

1.1 Used as prophylaxis of invasive fungal infections caused by ONE of the following:

- Aspergillus
- Candida

AND

1.2 One of the following conditions:

1.2.1 Patient is at high risk of infections due to severe immunosuppression from ONE of the following conditions:

- Hematopoietic stem cell transplant (HSCT) with graft-versus-host disease (GVHD)
- Hematologic malignancies with prolonged neutropenia from chemotherapy [eg, acute myeloid leukemia (AML), myelodysplastic syndromes (MDS)]

OR

1.2.2 Patient has a prior fungal infection requiring secondary prophylaxis

| Product Name: Noxafil Suspension | |
|----------------------------------|--|
| Diagnosis | Prophylaxis of Aspergillus or Candida Infections |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - BOTH of the following:

1.1 Used as prophylaxis of invasive fungal infections caused by ONE of the following:

- Aspergillus
- Candida

AND

1.2 One of the following conditions:

1.2.1 Patient is at high risk of infections due to severe immunosuppression from ONE of the following conditions:

- Hematopoietic stem cell transplant (HSCT) with graft-versus-host disease (GVHD)
- Hematologic malignancies with prolonged neutropenia from chemotherapy [eg, acute myeloid leukemia (AML), myelodysplastic syndromes (MDS)]

OR

1.2.2 Patient has a prior fungal infection requiring secondary prophylaxis

| Product Name: Noxafil | Suspension | |
|--|--|--|
| Diagnosis | Oropharyngeal Candidiasis (OPC) | |
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - BOTH of the following: | | |
| 1.1 Diagnosis of oropharyngeal candidiasis (OPC) | | |
| | AND | |
| | history of failure, contraindication, intolerance, or resistance to TWO of ced by submission of medical records or claims history: | |
| Fluconazole* (g Itraconazole* (g Clotrimazole Lo | eneric Sporanox) | |
| | | |

| Product Name: Cresemba | |
|------------------------|---------------------|
| Approval Length | 3 month(s) |
| Guideline Type | Prior Authorization |

*Drug may require PA

Notes

| Approval Criteria | | |
|--|----------------------|--|
| 1 - One of the following | : | |
| 1.1 Both of the following | ng: | |
| 1.1.1 Diagnosis of inv | asive aspergillosis | |
| | AND | |
| 1.1.2 Patient has a history of failure, contraindication, intolerance, or resistance to voriconazole* (generic Vfend) as evidenced by submission of medical records or claims history | | |
| | OR | |
| 1.2 Diagnosis of invas | ive mucormycosis | |
| Notes | *Drug may require PA | |

| Product Name: Tolsura | |
|-----------------------|---------------------|
| Approval Length | 3 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

- **1** Both of the following:
- **1.1** Diagnosis of ONE of the following fungal infections:
 - Blastomycosis
 - Histoplasmosis
 - Aspergillosis

AND

1.2 Patient has a history of failure, contraindication, intolerance, or resistance to

| itraconazole* capsules (generic Sporanox) as evidenced by submission of medical records or |
|--|
| claims history |
| |

| Notes | *Drug may require PA |
|-------|----------------------|
|-------|----------------------|

Product Name: Brand Sporanox capsules, generic itraconazole capsules, Brand Sporanox oral solution, generic itraconazole oral solution, Brand Vfend tablets, generic voriconazole tablets, Brand Vfend powder for oral suspension, generic voriconazole powder for oral suspension, Brand Noxafil tablets, generic posaconazole tablets, Noxafil oral suspension, Cresemba, Tolsura

| Diagnosis | All Other Diagnoses |
|----------------|---------------------|
| Guideline Type | Prior Authorization |

Approval Criteria

1 - The use of this drug is supported by information from ONE of the following appropriate compendia of current literature:

- Food and Drug Administration (FDA) approved indications and limits
- Published practice guidelines and treatment protocols
- Comparative data evaluating the efficacy, type and frequency of side effects and potential drug interactions among alternative products as well as the risks, benefits and potential member outcomes
- Drug Facts and Comparisons
- American Hospital Formulary Service Drug Information
- United States Pharmacopeia Drug Information
- DRUGDEX Information System
- UpToDate
- MicroMedex
- Peer-reviewed medical literature, including randomized clinical trials, outcomes, research data and pharmacoeconomic studies
- Other drug reference resources

AND

2 - The medication is being prescribed by or in consultation with an infectious disease specialist.

| Notes | *Authorization duration based on provider recommended treatment du |
|-------|--|
| | rations, not to exceed 12 months |

| Date | Notes |
|-----------|--|
| 9/30/2021 | UM criteria update per SN TSK003786763 eff 10.15.2021 |

Baxdela

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99516 Baxdela

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Baxdela | |
|---|--|
| Diagnosis | Community-Acquired Bacterial Pneumonia |
| Approval Length | 10 Days* |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - For continuation of therapy upon hospital discharge | |

OR

2 - As continuation of therapy when transitioning from intravenous antibiotics that are shown to be sensitive to the cultured organism for the requested indication

OR

3 - All of the following:

3.1 Diagnosis of community-acquired bacterial pneumonia (CABP)

AND

3.2 Infection caused by an organism that is confirmed to be or likely to be susceptible to treatment with Baxdela

AND

3.3 History of failure, contraindication, or intolerance to THREE of the following antibiotics or antibiotic regimens:

- Amoxicillin**
- A macrolide**
- Doxycycline**
- A fluoroquinolone**
- Combination therapy with amoxicillin/clavulanate or cephalosporin AND a macrolide or doxycycline

| Notes | *Note: Authorization will be issued for up to 10 days. **Drug may requi |
|-------|---|
| | re PA |

| Product Name: Baxdela | |
|-----------------------|--|
| Diagnosis | Acute Bacterial Skin and Skin Structure Infections |
| Approval Length | 14 Days* |
| Guideline Type | Prior Authorization |

Approval Criteria 1 - For continuation of therapy upon hospital discharge OR **2** - As continuation of therapy when transitioning from intravenous antibiotics that are shown to be sensitive to the cultured organism for the requested indication OR **3** - All of the following: 3.1 One of the following diagnoses: **3.1.1** Both of the following **3.1.1.1** Acute bacterial skin and skin structure infections AND 3.1.1.2 Infection caused by methicillin-resistant Staphylococcus aureus (MRSA) documented by culture and sensitivity report OR **3.1.2** Both of the following: 3.1.2.1 Empirical treatment of patients with acute bacterial skin and skin structure infections AND 3.1.2.2 Presence of MRSA infection is likely AND

3.2 History of failure, contraindication, or intolerance to linezolid (generic Zyvox) AND **3.3** History of failure, contraindication, or intolerance to ONE of the following antibiotics: Sulfamethoxazole-trimethoprim (SMZ-TMP)** ٠ A tetracycline** • Clindamycin** • OR 4 - All of the following: 4.1 Diagnosis of acute bacterial skin and skin structure infections AND **4.2** Infection caused by an organism that is confirmed to be or likely to be susceptible to treatment with Baxdela AND **4.3** History of failure, contraindication, or intolerance to THREE of the following antibiotics: A penicillin** • A cephalosporin** • A tetracycline** • Sulfamethoxazole-trimethoprim (SMZ-TMP)** Clindamycin** • *Note: Authorization will be issued for up to 14 days. **Drug may requi Notes

| Product Name: Baxdela | |
|-----------------------|---------------------|
| Diagnosis | Off-Label Uses* |
| Guideline Type | Prior Authorization |

re PA

Approval Criteria

1 - For continuation of therapy upon hospital discharge

OR

2 - As continuation of therapy when transitioning from intravenous antibiotics that are shown to be sensitive to the cultured organism for the requested indication

| Notes | *Note: Authorization duration based on provider recommended treatm |
|-------|--|
| | ent durations, up to 6 months. |

| Date | Notes |
|-----------|-------------------------------------|
| 5/12/2021 | Arizona Medicaid 7.1 Implementation |

Belbuca, Butrans - AZM

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-112074 Belbuca, Butrans - AZM

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 8/22/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Belbuca, Brand Butrans, generic buprenorphine patches * | |
|---|--|
| Cancer/Hospice/End of Life related pain | |
| 12 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - The patient is being treated for cancer, hospice, or end of life related pain

2 - If the request is for Belbuca or generic Butrans BOTH of the following:

2.1 Prescriber attests the information provided is true and accurate to the best of their knowledge and they understand that a routine audit may be performed; and medical information necessary to verify the accuracy of the information provided may be requested

AND

2.2 The patient has a history of failure, contraindication or intolerance to BRAND Butrans

| Notes | * If the member is currently taking the requested long-acting opioid for at least 30 days and does not meet the medical necessity authorizati on criteria requirements for treatment with an opioid, a denial should b e issued and a maximum 60-day authorization may be authorized one time for the requested drug/strength combination up to the requested quantity for transition to an alternative treatment. If the member is curr ently taking the requested long-acting opioid for at least 30 days and h as met the medical necessity authorization criteria requirements for tr eatment with an opioid, but has not tried brand buprenorphine patches a denial should be issued and a maximum 60-day authorization may be authorized one time for the requested drug/strength combination u p to the requested quantity for transition to an alternative treatment. A dditionally, a 12 month authorization should be entered for brand bupr |
|-------|--|
| | dditionally, a 12 month authorization should be entered for brand bupr enorphine patches. |

| Product Name: Brand Belbuca, Brand Butrans, generic buprenorphine patches | |
|---|---|
| Diagnosis | Cancer/Hospice/End of Life related pain |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - The patient is being treated for cancer, hospice, or end of life related pain (Document diagnosis and date of diagnosis)

2 - If the request is for Belbuca or generic Butrans ONLY: Prescriber attests the information provided is true and accurate to the best of their knowledge and they understand that a routine audit may be performed; and medical information necessary to verify the accuracy of the information provided may be requested

| Product Name: Brand Belbuca, Brand Butrans, generic buprenorphine patches * | |
|---|---|
| Diagnosis | Non-cancer pain/Non-hospice/Non-end of life care pain |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Prescriber attests to ALL of the following:

1.1 The information provided is true and accurate to the best of their knowledge and they understand that a routine audit may be performed; and medical information necessary to verify the accuracy of the information provided may be requested

AND

1.2 Treatment goals are defined, including estimated duration of treatment

AND

1.3 Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic intervention

AND

1.4 Patient has been screened for substance abuse/opioid dependence

1.5 If used in patients with medical comorbidities or if used concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression

AND

1.6 Pain is moderate to severe and expected to persist for an extended period of time

AND

1.7 Pain is chronic

AND

1.8 Pain is not postoperative (unless the patient is already receiving chronic opioid therapy prior to surgery, or if the postoperative pain is expected to be moderate to severe and persist for an extended period of time)

AND

1.9 Pain management is required around the clock with a long-acting opioid

AND

2 - The patient has a history of failure, contraindication, or intolerance to a trial of tramadol IR (immediate release), unless the patient is already receiving chronic opioid therapy prior to surgery for postoperative pain, or if the postoperative pain is expected to be moderate to severe and persist for an extended period of time (Drug may require PA)

AND

3 - If the request is for neuropathic pain (examples of neuropathic pain include neuralgias, neuropathies, fibromyalgia), BOTH of the following must be met:

3.1 Unless it is contraindicated, the patient has not exhibited an adequate response to 8 weeks of treatment with gabapentin titrated to a therapeutic dose (document date of trial)

AND

3.2 Unless it is contraindicated, the patient has not exhibited an adequate response to at least 6 weeks of treatment with a tricyclic antidepressant titrated to the maximum tolerated dose (document drug and date of trial)

AND

4 - If the request is for Belbuca or generic Butrans, the patient has a history of failure, contraindication or intolerance to BRAND Butrans

| at least 30 days and does not meet the medical necessity authoriz on criteria requirements for treatment with an opioid, a denial shoul e issued and a maximum 60-day authorization may be authorized of time for the requested drug/strength combination up to the request quantity for transition to an alternative treatment. If the member is of ently taking the requested long-acting opioid for at least 30 days ar as met the medical necessity authorization criteria requirements for eatment with an opioid, but has not tried brand buprenorphine patc a denial should be issued and a maximum 60-day authorization m be authorized one time for the requested drug/strength combination p to the requested quantity for transition to an alternative treatment | Notes | * If the member is currently taking the requested long-acting opioid for at least 30 days and does not meet the medical necessity authorizati on criteria requirements for treatment with an opioid, a denial should b e issued and a maximum 60-day authorization may be authorized one time for the requested drug/strength combination up to the requested quantity for transition to an alternative treatment. If the member is curr ently taking the requested long-acting opioid for at least 30 days and h as met the medical necessity authorization criteria requirements for tr eatment with an opioid, but has not tried brand buprenorphine patches a denial should be issued and a maximum 60-day authorization may be authorized one time for the requested drug/strength combination u p to the requested quantity for transition to an alternative treatment. A dditionally, a 6 month authorization should be entered for brand bupre norphine patches. |
|---|-------|---|
|---|-------|---|

| Product Name: Brand Belbuca, Brand Butrans, generic buprenorphine patches * | |
|---|---|
| Diagnosis | Non-cancer pain/Non-hospice/Non-end of life care pain |
| Approval Length | 6 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Patient demonstrates meaningful improvement in pain and function (document improvement in function or pain score improvement) AND 2 - Identify rationale for not tapering and discontinuing opioid (document rationale) AND 3 - Prescriber attests to ALL of the following: **3.1** The information provided is true and accurate to the best of their knowledge and they understand that a routine audit may be performed; and medical information necessary to verify the accuracy of the information provided may be requested AND 3.2 Treatment goals are defined, including estimated duration of treatment AND 3.3 Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic intervention AND 3.4 Patient has been screened for substance abuse/opioid dependence AND 3.5 If used in patients with medical comorbidities or if used concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression

3.6 Pain is moderate to severe and expected to persist for an extended period of time

AND

3.7 Pain is chronic

AND

3.8 Pain is not postoperative (unless the patient is already receiving chronic opioid therapy prior to surgery, or if the postoperative pain is expected to be moderate to severe and persist for an extended period of time)

AND

3.9 Pain management is required around the clock with a long-acting opioid

AND

4 - If the request is for Belbuca or generic Butrans, the patient has a history of failure, contraindication, or intolerance to BRAND Butrans

| Notes | * If the member is currently taking the requested long-acting opioid for at least 30 days and does not meet the medical necessity authorizati on criteria requirements for treatment with an opioid, a denial should b e issued and a maximum 60-day authorization may be authorized one time for the requested drug/strength combination up to the requested quantity for transition to an alternative treatment. If the member is curr ently taking the requested long-acting opioid for at least 30 days and h as met the medical necessity authorization criteria requirements for tr eatment with an opioid, but has not tried brand buprenorphine patches a denial should be issued and a maximum 60-day authorization may be authorized one time for the requested drug/strength combination u p to the requested quantity for transition to an alternative treatment. A dditionally, a 6 month authorization should be entered for brand bupre norphine patches. |
|-------|---|
|-------|---|

| Product Name: Brand I | Belbuca, Brand Butrans, generic buprenorphine patches * |
|---|--|
| Guideline Type | Quantity Limit |
| | |
| Approval Criteria | |
| 1 - The requested dose cannot be achieved by moving to a higher strength of the product | |
| AND | |
| 2 - The requested dose is within the FDA (Food and Drug Administration) maximum dose per day, where an FDA maximum dose per day exists | |
| Notes | *Approval durations: 12 months for cancer pain/hospice/end of life rel ated pain; 6 months for non-cancer pain/non-hospice/non-end of life r elated pain |

| Date | Notes |
|-----------|--|
| 8/22/2022 | Clerical updates, no clinical criteria changes |

Benlysta (belimumab)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-114489 | Benlysta (belimumab) |
|--------------|---|
| Formulary | Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) |
| Formulary No | te |

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Benlysta IV, Benlysta SQ | | |
|--|------------------------------|--|
| Diagnosis | Systemic Lupus Erythematosus | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |

1 - Diagnosis of systemic lupus erythematosus

2 - Patient is 5 years of age or older

AND

3 - Laboratory testing has documented the presence of autoantibodies [e.g., ANA, Anti-dsDNA, Anti-Sm, Anti-Ro/SSA, Anti-La/SSB]

AND

4 - Patient is currently receiving standard immunosuppressive therapy [e.g., hydroxychloroquine, chloroquine, prednisone, azathioprine, methotrexate]

AND

5 - Patient does NOT have severe active central nervous system lupus

AND

6 - Patient is not receiving Benlysta in combination with a biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab), Kineret (anakinra)]

| Product Name: Benlysta IV, Benlysta SQ | |
|--|------------------------|
| Diagnosis | Active Lupus Nephritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of active lupus nephritis

2 - Patient is 5 years of age or older

AND

3 - Patient is currently receiving standard immunosuppressive therapy for systemic lupus erythematosus [e.g., hydroxychloroquine, chloroquine, prednisone, azathioprine, methotrexate]

AND

4 - Patient does NOT have severe active central nervous system lupus

AND

5 - Patient is not receiving Benlysta in combination with a biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab), Kineret (anakinra)]

| Product Name: Benlysta IV, Benlysta SQ | | |
|--|--|--|
| Diagnosis | Systemic Lupus Erythematosus, Active Lupus Nephritis | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Documentation of positive clinical response to Benlysta therapy

AND

2 - Patient is not receiving Benlysta in combination with a biologic DMARD (disease modifying

anti-rheumatic drug) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab), Kineret (anakinra)]

| Date | Notes |
|-----------|--|
| 9/26/2022 | Updated age requirement. Added IV formulation as target. |

Benznidazole

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99434 Benznidazole

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Benznidazole | | |
|----------------------------|---|--|
| Diagnosis | Chagas disease (American trypanosomiasis) | |
| Approval Length | 60 Day(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |

1 - Diagnosis of Chagas disease (American trypanosomiasis) due to Trypanosoma cruzi

| Date | Notes |
|-----------|--|
| 3/10/2021 | Bulk Copy guidelines starting with B and C from C&S Arizona to Ariz ona Medicaid |

Berinert

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99661 Berinert

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Berinert | | |
|---------------------------------------|-----------------------|--|
| Diagnosis Hereditary angioedema (HAE) | | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |

Approval Criteria

1 - Diagnosis of hereditary angioedema (HAE) confirmed by ONE of the following:

1.1 C1 inhibitor (C1-INH) deficiency or dysfunction (Type I or II HAE) as documented by ONE of the following (per laboratory standard): C1-INH antigenic level below the lower limit of normal • C1-INH functional level below the lower limit of normal OR 1.2 HAE with normal C1 inhibitor levels and ONE of the following: Confirmed presence of a FXII, angiopoietin-1 or plasminogen gene mutation • Recurring angioedema attacks that are refractory to high-dose antihistamines with • confirmed family history of angioedema AND 2 - Prescribed for the treatment of acute HAE attacks AND **3** - Not used in combination with other approved treatments for acute HAE attacks (e.g. Firazyr, Ruconest) AND 4 - ONE of the following: **4.1** Submission of medical records documenting a history of failure, contraindication, or intolerance to Ruconest (C1 esterase inhibitor [recombinant]) OR 4.2 Patient is currently on Berinert therapy AND

5 - Prescribed by ONE of the following:

5.1 Immunologist

OR

5.2 Allergist

| Product Name: Berinert | | | |
|--|-----------------------------|--|--|
| Diagnosis | Hereditary angioedema (HAE) | | |
| Approval Length | 12 month(s) | | |
| Therapy Stage | Reauthorization | | |
| Guideline Type | Prior Authorization | | |
| Approval Criteria 1 - Documentation of positive clinical response | | | |
| | AND | | |
| 2 - Prescribed for the acute treatment of HAE (hereditary angioedema) attacks | | | |
| AND | | | |
| 3 - Not used in combination with other products indicated for the acute treatment of HAE attacks (e.g. Firazyr, Ruconest) | | | |
| AND | | | |
| 4 - Prescribed by ONE | of the following: | | |
| 4.1 Immunologist | | | |

| OR |
|---------------|
| 4.2 Allergist |

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk copy C&S Arizona Medicaid SP to Medicaid Arizona SP for eff 7 /1 |

Blood Glucose Monitors

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99566 | Blood Glucose Monitors |
|--------------|--|
| Formulary | Medicaid - Arizona (AZM, AZMREF, AZMDDD) |
| Formulary No | ote |

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Non-preferred Blood Glucose Monitors* | | |
|---|--|--|
| Approval Length | 12 month(s) | |
| Guideline Type Prior Authorization | | |
| | | |
| Approval Criteria | | |
| 1 - Patient is visually impaired | | |
| Notes | *Please reference background table for list of Non-preferred Blood Gl ucose Monitors **Approve Glucose Monitor at NDC Level | |

2. Background

| Benefit/Coverage/Program Information | | | |
|---------------------------------------|----------------------------|-----------------------------|--|
| Non-preferred Blood Glucose Monitors* | | | |
| CONTOUR KIT NEXT LNK | EASY TOUCH KIT MONITOR | EASYMAX V KIT SYSTEM | |
| CONTOUR NXT KIT LINK 2.4 | KROGER BGM KIT SYSTEM | EASYMAX NG KIT SYSTEM | |
| CONTOUR KIT NEXT EZ | ELEMENT AUTO KIT SYSTEM | MEIJER BGM KIT ESSENTIA | |
| CONTOUR KIT NEXT | SMARTEST KIT EJECT | MEIJER GLUCO KIT MONITOR | |
| CONTOUR KIT MONITOR | SMARTEST KIT PROTEGE | MEIJER BGM KIT PREMIUM | |
| RELION MICRO KIT | SMARTEST KIT PRONTO | FORA V30A KIT | |
| RELION KIT MONITOR | SMARTEST KIT PERSONA | FORA TN'G KIT VOICE | |
| BD LOGIC KIT MONITOR | GLUCOCOM KIT MONITOR | REFUAH PLUS KIT SYSTEM | |
| BD LATITUDE KIT | RIGHTEST SYS KIT GM300 | KROGER BGM KIT | |
| BD LATITUDE KIT SYSTEM | RIGHTEST SYS KIT GM100 | KROGER BGM KIT PREMIUM | |
| QUICKTEK KIT | RIGHTEST SYS KIT GM550 | CONTOUR KIT LINK 2.4 | |
| ADVANCE KIT INTUITIO | IGLUCOSE KIT | EASYMAX V KIT SYSTEM | |
| GLUCOCARD KIT SHNE CON | NOVA MAX KIT SYSTEM | EASYMAX NG KIT SYSTEM | |

| GLUCOCARD KIT SHNE EXP | WAVESENSE KIT KEYNOTE | MYGLUCOHEALT KIT SYSTEM |
|------------------------------|-----------------------------|----------------------------|
| GLUCOCARD KIT EXPRESSI | AGAMA JAZZ KIT WRLSS 2 | MICRODOT KIT SYSTEM |
| POCKETCHEM KIT EZ | AGAMATRIX KIT PRESTO | ONE TOUCH KIT VERIO FL |
| GLUCOCARD 01 KIT SYSTEM | WAVESENSE KIT AMP | RELION TRUE KIT MET AIR |
| GLUCOCARD 01 KIT MINI | SOLUS V2 KIT SYSTEM | VERASENS KIT |
| GLUCOCARD KIT X- METER | COOL MONITOR KIT | INFINITY KIT VOICE |
| GLUCOCARD KIT VITAL | TRUERESULT KIT MONITOR | OPTIUM KIT BL GLUC |
| RELION PREMI KIT COMP SYS | TRUERESULT KIT SYSTEM | PRECISION KIT XTRA |
| SMART SENSE KIT GLUC SYS | MEIJER BGM KIT ESSENTIA | PRECISION KIT LINK |
| CVS GLUCOSE KIT METER | MEIJER GLUCO KIT MONITOR | BIOTEL CARE KIT SYSTEM |
| INFINITY KIT SYSTEM | MEIJER BGM KIT PREMIUM | BIOTEL CARE KIT |
| EASYPRO KIT MONITOR | FORA V30A KIT | FREESTYLE KIT SIDEKICK |
| EASYPRO PLUS KIT | FORA TN'G KIT VOICE | FREESTYLE KIT FREEDOM |
| PRODIGY PCKT KIT METER | REFUAH PLUS KIT SYSTEM | KROGER BGM KIT PREMIUM |
| PRODIGY AUTO KIT MONITOR | KROGER BGM KIT | CONTOUR KIT LINK 2.4 |

| PRODIGY VOIC KIT METER | | |
|---------------------------|--|--|
| PRODIGY KIT NO CODIN | | |

| Date | Notes |
|-----------|-------------|
| 7/12/2021 | New Program |

Bonjesta and Diclegis

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99436 | Bonjesta and Diclegis |
|-----------|--|
| Formulary | Medicaid - Arizona (AZM, AZMREF, AZMDDD) |

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Bonjesta, Brand Diclegis, generic doxylamine/pyridoxine | |
|---|--|
| | |
| | |
| | |
| | |
| | |

Approval Criteria

1 - Diagnosis of nausea and vomiting associated with pregnancy

AND

2 - Documented failure or contraindication to lifestyle modifications (e.g., diet, avoidance of triggers)

AND

3 - Documented trial and failure or contraindication to a five day trial of over-the-counter doxylamine taken together with pyridoxine (i.e., not a combined dosage form, but separate formulations taken concomitantly)

| Date | Notes |
|-----------|--|
| 3/10/2021 | Bulk Copy guidelines starting with B and C from C&S Arizona to Ariz ona Medicaid |

Brand Over Generic Not Covered

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99590 Brand Over Generic Not Covered

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Generic products on a brand* over generic program | | |
|--|--|--|
| Guideline Type | Administrative | |
| | | |
| Approval Criteria | | |
| 1 - Requests for a generic product on a brand over generic program (presence of Brand over generic-Not Covered clinical program in formulary lookup) shall be denied. The plan's preferred product is the brand medication. | | |
| Notes | * Brand product may require prior authorization. | |

| Date | Notes |
|------------|-----------------------------------|
| 10/29/2021 | Changed effective date to 12/1/21 |

Brand Vascepa, Generic Icosapent Ethyl, Brand Lovaza, generic omega-3-acid ethyl esters

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99572 Brand Vascepa, Generic Icosapent Ethyl, Brand Lovaza, generic omega-3acid ethyl esters

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Vascepa, Generic Icosapent Ethyl, Brand Lovaza, generic omega-3- acid ethyl esters | |
|---|---------------------|
| Guideline Type | Prior Authorization |
| Approval Criteria 1 - Requests for brand Vascepa, generic Icosapent Ethyl, Brand Lovaza, generic omega-3- | |
| acid ethyl esters should be denied. The plan's preferred products are generic over-the-counter omega 3 fatty acids. | |

| Date | Notes |
|-----------|------------|
| 7/14/2021 | New Policy |

Breast Cancer - Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99541 | Breast Cancer - Arizona |
|----------|-------------------------|
|----------|-------------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Arimidex, generic anastrozole | |
|---|---------------------|
| Diagnosis | Breast Cancer |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - ONE of the following:

1.1 Adjuvant treatment of postmenopausal patients with hormone receptor-positive early breast cancer

OR

1.2 First-line treatment of postmenopausal patients with hormone receptor-positive or hormone receptor status unknown locally advanced or metastatic breast cancer

OR

1.3 Postmenopausal patients with disease progression following tamoxifen therapy

| Product Name: Brand Aromasin, generic exemestane | |
|--|---------------------|
| Diagnosis | Breast Cancer |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - ONE of the following:

1.1 Adjuvant treatment of postmenopausal patients with estrogen receptor-positive early breast cancer who have received 2 to 3 years of tamoxifen and are switched to exemestane for completion of a total of 5 consecutive years of adjuvant hormonal therapy

OR

1.2 Treatment of advanced breast cancer in postmenopausal patients whose disease has progressed following tamoxifen therapy

| Product Name: Brand Fareston, generic toremifene | |
|--|---------------------|
| Diagnosis | Breast Cancer |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Treatment of metastatic breast cancer in postmenopausal patients with estrogen receptor positive tumors or with tumors of unknown estrogen receptor status

| Product Name: Brand Arimidex, generic anastrozole, Brand Aromasin, generic exemestane, Brand Fareston, generic toremifene | |
|--|---|
| Diagnosis | National Comprehensive Cancer Network (NCCN) Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Use supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Brand Arimidex, generic anastrozole, Brand Aromasin, generic exemestane, Brand Fareston, generic toremifene | |
|--|---|
| Diagnosis | National Comprehensive Cancer Network (NCCN) Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to therapy

| Date | Notes |
|----------|-------------------------------------|
| 6/3/2021 | Arizona Medicaid 7.1 Implementation |

Breo Ellipta

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99564 Breo Ellipta

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Breo Ellipta | |
|----------------------------|---------------------|
| Diagnosis | Asthma, COPD |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Both of the following: | |
| 1.1 Diagnosis of asthma | |

AND

1.2 The patient has a history of failure, contraindication, or intolerance to treatment with ALL of the following preferred products:

- Advair Diskus (brand) or Advair HFA
- Dulera
- Symbicort

OR

2 - All of the following:

2.1 Diagnosis of chronic obstructive pulmonary disease (COPD)

AND

2.2 One of the following:

2.2.1 History of failure, contraindication, or intolerance to treatment with at least a 30 day trial of an orally inhaled anticholinergic agent (e.g. Spiriva, Atrovent, Combivent, Tudorza)

OR

2.2.2 History of failure, contraindication, or intolerance to treatment with at least a 30 day trial of an orally inhaled anticholinergic agent/long-acting beta-agonist combination agent (e.g. Anoro Ellipta, Stiolto Respimat)

AND

2.3 The patient has a history of failure, contraindication, or intolerance to treatment with ALL of the following preferred products:

- Advair Diskus (brand) or Advair HFA
- Dulera
- Symbicort

Brilinta and Effient

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99561 Brilinta and Effient

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|--------------------|-----------|
| P&T Approval Date: | |
| P&T Revision Date: | |

1. Criteria

| Product Name: Brand Brilinta, Brand Effient, Generic prasurgrel | |
|---|-------------------------------|
| Diagnosis | Acute coronary syndrome (ACS) |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| | |

Approval Criteria

1 - Diagnosis of acute coronary syndrome (ACS) [e.g., unstable angina (UA), non-ST

elevation myocardial infarction (NSTEMI) or ST-segment elevation myocardial infarction (STEMI)]

AND

2 - If request is for Effient (prasugrel), patient must be managed with percutaneous coronary intervention (PCI)

Buphenyl

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99600 Buphenyl

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Buphenyl oral powder, generic sodium phenylbutyrate oral powder | |
|---|-----------------------------|
| Diagnosis | Urea Cycle Disorders (UCDs) |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of urea cycle disorders (UCDs) | |

Product Name: Brand Buphenyl tablets, generic sodium phenylbutyrate tablets

| Diagnosis | Urea Cycle Disorders (UCDs) |
|-----------------|-----------------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of urea cycle disorders (UCDs)

AND

2 - Prescriber provides a reason or special circumstance the patient cannot use Buphenyl (sodium phenylbutyrate) powder for oral solution

| Product Name: Brand Buphenyl tablets, generic sodium phenylbutyrate tablets | | |
|---|-----------------------------|--|
| Diagnosis | Urea Cycle Disorders (UCDs) | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |

1 - Documentation of positive clinical response to Buphenyl (sodium phenylbutyrate) tablets

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona SP to Medicaid Arizona SP for 7/1 eff |

Buprenorphine Sublingual Tablet

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99576 | Buprenorphine | Sublingual Tablet |
|----------|----------------------|-------------------|
| | | - ···· |

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Buprenorphine Sublingual Tablet | | |
|---|---------------------|--|
| Approval Length | 6 Months* | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of opioid abuse/dependence. | | |
| | | |
| AND | | |
| | | |

2 - Prescriber attests they meet the DATA 2000 (Drug Addiction Treatment Act of 2000) requirements and has been assigned a unique identification number specific to the prescription of medication assisted therapy (DEA-X).

AND

3 - One of the following:

3.1 Patient is pregnant or breastfeeding;*

OR

3.2 Both of the following:

3.2.1 Patient had an intolerance or side effect to buprenorphine-naloxone sublingual tablet or film;

AND

3.2.2 Side effects or intolerances to buprenorphine-naloxone sublingual tablet or films were not resolved with a trial of anti-emetics (e.g. ondansetron) or non-opioid analgesics.

OR

3.3 Patient has a contraindication to naloxone.

OR

3.4 Both of the following:

3.4.1 Patient has a severe allergy to naloxone [e.g., Stevens-Johnson syndrome, DRESS (Drug Rash with Eosinophilia and Systemic Symptoms)];

AND

3.4.2 Provider has submitted a copy of the MedWatch Form 3500 to the Food and Drug Administration documenting the adverse reaction

AND

4 - Patient is not currently on ANY of the following:

- Benzodiazepines (e.g. Alprazolam, Diazepam, Lorazepam)
- Hypnotics (e.g. Temazepam, Rozerem, Zolpidem)
- Opioids (e.g. Oxycodone, Tramadol, Hydrocodone)

AND

5 - Prescriber attests that the Arizona State Board of Pharmacy Controlled Substance Prescription Drug Monitoring Program database has been reviewed and that patient has been warned about the dangers of ingesting concurrent sedating medications

| Notes | *Approve for 1 year if pregnant or breastfeeding |
|-------|--|
|-------|--|

Cablivi

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99601 Cablivi

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Cablivi | |
|-----------------------|---|
| Diagnosis | Acquired thrombotic thrombocytopenic purpura (aTTP) |
| Approval Length | 2 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| | |

Approval Criteria

1 - Diagnosis of acquired thrombotic thrombocytopenic purpura (aTTP)

AND

2 - Cablivi was initiated in the inpatient setting in combination with plasma exchange therapy

AND

3 - Cablivi will be used in combination with immunosuppressive therapy (e.g., corticosteroids)

AND

4 - Total treatment duration will be limited to 58 days beyond the last therapeutic plasma exchange

| Product Name: Cablivi | |
|-----------------------|---|
| Diagnosis | Acquired thrombotic thrombocytopenic purpura (aTTP) |
| Approval Length | 2 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Request is for a new (different) episode requiring the re-initiation of plasma exchange for the treatment of acquired thrombotic thrombocytopenic purpura (aTTP) (Documentation of date of prior episode and documentation date of new episode required)

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona SP to Medicaid Arizona SP for 7/1 eff |

Cabotegravir Containing Agents

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-113150 Cabotegravir Containing Agents

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Vocabria*, Cabenuva* | | |
|------------------------------------|------------------------------|--|
| Diagnosis | Treatment of HIV-1 Infection | |
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - All of the following: | | |
| 1.1 Diagnosis of HIV-1 infection | | |

AND

1.2 Patient is 12 years of age or older

AND

1.3 Patient's weight is greater than or equal to 35 kg

AND

1.4 Patient is currently virologically suppressed (HIV-1 RNA less than 50 copies/mL) on a stable, uninterrupted antiretroviral regimen for at least 6 months

AND

1.5 Patient has no history of treatment failure or known/suspected resistance to either cabotegravir or rilpivirine

AND

1.6 Provider attests that patient would benefit from long-acting injectable therapy over standard oral regimens

AND

1.7 Prescribed by or in consultation with a clinician with HIV expertise

OR

2 - For continuation of prior therapy

| Notes | *If patient meets criteria above, please approve both Vocabria and Ca |
|-------|---|
| | benuva at GPI list "CABOTEGRPA". |

| Product Name: Vocabria**, Apretude** | | |
|---|--|--|
| Diagnosis | HIV-1 Pre-Exposure Prophylaxis | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| Approval Criteria | | |
| sexually acquired HIV- | peing used for pre-exposure prophylaxis (PrEP) to reduce the risk of 1 infection | |
| | AND | |
| 2 - Patient's weight is g | reater than or equal to 35 kg | |
| AND | | |
| 3 - Documentation of both of the following U.S. Food and Drug (FDA)-approved test prior to use of Vocabria or Apretude: | | |
| Negative HIV-1 antigen/antibody test Negative HIV-1 RNA assay | | |
| AND | | |
| 4 - One of the following | i: | |
| 4.1 Contraindication or intolerance to generic emtricitabine-tenofovir disoproxil fumarate 200/300mg | | |
| | OR | |
| 4.2 Provider attests to | both of the following: | |
| Patient would benefit from long-acting injectable therapy over standard oral regimens Patient would be adherent to testing and dosing schedule | | |

| Notes | **If patient meets criteria above, please approve both Vocabria and A pretude at GPI list "APRETUDEPA" |
|-------|--|
| | |

| Product Name: Vocabria**, Apretude** | |
|--------------------------------------|--------------------------------|
| Diagnosis | HIV-1 Pre-Exposure Prophylaxis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Provider attests that patient is adherent to the testing appointments and scheduled injections of Apretude

AND

2 - Documentation of both of the following U.S. Food and Drug (FDA)-approved test prior to each maintenance injection of Apretude for HIV PrEP:

- Negative HIV-1 antigen/antibody test
- Negative HIV-1 RNA assay

| Notes **If patient meets criteria above, please approve both Vocabria and pretude at GPI list "APRETUDEPA" |
|--|
|--|

| Date | Notes |
|----------|-------------|
| 9/8/2022 | New Program |

Calcium/Vitamin D

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99531 Calcium/Vitamin D

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Approval Length | 12 month(s) | |
|--|--|--|
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Provider has submitted lab work documenting a Vitamin D deficiency | | |
| Notes | Calcium carbonate and calcium lactate are covered without the need f or prior authorization. | |

| Date | Notes |
|-----------|--------------------|
| 5/18/2021 | 7/1 Implementation |

Camzyos (mavacamten)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-114157 | Camzyos (mavacamten) |
|-----------|----------------------|
|-----------|----------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Camzyos | |
|-----------------------|-----------------------|
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of obstructive hypertrophic cardiomyopathy (HCM)

2 - Patient has New York Heart Association (NYHA) Class II or III symptoms (e.g., shortness of breath, chest pain)

AND

3 - Patient has a left ventricular ejection fraction of greater than or equal to 55%

AND

4 - Patient has valsalva left ventricular outflow tract (LVOT) peak gradient greater than or equal to 50 mmHg at rest or with provocation

AND

5 - Trial and failure, contraindication, or intolerance to both of the following at a maximally tolerated dose:

- non-vasodilating beta blocker (e.g., bisoprolol, propranolol)
- calcium channel blocker (e.g., verapamil, diltiazem)

AND

6 - Prescribed by or in consultation with a cardiologist

| Product Name: Camzyos | |
|-----------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Documentation of positive clinical response to therapy (e.g., improved symptom relief)

AND

2 - Patient has a left ventricular ejection fraction of greater than or equal to 50%

AND

3 - Prescribed by or in consultation with a cardiologist

| Date | Notes |
|-----------|-------------|
| 9/20/2022 | New Program |

Caplyta (lumateperone)

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-112751 Caplyta (lumateperone)

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 8/27/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Caplyta | |
|--------------------------------|---------------------|
| Diagnosis | Schizophrenia |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of schizophrenia | |

AND 2 - One of the following: 2.1 History of failure, contraindication or intolerance to at least THREE of the following preferred alternatives: Aripiprazole (e.g., generic Abilify, Abilify Maintena, Aristata) • Clozapine/clozapine ODT ٠ Latuda (lurasidone) • Olanzapine/olanzapine ODT ٠ Invega injectable formulations (e.g., Invega Sustenna, Invega Trinza, Hafyera) • Quetiapine ٠ Risperidone/risperidone ODT/Risperdal Consta (risperidone injection) • Ziprasidone • OR **2.2** One of the following: 2.2.1 The patient has been receiving treatment with Caplyta and is new to the plan (enrollment effective date within the past 90 days)

OR

2.2.2 The patient is currently receiving treatment with Caplyta in the hospital and must continue upon discharge

| Product Name: Caplyta | |
|-----------------------|---------------------|
| Diagnosis | Bipolar Disorder |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of bipolar I or II disorder (bipolar depression)

2 - One of the following:

2.1 History of failure, contraindication or intolerance to at least THREE of the following preferred alternatives:

- Bupropion IR/SR/XL (e.g., generic Wellbutrin IR/SR/XL)
- Citalopram tablets or solution
- Duloxetine 20 mg, 30 mg, or 60 mg
- Escitalopram tablets
- Fluoxetine tablets or solution
- Fluvoxamine tablets
- Mirtazapine
- Paroxetine IR tablets
- Sertraline tablets or oral concentrate for solution
- Trazodone
- Venlafaxine IR tablets or Venlafaxine ER capsules

OR

2.2 One of the following:

2.2.1 The patient has been receiving treatment with Caplyta and is new to the plan (enrollment effective date within the past 90 days)

OR

2.2.2 The patient is currently receiving treatment with Caplyta in the hospital and must continue upon discharge

| Product Name: Caplyta | |
|-----------------------|--|
| Diagnosis | Caplyta Requests Exceeding Quantity Limit* |
| Approval Length | 12 month(s) |
| Guideline Type | Quantity Limit |
| | |
| Approval Criteria | |

1 - ONE of the following:

1.1 The requested drug must be used for a Food and Drug Administration (FDA)-approved indication

OR

1.2 The use of this drug is supported by information from one of the following appropriate compendia of current literature:

- Food and Drug Administration (FDA) approved indications and limits
- Published practice guidelines and treatment protocols
- Comparative data evaluating the efficacy, type and frequency of side effects and potential drug interactions among alternative products as well as the risks, benefits and potential member outcomes
- Drug Facts and Comparisons
- American Hospital Formulary Service Drug Information
- United States Pharmacopeia Drug Information
- DRUGDEX Information System
- UpToDate
- MicroMedex
- Peer-reviewed medical literature, including randomized clinical trials, outcomes, research data and pharmacoeconomic studies
- Other drug reference resources

AND

2 - ONE of the following:

2.1 The drug is being prescribed within the manufacturer's published dosing guidelines

OR

2.2 The requested dose falls within dosing guidelines found in one of the following compendia of current literature:

- Food and Drug Administration (FDA) approved indications and limits
- Published practice guidelines and treatment protocols
- Comparative data evaluating the efficacy, type and frequency of side effects and potential drug interactions among alternative products as well as the risks, benefits and potential member outcomes
- Drug Facts and Comparisons

| United States Ph DRUGDEX Infor UpToDate MicroMedex Peer-reviewed n | nedical literature, including randomized clinical trials, outcomes, nd pharmacoeconomic studies | |
|--|---|--|
| | AND | |
| 3 - The requested dosaged ifferent dose or formula | ge cannot be achieved using the plan accepted quantity limit of a ation | |
| | AND | |
| 0 | escribed for a medically accepted indication that is recognized as a applicable health plans' program | |
| AND | | |
| (42 milligrams [mg]) per | led rationale for needing to exceed the quantity limit of one capsule day (NOTE: The treatment effect of Caplyta 84mg daily versus tically significant in clinical trials.) | |
| | *Caplyta requests should be reviewed using the above Non-Preferred criteria. This section is for Caplyta quantity limit requests only. | |

| Date | Notes |
|-----------|--|
| 8/26/2022 | Removed basic info notes, replaced "NP behavioral health drug" with "Caplyta". |

Caprelsa

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99678 Caprelsa

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Caprelsa | |
|---|--------------------------------|
| Diagnosis | Medullary thyroid cancer (MTC) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of medullary thyroid cancer (MTC) | |

2 - ONE of the following:

- Unresectable locally advanced disease •
- Metastatic disease •

AND

- **3** ONE of the following:
 - Patient has symptomatic disease Patient has progressive disease ٠
 - •

| edullary thyroid cancer (MTC) |
|-------------------------------|
| |
| t month(s) |
| eauthorization |
| ior Authorization |
| ea |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Caprelsa therapy

| Product Name: Caprelsa | |
|------------------------|---|
| Diagnosis | Follicular carcinoma, Hürthle cell carcinoma, Papillary carcinoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - One of the following diagnoses:

- Hürthle cell carcinoma •
- Papillary carcinoma •

2 - One of the following:

- Unresectable recurrent disease •
- Persistent locoregional disease •
- Metastatic disease •

AND

3 - One of the following:

- Patient has symptomatic disease Patient has progressive disease ٠
- •

AND

4 - Disease is refractory to radioactive iodine treatment

| Product Name: Caprelsa | |
|------------------------|---|
| Diagnosis | Follicular carcinoma, Hürthle cell carcinoma, Papillary carcinoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Caprelsa therapy

| Product Name: Caprelsa | |
|------------------------|------------------------------------|
| Diagnosis | Non-Small Cell Lung Cancer (NSCLC) |

| Approval Length | 12 month(s) |
|-----------------|-----------------------|
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of Non-Small Cell Lung Cancer (NSCLC)

AND

2 - Disease is positive for RET gene rearrangement

| Product Name: Caprelsa | |
|------------------------|------------------------------------|
| Diagnosis | Non-Small Cell Lung Cancer (NSCLC) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Caprelsa therapy

| Product Name: Caprelsa | |
|------------------------|---|
| Diagnosis | National Comprehensive Cancer Network (NCCN) Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Use supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Caprelsa | |
|------------------------|---|
| Diagnosis | National Comprehensive Cancer Network (NCCN) Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| Approval Criteria | |

1 - Documentation of positive clinical response to Caprelsa therapy

| Date | Notes |
|----------|--------------------|
| 4/8/2021 | 7/1 Implementation |

Carbaglu (carglumic acid)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-104872 | Carbaglu | (carglumic acid) |
|-----------|----------|------------------|
|-----------|----------|------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 4/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Brand Carbaglu, Generic carglumic acid | |
|--|--|
| Diagnosis | Acute Hyperammonemia due to N-acetylglutamate Synthase (NAGS) Deficiency |
| Approval Length | 3 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of acute hyperammonemia due to N-acetylglutamate synthase (NAGS) deficiency

2 - Medication will be used as adjunctive therapy to other ammonia lowering therapies (e.g., protein restriction, ammonia scavengers, dialysis)

AND

3 - Prescribed by or in consultation with a specialist focused in the treatment of metabolic disorders

| Product Name: Brand Carbaglu, Generic carglumic acid | |
|--|--|
| Diagnosis | Acute Hyperammonemia due to Propionic Acidemia (PA) or Methylmalonic Acidemia (MMA) |
| Approval Length | 1 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of acute hyperammonemia due to propionic acidemia (PA) or methylmalonic acidemia (MMA)

AND

2 - Medication will be used as adjunctive therapy to other ammonia lowering therapies (e.g. intravenous glucose, insulin, protein restriction, dialysis)

AND

3 - Patient's plasma ammonia level is greater than or equal to 50 micromol/L

AND

4 - Medication will be used for a maximum duration of 7 days

AND

5 - Prescribed by or in consultation with a specialist focused in the treatment of metabolic disorders

| Product Name: Brand (| Product Name: Brand Carbaglu, Generic carglumic acid | | |
|--|--|--|--|
| | | | |
| Diagnosis | Chronic Hyperammonemia due to N-acetylglutamate Synthase (NAGS) Deficiency | | |
| Approval Length | 12 month(s) | | |
| Therapy Stage | Initial Authorization | | |
| Guideline Type | Prior Authorization | | |
| | | | |
| Approval Criteria | | | |
| 1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of chronic hyperammonemia due to N-acetylglutamate synthase (NAGS) deficiency | | | |
| AND | | | |
| 2 - NAGS deficiency has been confirmed by genetic/mutational analysis | | | |
| AND | | | |
| 3 - Medication will be used as maintenance therapy | | | |
| AND | | | |
| 4 - Prescribed by or in consultation with a specialist focused in the treatment of metabolic disorders | | | |

Product Name: Brand Carbaglu, Generic carglumic acid

| Diagnosis | Chronic Hyperammonemia due to N-acetylglutamate Synthase (NAGS) Deficiency | |
|-----------------|--|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting a positive clinical response to therapy (e.g., plasma ammonia level within the normal range)

| Date | Notes |
|-----------|--|
| 3/31/2022 | New program for Carbaglu, mirrors ORx LOB. Added submission of MR to each section. |

Cayston

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99603 Cayston

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Cayston | |
|---------------------------------------|----------------------|
| Diagnosis | Cystic Fibrosis (CF) |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of cystic fibrosis (CF) | |

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona SP to Medicaid Arizona SP for 7/1 eff |

CGRP - AZ

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99517 CGRP - AZ

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Ajovy, Emgality 120mg | | |
|-------------------------------------|-----------------------|--|
| Diagnosis | Episodic Migraine | |
| Approval Length | 6 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

- **1** Diagnosis of episodic migraines with BOTH of the following:
 - Less than 15 headache days per month

• Patient has 4 to 14 migraine days per month

AND

2 - Trial and failure (after a trial of at least two months), contraindication, or intolerance to TWO of the following prophylactic therapies from the list below:

- Amitriptyline (Elavil)*
- ONE of the following beta-blockers: atenolol, metoprolol, nadolol, propranolol, or timolol*
- Divalproex sodium [Depakote/Depakote ER (extended-release)]*
- Topiramate (Topamax)*
- Venlafaxine [Effexor/Effexor XR (extended-release)*]

AND

3 - Medication will NOT be used in combination with another biologic CGRP (calcitonin generelated peptide) antagonist or inhibitor [e.g. Aimovig, Vyepti (eptinezumab-jjmr)]

| Notes | *Drug may require PA |
|-------|----------------------|
|-------|----------------------|

| Product Name: Aimovig | | |
|-----------------------|-----------------------|--|
| Diagnosis | Episodic Migraine | |
| Approval Length | 6 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Diagnosis of episodic migraines with BOTH of the following:

- Less than 15 headache days per month
- Patient has 4 to 14 migraine days per month

AND

2 - Trial and failure (after a trial of at least two months), contraindication, or intolerance to TWO of the following prophylactic therapies from the list below:

- Amitriptyline (Elavil)*
- ONE of the following beta-blockers: atenolol, metoprolol, nadolol, propranolol, or timolol*
- Divalproex sodium [Depakote/Depakote ER (extended-release)]*
- Topiramate (Topamax)*
- Venlafaxine [Effexor/Effexor XR (extended-release)]*

3 - Medication will NOT be used in combination with another biologic CGRP (calcitonin generelated peptide) antagonist or inhibitor (e.g. Ajovy, Emgality, Vyepti)

AND

4 - The patient has a history of failure, contraindication, or intolerance to BOTH of the following (document date tried):

Ajovy*

Emgality 120 milligrams*

| ļ | Notes | *Drug may require PA |
|---|-------|----------------------|

| Product Name: Aimovig, Ajovy, Emgality 120mg | | |
|--|------------------------|--|
| Diagnosis | osis Episodic Migraine | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Patient has experienced a positive response to therapy, demonstrated by a reduction in headache frequency and/or intensity

AND

2 - Medication will NOT be used in combination with another biologic CGRP (calcitonin generelated peptide) antagonist or inhibitor (e.g. Vyepti)

| Product Name: Ajovy or Emgality 120mg | |
|---------------------------------------|-----------------------|
| Diagnosis | Chronic Migraine |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- 1 Diagnosis of chronic migraines with BOTH of the following:
 - Greater than or equal to 15 headache days per month
 - Greater than or equal to 8 migraine days per month

AND

2 - Trial and failure (after a trial of at least two months), contraindication, or intolerance to TWO of the following prophylactic therapies from the list below:

- Amitriptyline (Elavil)*
- ONE of the following beta-blockers: atenolol, metoprolol, nadolol, propranolol, or timolol *
- Divalproex sodium [Depakote/Depakote ER (extended-release)*]
- Topiramate (Topamax)*
- Venlafaxine [Effexor/Effexor XR (extended-release)*]

AND

3 - Medication will NOT be used in combination with another biologic CGRP (calcitonin generelated peptide) antagonist or inhibitor

| Notes | *Drug may require PA |
|-------|----------------------|
|-------|----------------------|

| Product Name: Aimovig | |
|-----------------------|------------------|
| Diagnosis | Chronic Migraine |

| Approval Length | 6 month(s) |
|-----------------|-----------------------|
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of chronic migraines with BOTH of the following:

- Greater than or equal to 15 headache days per month
- Greater than or equal to 8 migraine days per month

AND

2 - Trial and failure (after a trial of at least two months), contraindication, or intolerance to TWO of the following prophylactic therapies from the list below (document name and date tried): prophylactic therapies from the list below:

- Amitriptyline (Elavil)*
- ONE of the following beta-blockers: atenolol, metoprolol, nadolol, propranolol, or timolol*
- Divalproex sodium [Depakote/Depakote ER (extended-release)]*
- OnabotulinumtoxinA (Botox)*
- Topiramate (Topamax)*
- Venlafaxine [Effexor/Effexor XR (extended-release)]*

AND

3 - Medication will NOT be used in combination with another biologic CGRP (calcitonin generelated peptide) antagonist or inhibitor (e.g., Ajovy Emgality, Vyepti)

AND

4 - The patient has a history of failure, contraindication, or intolerance to BOTH of the following (document date tried):

- Ajovy*
- Emgality 120 milligrams*

| Notes *Drug may require PA |
|----------------------------|
|----------------------------|

| Product Name: Aimovig, Ajovy, or Emgality 120mg | |
|---|---------------------|
| Diagnosis | Chronic Migraine |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient has experienced a positive response to therapy, demonstrated by a reduction in headache frequency and/or intensity

AND

2 - Medication will NOT be used in combination with another biologic CGRP (calcitonin generelated peptide) antagonist or inhibitor (e.g., Vyepti)

| Product Name: Emgality 100mg | |
|------------------------------|---------------------------|
| Diagnosis | Episodic Cluster Headache |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of episodic cluster headache

AND

2 - Patient has experienced at least 2 cluster periods lasting from 7 days to 365 days, separated by pain-free periods lasting at least three months

AND

3 - Medication will NOT be used in combination with another biologic CGRP (calcitonin generelated peptide) antagonist or inhibitor (e.g., Aimovig, Ajovy, Vyepti)

| Product Name: Emgality 100mg | |
|------------------------------|---------------------------|
| Diagnosis | Episodic Cluster Headache |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient has experienced a positive response to therapy, demonstrated by a reduction in headache frequency and/or intensity

AND

2 - Medication will NOT be used in combination with another biologic CGRP (calcitonin generelated peptide) antagonist or inhibitor (e.g., Aimovig, Ajovy, Vyepti)

| Date | Notes |
|-----------|-------------------------------------|
| 5/13/2021 | Arizona Medicaid 7.1 Implementation |

Cholbam

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99700 Cholbam

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Cholbam | |
|-----------------------|------------------------------|
| Diagnosis | Bile Acid Synthesis Disorder |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of a bile acid synthesis disorder

2 - It is due to single enzyme defects

| Product Name: Cholbam | |
|-----------------------|--|
| Diagnosis | Peroxisomal Disorders Including Zellweger Spectrum Disorders |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | · |

Approval Criteria

1 - Diagnosis of peroxisomal disorders including Zellweger spectrum disorders

AND

2 - Patient exhibits manifestations of liver disease, steatorrhea, or complications from decreased fat soluble vitamin absorption

AND

3 - It is being used as adjunctive treatment

| Product Name: Cholbam | | |
|-----------------------|---------------------|--|
| Diagnosis | All Indications | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |
| | | |

Approval Criteria

1 - Documentation of positive clinical response to Cholbam therapy

| Date | Notes |
|-----------|--------------------|
| 4/10/2021 | 7/1 Implementation |

Cialis for BPH - AZM

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-105174 Cialis for BPH - AZM

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 4/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Brand Cialis 5mg, generic tadalafil 5mg | | | |
|--|--|--|--|
| Diagnosis Benign Prostatic Hyperplasia (BPH) | | | |
| Approval Length 12 month(s) | | | |
| Guideline Type Prior Authorization | | | |
| | | | |
| Approval Criteria | | | |
| 1 - All of the following: | | | |
| 1.1 The patient has a diagnosis of benign prostatic hyperplasia (BPH) | | | |

1.2 History of failure, intolerance, or contraindication to BOTH of the following:

- Alpha Blockers (e.g., tamsulosin, alfuzosin ER, doxazosin, or terazosin)
- 5-alpha reductase inhibitors (e.g., finasteride)

AND

1.3 Dose does not exceed 5 milligrams once daily

AND

2 - Provider attests that patient is not using any form of organic nitrate (for example, nitroglycerin, isosorbide dinitrate, isosorbide mononitrate or amyl nitrate) or Adempas

| Date | Notes |
|-----------|--|
| 3/24/2022 | Added physician attestation re: patient not using nitrates |

Cibinqo (abrocitinib)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-104977 | Cibinqo (abrocitinib) |
|-----------|-----------------------|
|-----------|-----------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 4/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Cibinqo | | |
|-----------------------|--|--|
| 6 month(s) | | |
| Initial Authorization | | |
| Prior Authorization | | |
| | | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of moderate to severe atopic dermatitis

2 - One of the following:

- Involvement of at least 10% body surface area (BSA)
- SCORing Atopic Dermatitis (SCORAD) index value of at least 25 [A]

AND

3 - Prescribed by or in consultation with one of the following:

- Dermatologist
- Allergist/Immunologist

AND

4 - Trial and failure of a minimum 30-day supply (14-day supply for topical corticosteroids), contraindication, or intolerance to at least TWO of the following:

- Medium or higher potency topical corticosteroid
- Pimecrolimus cream*
- Tacrolimus ointment
- Eucrisa (crisaborole) ointment*

AND

5 - One of the following:

5.1 Trial and failure of a minimum 12-week supply of at least one systemic drug product for the treatment of atopic dermatitis (examples include, but are not limited to, Adbry [tralokinumab-ldrm], Dupixent [dupilumab], etc.)

OR

5.2 Patient has a contraindication, intolerance, or treatment is inadvisable with both of the following FDA-approved atopic dermatitis therapies:

• Adbry (tralokinumab-ldrm)

• Dupixent (dupilumab)

AND

6 - Not used in combination with biologic immunomodulators (e.g., Dupixent, Adbry) or other immunosuppressants (e.g., azathioprine, cyclosporine)

| Notes | *Product may require step therapy |
|-------|-----------------------------------|
|-------|-----------------------------------|

| Product Name: Cibinqo | | |
|-----------------------|---------------------|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting a positive clinical response to therapy as evidenced by at least ONE of the following:

- Reduction in body surface area involvement from baseline
- Reduction in SCORing Atopic Dermatitis (SCORAD) index value from baseline [A]

AND

2 - Not used in combination with biologic immunomodulators (e.g., Dupixent, Adbry) or other immunosuppressants (e.g., azathioprine, cyclosporine)

2. Background

| Clinical Practice | Guidelines | | |
|--|------------|-------------|----------|
| Table 1. Relative potencies of topical corticosteroids [2] | | | |
| | | | |
| Class | Drug | Dosage Form | Strength |
| | - | - | (%) |
| | | | |

| Very high potency | Augmented betamethasone dipropionate | Ointment, gel | 0.05 |
|-------------------|--------------------------------------|---------------------------------|-------|
| | Clobetasol propionate | Cream, foam, ointment | 0.05 |
| | Diflorasone diacetate | Ointment | 0.05 |
| | Halobetasol propionate | Cream, ointment | 0.05 |
| High | Amcinonide | Cream, lotion, ointment | 0.1 |
| Potency | Augmented betamethasone dipropionate | Cream, lotion | 0.05 |
| | Betamethasone dipropionate | Cream, foam, ointment, solution | 0.05 |
| | Desoximetasone | Cream, ointment | 0.25 |
| | Desoximetasone | Gel | 0.05 |
| | Diflorasone diacetate | Cream | 0.05 |
| | Fluocinonide | Cream, gel, ointment, solution | 0.05 |
| | Halcinonide | Cream, ointment | 0.1 |
| | Mometasone furoate | Ointment | 0.1 |
| | Triamcinolone acetonide | Cream, ointment | 0.5 |
| Medium | Betamethasone valerate | Cream, foam, lotion, ointment | 0.1 |
| potency | Clocortolone pivalate | Cream | 0.1 |
| | Desoximetasone | Cream | 0.05 |
| | Fluocinolone acetonide | Cream, ointment | 0.025 |
| | Flurandrenolide | Cream, ointment, lotion | 0.05 |
| | Fluticasone propionate | Cream | 0.05 |
| | Fluticasone propionate | Ointment | 0.005 |
| | Mometasone furoate | Cream, lotion | 0.1 |
| | Triamcinolone acetonide | Cream, ointment, lotion | 0.1 |
| | Hydrocortisone butyrate | Cream, ointment, solution | 0.1 |
| | Hydrocortisone probutate | Cream | 0.1 |

| Lower- medium potency | Hydrocortisone valerate | Cream, ointment | 0.2 |
|-----------------------------|----------------------------|-----------------------------------|--------------|
| | Prednicarbate | Cream | 0.1 |
| Low potency | Alclometasone dipropionate | Cream, ointment | 0.05 |
| | Desonide | Cream, gel, foam, ointment | 0.05 |
| | Fluocinolone acetonide | Cream, solution | 0.01 |
| Lowest potency | Dexamethasone | Cream | 0.1 |
| | Hydrocortisone | Cream, lotion, ointment, solution | 0.25, 0.5, 1 |
| | Hydrocortisone acetate | Cream, ointment | 0.5-1 |

3. Endnotes

A. The Scoring Atopic Dermatitis (SCORAD) index is a clinical tool for assessing the severity of atopic dermatitis lesions based on affected body area and intensity of plaque characteristics. [3, 4] The extent and severity of AD over the body area (A) and the severity of 6 specific symptoms (erythema, edema/papulation, excoriations, lichenification, oozing/crusts, and dryness) (B) are assessed and scored by the Investigator. Subjective assessment of itch and sleeplessness is scored by the patient (C). The SCORAD score is a combined score (A/5 + 7B/2 + C) with a maximum of 103. Higher scores indicate greater severity/worsened state. A score of 25 to 50 indicates moderate disease severity and greater than 50 indicates severe disease. [5]

| Date | Notes |
|-----------|---|
| 3/22/2022 | New Program mirrors ORx with Submission of Records added to initi al and reauth |

Cimzia- Arizona

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99712 Cimzia- Arizona

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Cimzia | | |
|----------------------|-----------------------|--|
| Diagnosis | Crohn's Disease | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |

Approval Criteria

1 - One of the following:

1.1 All of the following:

1.1.1 Diagnosis of moderately to severely active Crohn's disease

AND

1.1.2 History of failure to ONE of the following conventional therapies at maximally indicated doses within the last 3 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial):*

- Corticosteroids (e.g., prednisone, methylprednisolone, budesonide)
- 6-mercaptopurine (Purinethol)
- Azathioprine (Imuran)
- Methotrexate (Rheumatrex, Trexall)

AND

1.1.3 Patient is NOT receiving Cimzia in combination with ONE of the following:

- Biologic DMARD (disease modifying antirheumatic drug) [e.g., Enbrel (etanercept), Humira (adalimumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

1.1.4 History of failure, contraindication, or intolerance to Humira (adalimumab)

AND

1.1.5 Prescribed by or in consultation with a gastroenterologist

OR

1.2 All of the following:

1.2.1 Patient is currently on Cimzia therapy as documented by claims history or medical records (document drug, date, and duration of therapy)

AND 1.2.2 Diagnosis of Crohn's disease AND **1.2.3** Patient is NOT receiving Cimzia in combination with ONE of the following: Biologic DMARD [e.g., Enbrel (etanercept), Humira (adalimumab), Simponi • (golimumab)] Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)] • Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)] • AND 1.2.4 Prescribed by or in consultation with a gastroenterologist Notes *Note: Claims history may be used in conjunction as documentation of drug, date, and duration of trial

| Product Name: Cimzia | |
|----------------------|---------------------|
| Diagnosis | Crohn's Disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Cimzia therapy

AND

2 - Patient is NOT receiving Cimzia in combination with ONE of the following:

- Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Enbrel (etanercept), Humira (adalimumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

3 - Prescribed by or in consultation with a gastroenterologist

| Product Name: Cimzia | |
|----------------------|---------------------------|
| Diagnosis | Rheumatoid Arthritis (RA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- **1** One of the following:
- **1.1** All of the following:

1.1.1 Diagnosis of moderately to severely active rheumatoid arthritis (RA)

AND

1.1.2 History of failure to a 3 month trial of one non-biologic disease modifying antirheumatic drug (DMARD) [eg, methotrexate, leflunomide, sulfasalazine, hydroxychloroquine] at maximally indicated doses within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*

AND

1.1.3 Patient is NOT receiving Cimzia in combination with ONE of the following:

- Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Enbrel (etanercept), Humira (adalimumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]

Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)] • AND **1.1.4** History of failure, contraindication, or intolerance to ALL of the following: Humira (adalimumab) • Enbrel (etanercept) ٠ Xeljanz (tofacitinib) • AND **1.1.5** Prescribed by or in consultation with a rheumatologist OR **1.2** All of the following: **1.2.1** Patient is currently on Cimzia therapy as documented by claims history or medical records (document drug, date, and duration of therapy) AND 1.2.2 Diagnosis of moderately to severely active RA AND **1.2.3** Patient is NOT receiving Cimzia in combination with ONE of the following: Biologic DMARD [e.g., Enbrel (etanercept), Humira (adalimumab), Simponi ٠ (golimumab)] Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)] ٠ Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)] • AND 1.2.4 Prescribed by or in consultation with a rheumatologist

| Notes | *Note: Claims history may be used in conjunction as documentation of drug, date, and duration of trial |
|-------|--|
| | drug, date, and duration of that |

| Product Name: Cimzia | |
|----------------------|---------------------------|
| Diagnosis | Rheumatoid Arthritis (RA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Cimzia therapy

AND

- **2** Patient is NOT receiving Cimzia in combination with ONE of the following:
 - Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Enbrel (etanercept), Humira (adalimumab), Simponi (golimumab)]
 - Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
 - Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with a rheumatologist

| Product Name: Cimzia | |
|----------------------|-----------------------|
| Diagnosis | Psoriatic Arthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | • |

Approval Criteria

1 - One of the following:

1.1 All of the following:

1.1.1 Diagnosis of active psoriatic arthritis

AND

1.1.2 History of failure to a 3 month trial of methotrexate at the maximally indicated dose within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*

AND

1.1.3 Patient is NOT receiving Cimzia in combination with ONE of the following:

- Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Enbrel (etanercept), Humira (adalimumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

1.1.4 History of failure, contraindication, or intolerance to THREE of the following:

- Humira (adalimumab)
- Enbrel (etanercept)
- Otezla (apremilast)
- Xeljanz (tofacitinib)

AND

1.1.5 Prescribed by or in consultation with ONE of the following:

- Rheumatologist
- Dermatologist

1.2 All of the following:

| 1.2.1 Patient is currently on Cimzia therapy a records (document drug, date, and duration of the second s | |
|---|---|
| A | ۱D |
| 1.2.2 Diagnosis of active psoriatic arthritis | |
| 1A | ND |
| 1.2.3 Patient is NOT receiving Cimzia in com | bination with ONE of the following: |
| Biologic DMARD [e.g., Enbrel (etanerce (golimumab)] Janus kinase inhibitor [e.g., Xeljanz (tof Phosphodiesterase 4 (PDE4) inhibitor [etanerce] | acitinib)] |
| A | ۱D |
| 1.2.4 Prescribed by or in consultation with ON | IE of the following: |
| RheumatologistDermatologist | |
| Notes *Note: Claims history m drug, date, and duration | nay be used in conjunction as documentation of on of trial |

| Product Name: Cimzia | |
|----------------------|---------------------|
| Diagnosis | Psoriatic Arthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

Г

1 - Documentation of positive clinical response to Cimzia therapy

2 - Patient is NOT receiving Cimzia in combination with ONE of the following:

- Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Enbrel (etanercept), Humira (adalimumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with ONE of the following:

- Rheumatologist
- Dermatologist

| Product Name: Cimzia | |
|----------------------|--|
| Diagnosis | Ankylosing Spondylitis or Non-Radiographic Axial Spondyloarthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - One of the following:

1.1 All of the following:

1.1.1 Diagnosis of active ankylosing spondylitis or non-radiographic axial spondyloarthritis

AND

1.1.2 History of failure to two NSAIDs (non-steroidal anti-inflammatory drugs; e.g., ibuprofen, naproxen) at maximally indicated doses, each used for at least 4 weeks within the last 3 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trials)*

| AND |
|--|
| |
| 1.1.3 Patient is NOT receiving Cimzia in combination with ONE of the following: |
| Biologic DMARD (disease modifying antirheumatic drug) [e.g., Enbrel (etanercept), Humira (adalimumab), Simponi (golimumab)] Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)] Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)] |
| AND |
| |
| 1.1.4 History of failure, contraindication, or intolerance to BOTH of the following: |
| Humira (adalimumab)Enbrel (etanercept) |
| AND |
| |
| 1.1.5 Prescribed by or in consultation with a rheumatologist |
| OR |
| 1.2 All of the following: |
| 1.2.1 Patient is currently on Cimzia therapy as documented by claims history or medical records (document drug, date, and duration of therapy) |
| AND |
| |
| 1.2.2 Diagnosis of active ankylosing spondylitis or non-radiographic axial spondyloarthritis |
| AND |
| 1.2.3 Patient is NOT receiving Cimzia in combination with ONE of the following: |
| |

• Biologic DMARD [e.g., Enbrel (etanercept), Humira (adalimumab), Simponi (golimumab)]

- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

1.2.4 Prescribed by or in consultation with a rheumatologist

| *Note: Claims history may be used in conjunction as documentation of |
|--|
| drug, date, and duration of trials |

| Product Name: Cimzia | |
|----------------------|--|
| Diagnosis | Ankylosing Spondylitis or Non-Radiographic Axial Spondyloarthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Cimzia therapy

AND

- 2 Patient is NOT receiving Cimzia in combination with ONE of the following:
 - Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Enbrel (etanercept), Humira (adalimumab), Simponi (golimumab)]
 - Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
 - Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with a rheumatologist

Product Name: Cimzia

| Diagnosis | Plaque Psoriasis |
|-----------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - One of the following:

1.1 All of the following:

1.1.1 Diagnosis of moderate to severe plaque psoriasis

AND

1.1.2 Greater than or equal to 3% body surface area involvement, palmoplantar, facial, or genital involvement, or severe scalp psoriasis

AND

1.1.3 History of failure to one of the following topical therapies, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial):*

- Corticosteroids (e.g., betamethasone, clobetasol, desonide)
- Vitamin D analogs (e.g., calcitriol, calcipotriene)
- Tazarotene
- Calcineurin inhibitors (e.g., tacrolimus, pimecrolimus)
- Anthralin
- Coal tar

AND

1.1.4 History of failure of a 3 month trial of methotrexate at the maximally indicated dose within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*

AND

| 1.1.5 Patient is NOT receiving Cimzia in combination with ONE of the following: | |
|--|--|
| Biologic DMARD (disease modifying antirheumatic drug) [e.g., Enbrel (etanercept), Humira (adalimumab), Simponi (golimumab)] Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)] Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)] | |
| AND | |
| 1.1.6 History of failure, contraindication, or intolerance to ALL of the following: | |
| Humira (adalimumab) Enbrel (etanercept) Otezla (apremilsat) | |
| AND | |
| 1.1.7 Prescribed by or in consultation with a dermatologist | |
| OR | |
| 1.2 All of the following: | |
| 1.2.1 Patient is currently on Cimzia therapy as documented by claims history or medical records (document drug, date, and duration of therapy) | |
| AND | |
| 1.2.2 Diagnosis of moderate to severe plaque psoriasis | |
| AND | |
| 1.2.3 Patient is NOT receiving Cimzia in combination with ONE of the following: | |
| Biologic DMARD [e.g., Enbrel (etanercept), Humira (adalimumab), Simponi (golimumab)] Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)] | |
| | |

| Phosphodiester | ase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)] | |
|---|---|--|
| | AND | |
| 1.2.4 Prescribed by or in consultation with a dermatologist | | |
| Notes | *Note: Claims history may be used in conjunction as documentation of drug, date, and duration of trials | |
| | 5 | |

| Product Name: Cimzia | |
|----------------------|---------------------|
| Diagnosis | Plaque Psoriasis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Cimzia therapy

AND

- **2** Patient is NOT receiving Cimzia in combination with ONE of the following:
 - Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Enbrel (etanercept), Humira (adalimumab), Simponi (golimumab)]
 - Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
 - Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with a dermatologist

2. Revision History

| Date | Notes |
|------|-------|
|------|-------|

| 5/19/2021 |
|-----------|
|-----------|

Cinryze

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99662 Cinryze

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Cinryze | |
|-----------------------|-----------------------------|
| Diagnosis | Hereditary angioedema (HAE) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of hereditary angioedema (HAE) as confirmed by ONE of the following:

1.1 C1 inhibitor (C1-INH) deficiency or dysfunction (Type I or II HAE) as documented by one of the following (per laboratory standard): C1-INH antigenic level below the lower limit of normal • C1-INH functional level below the lower limit of normal OR **1.2** HAE with normal C1 inhibitor levels and one of the following: Confirmed presence of a FXII, angiopoietin-1 or plasminogen gene mutation • Recurring angioedema attacks that are refractory to high-dose antihistamines with • confirmed family history of angioedema AND **2** - Prescribed for the prophylaxis of HAE attacks AND 3 - Not used in combination with other approved C1 esterase inhibitors indicated for prophylaxis against HAE attacks (e.g., Haegarda) AND 4 - Prescriber attests that the patient has experienced attacks of a severity and/or frequency such that they would clinically benefit from prophylactic therapy with Cinryze AND 5 - ONE of the following: 5.1 Submission of medical records documenting a history of failure, contraindication, or intolerance to Haegarda (C1 esterase inhibitor, human) OR

5.2 Patient is currently on Cinryze therapy

AND

6 - Prescribed by ONE of the following:

- Immunologist
- Allergist

| Product | Name: | Cinry | vze |
|----------|-------|-------|-------|
| 1 100000 | numo. | | y 2 0 |

| Hereditary angioedema (HAE) | |
|-----------------------------|--|
| 12 month(s) | |
| Reauthorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Documentation of positive clinical response, defined as a clinically significant reduction in the rate and/or number of HAE attacks, while on Cinryze therapy

AND

2 - Reduction in the utilization of on-demand therapies used for acute attacks (e.g., Berinert, Firazyr, Ruconest) as determined by claims information, while on Cinryze therapy

AND

3 - Prescribed for the prophylaxis of HAE attacks

AND

4 - Not used in combination with other products indicated for prophylaxis against HAE attacks (e.g., Haegarda, Takhzyro)

AND 5 - Prescribed by ONE of the following: • Immunologist • Allergist

2. Revision History

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk copy C&S Arizona Medicaid SP to Medicaid Arizona SP for eff 7 /1 |

CMV and Herpes Virus Agents- Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99518 CMV and Herpes Virus Agents- Arizona

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Valcyte tabs/oral soln, generic valganciclovir tabs/oral soln, Brand Cytovene inj, generic ganciclovir inj, Foscavir inj | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Medication is being used for ONE of the following:

1.1 Cytomegalovirus (CMV) disease prophylaxis

OR

1.2 Cytomegalovirus (CMV) retinitis

OR

1.3 Cytomegalovirus (CMV) retinitis prophylaxis

OR

1.4 BOTH of the following:

1.4.1 The use of this drug is supported by information from ONE of the following appropriate compendia of current literature:

- American Hospital Formulary Service Drug Information
- National Comprehensive Cancer Network Drugs and Biologics Compendium
- Thomson Micromedex DrugDex
- Clinical pharmacology

AND

1.4.2 The drug is being prescribed for a medically accepted indication that is recognized as a covered benefit by the applicable health plan's program*

| *Note: Medications used solely for anti-obesity/weight loss, cosmetic (e.g., alopecia, actinic keratosis, vitiligo), erectile dysfunction, and sexu al dysfunction purposes are NOT medically accepted indications and |
|---|
| are NOT recognized as a covered benefit. Erectile dysfunction drugs (Cialis/Tadalafil) are covered for clinical diagnoses other than ED. |

| Product Name: cidofovir inj | |
|-----------------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Medication is being used for ONE of the following:

1.1 Cytomegalovirus (CMV) retinitis

OR

1.2 Cytomegalovirus (CMV) retinitis prophylaxis

OR

1.3 BOTH of the following:

1.3.1 The use of this drug is supported by information from ONE of the following appropriate compendia of current literature:

- American Hospital Formulary Service Drug Information
- National Comprehensive Cancer Network Drugs and Biologics Compendium
- Thomson Micromedex DrugDex
- Clinical pharmacology

AND

1.3.2 The drug is being prescribed for a medically accepted indication that is recognized as a covered benefit by the applicable health plan's program*

| *Note: Medications used solely for anti-obesity/weight loss, cosmetic (e.g., alopecia, actinic keratosis, vitiligo), erectile dysfunction, and sexu al dysfunction purposes are NOT medically accepted indications and are NOT recognized as a covered benefit. Erectile dysfunction drugs (Cialis/Tadalafil) are covered for clinical diagnoses other than ED |
|--|
| Cialis/Tadalafil) are covered for clinical diagnoses other than ED. |

| Product Name: famciclovir tabs | |
|--------------------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Medication is being used for ONE of the following:

| 1.1 Herpes genitalis |
|---|
| OR |
| 1.2 Herpes genitalis prophylaxis |
| OR |
| 1.3 Herpes labialis |
| OR |
| 1.4 Herpes simplex virus infection |
| OR |
| 1.5 Herpes zoster (shingles) infection |
| OR |
| 1.6 BOTH of the following: |
| 1.6.1 The use of this drug is supported by information from ONE of the following appropriate compendia of current literature: |
| American Hospital Formulary Service Drug Information National Comprehensive Cancer Network Drugs and Biologics Compendium Thomson Micromedex DrugDex Clinical pharmacology |
| AND |
| 1.6.2 The drug is being prescribed for a medically accepted indication that is recognized as a covered benefit by the applicable health plan's program* |

| Notes | *Note: Medications used solely for anti-obesity/weight loss, cosmetic (e.g., alopecia, actinic keratosis, vitiligo), erectile dysfunction, and sexu al dysfunction purposes are NOT medically accepted indications and are NOT recognized as a covered benefit. Erectile dysfunction drugs (|
|-------|--|
| | Cialis/Tadalafil) are covered for clinical diagnoses other than ED. |

| Product Name: Brand Valtrex tabs, generic valacyclovir tabs | | |
|---|---------------------|--|
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Medication is being used for ONE of the following: | | |
| 1.1 Herpes genitalis | | |
| | | |
| | OR | |
| 1.2 Herpes genitalis prophylaxis | | |
| | OR | |
| | | |
| 1.3 Herpes labialis | | |
| | OR | |
| | | |
| 1.4 Herpes simplex virus infection | | |
| | OR | |
| | | |
| 1.5 Herpes zoster (shingles) infection | | |
| | OR | |
| 1.6 Varicella (chicken | pox) infection | |

1.7 BOTH of the following

1.7.1 The use of this drug is supported by information from ONE of the following appropriate compendia of current literature:

- American Hospital Formulary Service Drug Information
- National Comprehensive Cancer Network Drugs and Biologics Compendium
- Thomson Micromedex DrugDex
- Clinical pharmacology

AND

1.7.2 The drug is being prescribed for a medically accepted indication that is recognized as a covered benefit by the applicable health plan's program*

| *Note: Medications used solely for anti-obesity/weight loss, cosmetic (e.g., alopecia, actinic keratosis, vitiligo), erectile dysfunction, and sexu al dysfunction purposes are NOT medically accepted indications and are NOT recognized as a covered benefit. Erectile dysfunction drugs (Cialis/Tadalafil) are covered for clinical diagnoses other than ED |
|--|
| Cialis/Tadalafil) are covered for clinical diagnoses other than ED. |
| |

2. Revision History

| Date | Notes |
|-----------|-------------------------------------|
| 5/13/2021 | Arizona Medicaid 7.1 Implementation |

Colony Stimulating Factors - Arizona

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99802 Colony Stimulating Factors - Arizona

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Neupogen, Nivestym | |
|----------------------------------|----------------------------------|
| Diagnosis | Bone Marrow/Stem Cell Transplant |
| Approval Length | 3 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - ONE of the following:

1.1 Patient has non-myeloid malignancies and is undergoing myeloablative chemotherapy followed by autologous or allogeneic bone marrow transplant (BMT)

OR

1.2 Used for mobilization of hematopoietic progenitor cells into the peripheral blood for collection by leukapheresis

OR

1.3 Patient has had a peripheral stem cell transplant (PSCT) and has received myeloablative chemotherapy

AND

2 - Prescribed by, or in consultation with, a hematologist or oncologist

| Product Name: Leukine, Zarxio | |
|-------------------------------|----------------------------------|
| Diagnosis | Bone Marrow/Stem Cell Transplant |
| Approval Length | 3 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - ONE of the following:

1.1 Patient has non-myeloid malignancies and is undergoing myeloablative chemotherapy followed by autologous or allogeneic bone marrow transplant (BMT)

OR

1.2 Used for mobilization of hematopoietic progenitor cells into the peripheral blood for collection by leukapheresis

1.3 Patient has had a peripheral stem cell transplant (PSCT) and has received myeloablative chemotherapy

AND

2 - Prescribed by, or in consultation with, a hematologist or oncologist

AND

3 - Patient has a history of failure, contraindication, or intolerance to Neupogen and Nivestym

| Product Name: Neupogen, Nivestym | |
|----------------------------------|--|
| Diagnosis | AML Induction or Consolidation Therapy |
| Approval Length | 3 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of acute myeloid leukemia (AML)

AND

2 - Patient has completed either induction or consolidation chemotherapy

AND

3 - Prescribed by, or in consultation with, a hematologist or oncologist

| Product Name: Zarxio, Leukine | |
|-------------------------------|--|
| Diagnosis | AML Induction or Consolidation Therapy |
| Approval Length | 3 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of acute myeloid leukemia (AML)

AND

2 - Patient has completed either induction or consolidation chemotherapy

AND

3 - Prescribed by, or in consultation with, a hematologist or oncologist

AND

4 - Patient has a history of failure, contraindication, or intolerance to Neupogen and Nivestym

| Product Name: Fulphila, Udenyca, Neupogen, Nivestym, Nyvepria | |
|---|--|
| Diagnosis | Neutropenia Associated with Cancer Chemotherapy –Dose Dense Chemotherapy |
| Approval Length | 3 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - ONE of the following:

1.1 Patient is receiving National Cancer Institute's Breast Intergroup, INT C9741 dose dense chemotherapy protocol for primary breast cancer

OR

1.2 Patient is receiving a dose-dense chemotherapy regimen for which the incidence of febrile neutropenia (FN) is unknown

2 - Prescribed by, or in consultation with, a hematologist or oncologist

| Product Name: Leukine, Zarxio | |
|-------------------------------|---|
| Diagnosis | Neutropenia Associated with Cancer Chemotherapy –Dose Dense Chemotherapy |
| Approval Length | 3 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - ONE of the following:

1.1 Patient is receiving National Cancer Institute's Breast Intergroup, INT C9741 dose dense chemotherapy protocol for primary breast cancer

OR

1.2 Patient is receiving a dose-dense chemotherapy regimen for which the incidence of febrile neutropenia (FN) is unknown

AND

2 - Prescribed by, or in consultation with, a hematologist or oncologist

AND

3 - Patient has a history of failure, contraindication, or intolerance to Neupogen and Nivestym

| Product Name: Neulasta, Neulasta Onpro, Ziextenzo | |
|---|---|
| Diagnosis | Neutropenia Associated with Cancer Chemotherapy –Dose Dense Chemotherapy |

| Approval Length | 3 month(s) |
|---|--|
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - ONE of the follow | <i>v</i> ing: |
| | ving National Cancer Institute's Breast Intergroup, INT C9741 dose dense col for primary breast cancer |
| | OR |
| 1.2 Patient is receiv febrile neutropenia (| ving a dose-dense chemotherapy regimen for which the incidence of FN) is unknown |
| | AND |
| 2 - Prescribed by, or | in consultation with, a hematologist or oncologist |
| | AND |
| 3 - Patient has a his | ory of failure, contraindication, or intolerance to Fulphila, Udenyca, and |

| | • |
|----------|---|
| Nyvepria | |

| Product Name: Fulphila, Udenyca, Neupogen, Nivestym, Nyvepria | |
|---|--|
| Diagnosis | Primary Prophylaxis of Chemotherapy-Induced Febrile Neutropenia (FN) |
| Approval Length | 3 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - ONE of the following:

1.1 Patient is receiving chemotherapy regimen(s) associated with greater than 20 percent incidence of febrile neutropenia (FN)

OR

1.2 BOTH of the following:

- Patient is receiving chemotherapy regimen(s) associated with 10-20 percent incidence of FN
- Patient has one or more risk factors associated with chemotherapy-induced infection, FN, or neutropenia

AND

2 - Prescribed by, or in consultation with, a hematologist or oncologist

| Product Name: Granix,Zarxio | |
|-----------------------------|--|
| Diagnosis | Primary Prophylaxis of Chemotherapy-Induced Febrile Neutropenia (FN) |
| Approval Length | 3 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - ONE of the following:

1.1 Patient is receiving chemotherapy regimen(s) associated with greater than 20 percent incidence of febrile neutropenia (FN)

OR

1.2 BOTH of the following:

- Patient is receiving chemotherapy regimen(s) associated with 10-20 percent incidence of FN
- Patient has one or more risk factors associated with chemotherapy-induced infection, FN, or neutropenia

2 - Prescribed by, or in consultation with, a hematologist or oncologist

AND

3 - Patient has a history of failure, contraindication, or intolerance to Neupogen and Nivestym

| Product Name: Neulasta, Neulasta Onpro, Ziextenzo | |
|---|--|
| Diagnosis | Primary Prophylaxis of Chemotherapy-Induced Febrile Neutropenia (FN) |
| Approval Length | 3 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - ONE of the following:

1.1 Patient is receiving chemotherapy regimen(s) associated with greater than 20 percent incidence of febrile neutropenia (FN)

OR

1.2 BOTH of the following:

- Patient is receiving chemotherapy regimen(s) associated with 10-20 percent incidence of FN
- Patient has one or more risk factors associated with chemotherapy-induced infection, FN, or neutropenia

AND

2 - Prescribed by, or in consultation with, a hematologist or oncologist

3 - Patient has a history of failure, contraindication, or intolerance to Fulphila,Udenyca, and Nyvepria

| Product Name: Fulphila, Udenyca, Neupogen, Nivestym, Nyvepria | |
|---|---|
| Diagnosis | Secondary Prophylaxis of Febrile Neutropenia (FN) |
| Approval Length | 3 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient is receiving myelosuppressive anti-cancer drugs associated with neutropenia (absolute neutrophil count [ANC] less than or equal to 500 cells per mm3)

AND

2 - Patient has a history of febrile neutropenia (FN) during a previous course of chemotherapy

AND

3 - Prescribed by, or in consultation with, a hematologist or oncologist

| Product Name: Granix,Zarxio | |
|-----------------------------|---|
| Diagnosis | Secondary Prophylaxis of Febrile Neutropenia (FN) |
| Approval Length | 3 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient is receiving myelosuppressive anti-cancer drugs associated with neutropenia (absolute neutrophil count [ANC] less than or equal to 500 cells per mm3)

2 - Patient has a history of febrile neutropenia (FN) during a previous course of chemotherapy

AND

3 - Prescribed by, or in consultation with, a hematologist or oncologist

AND

4 - Patient has a history of failure, contraindication, or intolerance to Neupogen, Fulphila, Udenyca, Nivestym, Nyvepria *

| Product Name: Neulasta, Neulasta Onpro, Ziextenzo | | |
|---|---|--|
| Diagnosis | Secondary Prophylaxis of Febrile Neutropenia (FN) | |
| Approval Length | 3 month(s) | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Patient is receiving myelosuppressive anti-cancer drugs associated with neutropenia (absolute neutrophil count [ANC] less than or equal to 500 cells per mm3)

AND

2 - Patient has a history of febrile neutropenia (FN) during a previous course of chemotherapy

AND

3 - Prescribed by, or in consultation with, a hematologist or oncologist

4 - Patient has a history of failure, contraindication, or intolerance to Fulphila,Udenyca, and Nyvepria

| Product Name: Fulphila, Udenyca, Neupogen, Nyvepria, Nivestym | | |
|---|---|--|
| Diagnosis | Treatment of Febrile Neutropenia (FN) (off-label) | |
| Approval Length | 1 month(s) | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Patient is receiving myelosuppressive anti-cancer drugs associated with neutropenia (absolute neutrophil count [ANC] less than or equal to 500 cells per mm3)

AND

2 - Diagnosis of febrile neutropenia (FN) and patient is considered high risk for infectionassociated complications

AND

3 - Prescribed by, or in consultation with, a hematologist or oncologist

| Product Name: Leukine, Zarxio | | |
|---|--|--|
| Treatment of Febrile Neutropenia (FN) (off-label) | | |
| 1 month(s) | | |
| Prior Authorization | | |
| | | |

Approval Criteria

1 - Patient is receiving myelosuppressive anti-cancer drugs associated with neutropenia (absolute neutrophil count [ANC] less than or equal to 500 cells per mm3)

2 - Diagnosis of febrile neutropenia (FN) and patient is considered high risk for infectionassociated complications

AND

3 - Prescribed by, or in consultation with, a hematologist or oncologist

AND

4 - Patient has a history of failure, contraindication, or intolerance to Fulphila, Udenyca, Neupogen, Nivestym, Nyvepria*

| Product Name: Neulasta, Neulasta Onpro, Ziextenzo | | |
|---|---|--|
| Diagnosis | Treatment of Febrile Neutropenia (FN) (off-label) | |
| Approval Length | 1 month(s) | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Patient is receiving myelosuppressive anti-cancer drugs associated with neutropenia (absolute neutrophil count [ANC] less than or equal to 500 cells per mm3)

AND

2 - Diagnosis of febrile neutropenia (FN) and patient is considered high risk for infectionassociated complications

AND

3 - Prescribed by, or in consultation with, a hematologist or oncologist

AND

4 - Patient has a history of failure, contraindication, or intolerance to Fulphila and Udenyca and Nyvepria

| Product Name: Neupogen, Nivestym | |
|----------------------------------|----------------------------------|
| Diagnosis | Severe Chronic Neutropenia (SCN) |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of severe chronic neutropenia (SCN) (i.e., congenital, cyclic, and idiopathic neutropenias with chronic absolute neutrophil count [ANC] less than or equal to 500 cells per mm3)

AND

2 - Prescribed by, or in consultation with, a hematologist or oncologist

| Product Name: Zarxio | |
|----------------------|----------------------------------|
| Diagnosis | Severe Chronic Neutropenia (SCN) |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of severe chronic neutropenia (SCN) (i.e., congenital, cyclic, and idiopathic neutropenias with chronic absolute neutrophil count [ANC] less than or equal to 500 cells per mm3)

2 - Prescribed by, or in consultation with, a hematologist or oncologist

AND

3 - Patient has a history of failure, contraindication, or intolerance to Neupogen and Nivestym

| Product Name: Neupogen, Nivestym | |
|----------------------------------|-------------------------------------|
| Diagnosis | HIV-Related Neutropenia (off-label) |
| Approval Length | 6 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of human immunodeficiency virus (HIV) infection

AND

2 - Patient has an absolute neutrophil count (ANC) less than or equal to 1,000 cells per mm3

AND

3 - Prescribed by, or in consultation with, ONE of the following:

- Hematologist
- Oncologist
- Infectious disease specialist

| HV-Related Neutropenia (off-label) |
|------------------------------------|
| |
| S month(s) |
| Prior Authorization |
| |
| |

4 - Patient has a history of failure, contraindication, or intolerance to Neupogen and Nivestym

| Product Name: Neupogen, Nivestym | |
|----------------------------------|---|
| Diagnosis | Hepatitis C Treatment Related Neutropenia (off-label) |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | · |

Approval Criteria

- **1** ONE of the following:
- **1.1** ALL of the following:
 - Diagnosis of hepatitis C virus
 - Patient is undergoing treatment with Peg-Intron (peginterferon alfa-2b) or Pegasys (peginterferon alfa-2a)
 - Documentation of neutropenia (absolute neutrophil count [ANC] less than or equal to 500 cells per mm3) after dose reduction of Peg-Intron or Pegasys

OR

1.2 BOTH of the following:

1.2.1 Documentation of interferon-induced neutropenia (ANC less than or equal to 500 cells per mm3) due to treatment with Peg-Intron (peginterferon alfa-2b) or Pegasys (peginterferon alfa-2a)

AND

1.2.2 ONE of the following:

- Diagnosis of human immunodeficiency virus (HIV) co-infection
- Status post liver transplant
- Diagnosis of established cirrhosis

AND

2 - Prescribed by, or in consultation with, a hematologist, oncologist, gastroenterologist, hepatologist, or infectious disease specialist

| Product Name: Zarxio | |
|---|--|
| Hepatitis C Treatment Related Neutropenia | |
| 12 month(s) | |
| Prior Authorization | |
| | |

Approval Criteria

- **1** ONE of the following:
- **1.1** ALL of the following:
 - Diagnosis of hepatitis C virus
 - Patient is undergoing treatment with Peg-Intron (peginterferon alfa-2b) or Pegasys (peginterferon alfa-2a)
 - Documentation of neutropenia (absolute neutrophil count [ANC] less than or equal to 500 cells per mm3) after dose reduction of Peg-Intron or Pegasys

OR

1.2 BOTH of the following:

1.2.1 Documentation of interferon-induced neutropenia (ANC less than or equal to 500 cells per mm3) due to treatment with Peg-Intron (peginterferon alfa-2b) or Pegasys (peginterferon alfa-2a)

AND

1.2.2 ONE of the following:

- Diagnosis of human immunodeficiency virus (HIV) co-infection
- Status post liver transplant
- Diagnosis of established cirrhosis

AND

2 - Prescribed by, or in consultation with, a hematologist, oncologist, gastroenterologist, hepatologist, or infectious disease specialist

AND

3 - Patient has a history of failure, contraindication, or intolerance to Neupogen and Nivestym

| Product Name: Neupogen, Fulphila, Udenyca, Nivestym, Nyverpria | |
|--|--|
| Diagnosis | Hematopoietic Syndrome of Acute Radiation Syndrome |
| Approval Length | 3 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient has been acutely exposed to myelosuppressive doses of radiation

AND

2 - Prescribed by, or in consultation with, a hematologist or oncologist

| Product Name: Leukine, Zarxio | |
|--|--|
| Hematopoietic Syndrome of Acute Radiation Syndrome | |
| 3 month(s) | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Patient has been acutely exposed to myelosuppressive doses of radiation

AND

2 - Prescribed by, or in consultation with, a hematologist or oncologist

AND

3 - Patient has a history of failure, contraindication, or intolerance to Neupogen and Nivestym

| Product Name: Neulasta, Neulasta Onpro, Ziextenzo | |
|---|--|
| Diagnosis | Hematopoietic Syndrome of Acute Radiation Syndrome |
| Approval Length | 3 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient has been acutely exposed to myelosuppressive doses of radiation

AND

2 - Prescribed by, or in consultation with, a hematologist or oncologist

AND

3 - Patient has a history of failure, contraindication, or intolerance to Fulphila, Udenyca, and Nyvepria

Combination Basal Insulin/GLP-1 Receptor Agonist

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99510 | Combination Basal Insulin/GLP-1 Receptor Agonist |
|----------|--|
|----------|--|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Soliqua | |
|--|--------------|
| Approval Length | 12 month(s) |
| Guideline Type | Step Therapy |
| | |
| Approval Criteria | |
| 1 - Inadequately controlled on BOTH of the following | |
| GLP-1 (glucagon-like peptide-1) receptor agonist [e.g. Adlyxin (lixisenatide), Trulicity | |

 GLP-1 (glucagon-like peptide-1) receptor agonist [e.g. Adlyxin (lixisenatide), Trulicity (dulaglutide), Victoza (liraglutide), Bydureon (exenatide extended-release), Byetta (exenatide)] • Basal insulin (e.g. insulin glargine, insulin degludec, insulin detemir)

| Product Name: Xultophy | |
|---|---|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| Approval Criteria 1 - Diagnosis of type 2 diabetes mellitus | |
| | AND |
| 2 - Inadequately controlled on BOTH of the following | |
| GLP-1 (glucagon-like peptide-1) receptor agonist [e.g. Adlyxin (lixisenatide), Trulicity (dulaglutide), Victoza (liraglutide), Bydureon (exenatide extended-release), Byetta (exenatide)] Basal insulin (e.g. insulin glargine, insulin degludec, insulin detemir) | |
| AND | |
| 3 - History of failure, in | tolerance, or contraindication to Soliqua |

| Product Name: Xultophy | |
|-----------------------------|--|
| Approval Length 12 month(s) | |
| Reauthorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Xultophy therapy

2. Revision History

| Date | Notes |
|-----------|-------------------------------------|
| 5/24/2021 | Arizona Medicaid 7.1 Implementation |

Combination DPP-4/SGLT-2 Inhibitors (Glyxambi, Qtern, Steglujan)

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-114543 Combination DPP-4/SGLT-2 Inhibitors (Glyxambi, Qtern, Steglujan)

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Glyxambi, Qtern, Steglujan | | |
|---|---------------------|--|
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of type 2 diabetes mellitus | | |
| | | |
| | AND | |
| | | |

| 2 - History of failure, intolerance, or contraindication to metformin at a minimum dose of 1500 milligrams (mg) daily for 90 days |
|--|
| AND |
| 3 - History and failure, intolerance, or contraindication to ALL of the following: Janumet Janumet XR Januvia Jentadueto Kombiglyze XR Onglyza Tradjenta Trijardy XR |

2. Revision History

| Date | Notes |
|-----------|---------------------------------------|
| 9/26/2022 | Combined all criteria, Glyxambi is NP |

Compounds and Bulk Powders

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99565 Compounds and Bulk Powders |
|-------------------------------------|
|-------------------------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Requests for Compounds or Bulk Powders | | |
|--|----------------|--|
| Approval Length | 2 month(s) | |
| Guideline Type | Administrative | |
| | | |
| Approval Criteria | | |
| 1 - One of the following: | | |
| 1.1 The compound is an antibiotic. | | |

OR

1.2 Each active ingredient in the compounded drug is a covered medication

AND

2 - ONE of the following:

2.1 Each active ingredient in the compounded drug is to be administered for an FDA (Food and Drug Administration)-approved indication

OR

2.2 The use of each active ingredient in the compounded drug is supported by information from ONE of the following appropriate compendia of current literature:

- American Hospital Formulary Service Drug Information
- National Comprehensive Cancer Network Drugs and Biologics Compendium
- Thomson Micromedex DrugDex
- Clinical pharmacology
- United States Pharmacopoeia-National Formulary (USP-NF)

AND

3 - If a drug included in the compound requires prior authorization and/or step therapy, all drug specific clinical criteria must also be met

AND

4 - The compounded drug must not include any ingredient that has been withdrawn or removed from the market due to safety reasons.

AND

5 - ONE of the following:

5.1 A unique vehicle is required for topically administered compounds

OR

5.2 A unique dosage form is required for a commercially available product due to patient's age, weight, or inability to take a solid dosage form

OR

5.3 A unique formulation is required for a commercially available product due to an allergy or intolerance to an inactive ingredient in the commercially available product

OR

5.4 There is a shortage of the commercially available product per the FDA Drug Shortage database or the ASHP Current Drug Shortages tracking log

AND

6 - Coverage for compounds and bulk powders will NOT be approved for any of the following:

6.1 For topical compound preparations (e.g. creams, ointments, lotions, or gels to be applied to the skin for transdermal, transcutaneous, or any other topical route), requested compound contains any FDA approved ingredient that is not FDA approved for TOPICAL use (see Table 1 in Background section)

OR

6.2 If the requested compound contains topical fluticasone, topical fluticasone will NOT be approved unless both of the following are met:

6.2.1 Topical fluticasone is intended to treat a dermatologic condition (scar treatments are considered cosmetic and will not be covered)

AND

6.2.2 Patient has a contraindication to all commercially available topical fluticasone formulations

OR

6.3 Requested compound contains any ingredients when used for cosmetic purposes (see Table 2 in Background section)

OR

6.4 Requested compound contains any ingredient(s) which are on the FDA's Do Not Compound List (see Table 3 in Background section)

2. Background

Benefit/Coverage/Program Information

Table 1: Example topical compound preparations that contain any FDA approved ingredient that are not FDA approved for TOPICAL use, including but NOT LIMITED TO the following:

- (1) Ketamine
- (2) Gabapentin
- (3) Flurbiprofen (topical ophthalmic use not included)
- (4) Ketoprofen
- (5) Morphine
- (6) Nabumetone
- (7) Oxycodone
- (8) Cyclobenzaprine
- (9) Baclofen

| (10) | Tramadol |
|------|--------------------------------|
| (11) | Hydrocodone |
| (12) | Meloxicam |
| (13) | Amitriptyline |
| (14) | Pentoxifylline |
| (15) | Orphenadrine |
| (16) | Piroxicam |
| (17) | Levocetirizine |
| (18) | Amantadine |
| (19) | Oxytocin |
| (20) | Sumatriptan |
| (21) | Chorionic gonadotropin (human) |
| (22) | Clomipramine |
| (23) | Dexamethasone |
| (24) | Hydromorphone |
| (25) | Methadone |
| (26) | Papaverine |
| (27) | Mefenamic acid |
| (28) | Promethazine |
| (29) | Succimer DMSA |
| (30) | Tizanidine |
| (31) | Apomorphine |
| (32) | Carbamazepine |
| | |

- (33) Ketorolac
- (34) Dimercaptopropane-sulfonate
- (35) Dimercaptosuccinic acid
- (36) Duloxetine
- (37) Fluoxetine
- (38) Bromfenac (topical ophthalmic use not included)
- (39) Nepafenac (topical ophthalmic use not included)

Table 2: Example compounds that contain ingredients for cosmetic purposes:

- (1) Hydroquinone
- (2) Acetyl hexapeptide-8
- (3) Tocopheryl Acid Succinate
- (4) PracaSil TM-Plus
- (5) Chrysaderm Day Cream
- (6) Chrysaderm Night Cream
- (7) PCCA Spira-Wash
- (8) Lipopen Ultra
- (9) Versapro
- (10) Fluticasone
- (11) Mometasone
- (12) Halobetasol
- (13) Betamethasone
- (14) Clobetasol
- (15) Triamcinolone

| (16) | Minoxidil | |
|---|---|--|
| (17) | Tretinoin | |
| (18) | Dexamethasone | |
| (19) | Spironolactone | |
| (20) | Cycloserine | |
| (21) | Tamoxifen | |
| (22) | Sermorelin | |
| (23) | Mederma Cream | |
| (24) | PCCA Cosmetic HRT Base | |
| (25) | Sanare Scar Therapy Cream | |
| (26) | Scarcin Cream | |
| (27) | Apothederm | |
| (28) | Stera Cream | |
| (29) | Copasil | |
| (30) | Collagenase | |
| (31) | Arbutin Alpha | |
| (32) | Nourisil | |
| (33) | Freedom Cepapro | |
| (34) | Freedom Silomac Andydrous | |
| (35) | Retinaldehyde | |
| (36) | Apothederm | |
| Table 3: Example ingredients on the FDA's Do Not Compound List: | | |
| (1) 2 | (1) 3 3' 4' 5 totrachlorosaliovlanilida | |

(1) 3,3',4',5-tetrachlorosalicylanilide

- (2) Adenosine phosphate
- (3) Adrenal cortex
- (4) Alatrofloxacin mesylate
- (5) Aminopyrine
- (6) Astemizole
- (7) Azaribine
- (8) Benoxaprofen
- (9) Bithionol
- (10) Camphorated oil
- (11) Carbetapentane citrate
- (12) Casein, iodinated
- (13) Cerivastatin sodium
- (14) Chlormadinone acetate
- (15) Chloroform
- (16) Cisapride
- (17) Defenfluramine hydrochloride
- (18) Diamthazole dihydrochloride
- (19) Dibromsalan
- (20) Dihydrostreptomycin sulfate
- (21) Dipyrone
- (22) Encainide hydrochloride
- (23) Etretinate
- (24) Fenfluramine hydrochloride

- (25) Flosequinan
- (26) Glycerol, iodinated
- (27) Grepafloxacin
- (28) Mepazine
- (29) Metabromsalan
- (30) Methapyrilene
- (31) Methopholine
- (32) Methoxyflurane
- (33) Mibefradil dihydrochloride
- (34) Nomifensine maleate
- (35) Novobiocin sodium
- (36) Oxyphenisatin acetate
- (37) Oxyphenisatin
- (38) Pemoline
- (39) Pergolide mesylate
- (40) Phenacetin
- (41) Phenformin hydrochloride
- (42) Phenylpropanolamine
- (43) Pipamazine
- (44) Potassium arsenite
- (45) Propoxyphene
- (46) Rapacuronium bromide
- (47) Rofecoxib

- (48) Sibutramine hydrochloride
- (49) Sparteine sulfate
- (50) Sulfadimethoxine
- (51) Sweet spirits of nitre
- (52) Tegaserod maleate
- (53) Temafloxacin hydrochloride
- (54) Terfenadine
- (55) Ticrynafen
- (56) Tribromsalan
- (57) Trichloroethane
- (58) Troglitazone
- (59) Trovafloxacin mesylate:
- (60) Urethane
- (61) Valdecoxib
- (62) Zomepirac sodium

3. Revision History

| Date | Notes |
|----------|------------------|
| 7/7/2021 | Update guideline |

Constipation Agents - AZM

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-107421 Constipation Agents - AZM

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 5/25/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Amitiza, Brand lubiprostone | | |
|---|-----------------------|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - One of the following: | | |
| 1.1 ONE of the following diagnoses: | | |
| Opioid-induced constipation in an adult with chronic, non-cancer pain | | |

• Opioid-induced constipation in patients with chronic pain related to prior cancer or its treatment who do not require frequent (e.g., weekly) opioid dosage escalation

• Chronic idiopathic constipation

OR

1.2 Both of the following:

- Diagnosis of irritable bowel syndrome with constipation
- Patient was female at birth

AND

2 - BOTH of the following:

2.1 Trial and failure, contraindication, or intolerance to an osmotic laxative e.g., (lactulose, polyethylene glycol, sorbitol)

AND

2.2 Trial and failure, contraindication, or intolerance to ONE of the following:

- Bulk Forming Laxatives (e.g., psyllium, fiber)
- Stimulant Laxatives (e.g., bisacodyl, senna)

| Product Name: Ibsrela | |
|-----------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of irritable bowel syndrome with constipation

AND

2 - History of failure, contraindication or intolerance to BOTH of the following:

- Lactulose
- Polyethylene glycol (Miralax)

AND

3 - History of failure, contraindication or intolerance to ONE of the following:

- Amitiza
- Linzess
- Trulance

| Product Name: Linzess | |
|-----------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- **1** ONE of the following diagnoses:
 - Chronic idiopathic constipation
 - Irritable bowel syndrome with constipation

AND

2 - Patient is greater than or equal to 18 years of age

AND

3 - Both of the following:

3.1 Trial and failure, contraindication, or intolerance to an osmotic laxative e.g., (lactulose, polyethylene glycol, sorbitol)

AND

3.2 Trial and failure, contraindication, or intolerance to ONE of the following:

- Bulk Forming Laxatives (e.g., psyllium, fiber)
- Stimulant Laxatives (e.g., bisacodyl, senna)

| Product Name: Trulance | | |
|------------------------|-----------------------|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Ammanual Critoria | | |

Approval Criteria

1 - ONE of the following diagnoses:

- Chronic idiopathic constipation
- Irritable bowel syndrome with constipation

AND

2 - Patient is greater than or equal to 18 years of age

| Product Name: Motegrity | | |
|-------------------------|-----------------------|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Diagnosis of chronic idiopathic constipation

AND

2 - Both of the following

2.1 History of failure, contraindication or intolerance to BOTH of the following:

- Lactulose
- Polyethylene glycol (Miralax)

AND

2.2 History of failure, contraindication, or intolerance to BOTH of the following:

- Linzess
- Amitiza

| Product Name: Movantik | |
|------------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - ONE of the following diagnoses:

- Opioid-induced constipation in patients being treated for chronic, non-cancer pain
- Opioid-induced constipation in patients with chronic pain related to prior cancer or its treatment who do not require frequent (e.g., weekly) opioid dosage escalation

| Product Name: Symproic | |
|------------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - ONE of the following diagnoses:

- Opioid-induced constipation in patients being treated for chronic, non-cancer pain
- Opioid-induced constipation in patients with chronic pain related to prior cancer or its treatment who do not require frequent (e.g., weekly) opioid dosage escalation

AND

2 - History of failure, contraindication or intolerance to BOTH of the following:

- Lactulose
- Polyethylene glycol (Miralax)

AND

3 - History of failure, contraindication or intolerance to Movantik

| Product Name: Zelnorm | | |
|---|-----------------------|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| Approval Criteria | | |
| 1 - Diagnosis of irritable bowel syndrome with constipation | | |
| | AND | |
| 2 - Patient was female at birth | | |
| AND | | |

3 - History of failure, contraindication or intolerance to BOTH of the following:

- Lactulose
- Polyethylene glycol (Miralax)

AND

4 - History of failure, contraindication or intolerance to ONE of the following:

- Amitiza
- Linzess
- Trulance

| Product Name: Brand Amitiza, generic lubiprostone, Ibsrela, Linzess, Motegrity, Movantik, Symproic, Trulance, Zelnorm | | |
|---|---------------------|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |
| | | |

Approval Criteria

1 - Documentation of positive clinical response to therapy

2. Revision History

| Date | Notes |
|-----------|-------------------------|
| 5/23/2022 | Added Ibsrela as target |

Copper Chelating Agents

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99604 | Copper Chelating Agents |
|--------------|---|
| Formulary | Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) |
| Formulary No | ote |

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Depen Titratab, generic penicillamine tablets | |
|---|------------------------------------|
| Diagnosis | Severe active rheumatoid arthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of severe active rheumatoid arthritis

| Product Name: Brand Depen Titratab, generic penicillamine tablets | |
|---|------------------------------------|
| Diagnosis | Severe active rheumatoid arthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Depen Titratabs therapy

| Product Name: Brand Depen Titratab, generic penicillamine tablets | |
|---|--|
| Diagnosis | Wilson's disease (i.e., hepatolenticular degeneration), Cystinuria |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | • |

Approval Criteria

- **1** Patient has ONE of the following diagnoses:
 - Diagnosis of Wilson's disease (i.e., hepatolenticular degeneration) Diagnosis of Cystinuria •
 - •

| Product Name: Brand Cuprimine, generic penicillamine capsules | |
|---|--|
| Diagnosis | Wilson's disease (i.e., hepatolenticular degeneration), Cystinuria, Severe active rheumatoid arthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- **1** Patient has ONE of the following diagnoses:
 - Wilson's disease (i.e., hepatolenticular degeneration) •
 - Cystinuria •

• Severe active rheumatoid arthritis

AND

2 - History of failure or intolerance to Depen (penicillamine)

| Product Name: Brand Cuprimine, generic penicillamine capsules | |
|---|--|
| Diagnosis | Wilson's disease (i.e., hepatolenticular degeneration), Cystinuria, Severe active rheumatoid arthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Cuprimine (penicillamine) therapy

| Product Name: Brand Syprine, generic trientine, generic Clovique | |
|--|--|
| Diagnosis | Wilson's disease (i.e., hepatolenticular degeneration) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of Wilson's disease (i.e., hepatolenticular degeneration)

AND

2 - History of failure, contraindication, or intolerance to Depen (penicillamine) or Cuprimine (penicillamine)

Product Name: Brand Syprine, generic trientine, generic Clovique

| Diagnosis | Wilson's disease (i.e., hepatolenticular degeneration) |
|-----------------|--|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Syprine (trientine) therapy

2. Revision History

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona SP to Medicaid Arizona SP for 7/1 eff |

Corlanor

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99441 Corlanor

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Corlanor | | |
|------------------------|-----------------------|--|
| Diagnosis | Chronic Heart Failure | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Worsening heart failure in a diagnosis of stable, symptomatic chronic (e.g. New York Heart Association (NYHA) class II, III or IV) heart failure

AND 2 - Patient has a left ventricular ejection fraction (EF) less than or equal to 35% AND 3 - The patient is in sinus rhythm AND 4 - Patient has a resting heart rate greater than or equal to 70 beats per minute AND 5 - ONE of the following: 5.1 Patient is on maximum tolerated doses of beta blockers (e.g., carvedilol, metoprolol succinate, bisoprolol) OR

5.2 Patient has a contraindication or intolerance to beta-blocker therapy

| Product Name: Corlanor | | |
|------------------------|---|--|
| Diagnosis | Heart Failure due to Dilated Cardiomyopathy (DCM) | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |

1 - Diagnosis of stable symptomatic heart failure due to dilated cardiomyopathy (DCM)

| | AND |
|--|-----|
| 2 - Patient is in sinus rhythm | |
| | AND |
| 3 - Patient has an elevated heart rate | |

| Product Name: Corlanor | | |
|------------------------|--|--|
| Diagnosis | Chronic Heart Failure, Heart Failure due to Dilated Cardiomyopathy (DCM) | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |

1 - Documentation of positive clinical response to Corlanor therapy

2. Revision History

| Date | Notes |
|-----------|--|
| 3/10/2021 | Bulk Copy guidelines starting with B and C from C&S Arizona to Ariz ona Medicaid |

Cosentyx (secukinumab)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-114520 | Cosentyx (secuki | numab) |
|-----------|------------------|--------|
|-----------|------------------|--------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Cosentyx | |
|------------------------|-----------------------|
| Diagnosis | Plaque Psoriasis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - One of the following:

1.1 All of the following:

1.1.1 Diagnosis of moderate to severe plaque psoriasis

AND

1.1.2 Greater than or equal to 3 percent body surface area involvement, palmoplantar, facial, or genital involvement, or severe scalp psoriasis

AND

1.1.3 Both of the following:

1.1.3.1 History of failure to TWO of the following topical therapies, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial):*

- Corticosteroids (e.g., betamethasone, clobetasol, desonide)
- Vitamin D analogs (e.g., calcitriol, calcipotriene)
- Tazarotene
- Calcineurin inhibitors (e.g., tacrolimus, pimecrolimus)
- Anthralin
- Coal tar

AND

1.1.3.2 History of failure to a 3 month trial of methotrexate at the maximally indicated dose within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document date, and duration of trial)

AND

1.1.4 Patient is not receiving Cosentyx in combination with ANY of the following:

- Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Taltz (ixekizumab), Orencia (abatacept)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

1.1.5 History of failure, contraindication, or intolerance to ALL of the following: Humira (adalimumab) • Enbrel (etanercept) • Otezla (apremilast) AND 1.1.6 Prescribed by or in consultation with a dermatologist OR **1.2** All of the following: **1.2.1** Patient is currently on Cosentyx therapy as documented by claims history or medical records (document date, and duration of therapy) AND **1.2.2** Diagnosis of moderate to severe plaque psoriasis AND **1.2.3** Patient is not receiving Cosentyx in combination with ANY of the following: Biologic DMARD [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi • (golimumab), Taltz (ixekizumab), Orencia (abatacept)] Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)] ٠ Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)] • AND 1.2.4 Prescribed by or in consultation with a dermatologist AND

| 2 - Patient is 6 years of | age or older |
|----------------------------------|---|
| Notes | *Note: Claims history may be used in conjunction as documentation of drug, date, and duration of trials |

| Product Name: Cosentyx | |
|------------------------|---------------------|
| Diagnosis | Plaque Psoriasis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Cosentyx therapy

AND

2 - Patient is not receiving Cosentyx in combination with ANY of the following:

- Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Taltz (ixekizumab), Orencia (abatacept)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with a dermatologist

| Product Name: Cosentyx | |
|------------------------|------------------------|
| Diagnosis | Ankylosing Spondylitis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - One of the following:

1.1 All of the following:

1.1.1 Diagnosis of active ankylosing spondylitis

AND

1.1.2 History of failure to two NSAIDs (non-steroidal anti-inflammatory drugs) (e.g., ibuprofen, naproxen) at maximally indicated doses, each used for at least 4 weeks within the last 3 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trials)*

AND

1.1.3 Patient is not receiving Cosentyx in combination with ANY of the following:

- Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Taltz (ixekizumab), Orencia (abatacept)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

1.1.4 History of failure, contraindication, or intolerance to TWO of the following:

- Humira (adalimumab)
- Enbrel (etanercept)
- Xeljanz (tofacitinib)

AND

1.1.5 Prescribed by or in consultation with a rheumatologist

OR

1.2 All of the following:

| 1.2.1 Patient is currently on Cosentyx therapy as documented by claims history or medical records (document date, and duration of therapy) | |
|---|---|
| | AND |
| 1.2.2 Diagnosis of ac | tive ankylosing spondylitis |
| | AND |
| 1.2.3 Patient is not re | eceiving Cosentyx in combination with ANY of the following: |
| Biologic DMARD [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Taltz (ixekizumab), Orencia (abatacept)] Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)] Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)] | |
| AND | |
| 1.2.4 Prescribed by or in consultation with a rheumatologist | |
| Notes | *Note: Claims history may be used in conjunction as documentation of drug, date, and duration of trials |

| Product Name: Cosentyx | |
|------------------------|------------------------|
| Diagnosis | Ankylosing Spondylitis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Cosentyx therapy

AND

2 - Patient is not receiving Cosentyx in combination with ANY of the following:

- Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Taltz (ixekizumab), Orencia (abatacept)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with a rheumatologist

| Product Name: Cosentyx | |
|------------------------|-----------------------|
| Diagnosis | Psoriatic Arthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - One of the following:

- 1.1 All of the following:
- **1.1.1** Diagnosis of active psoriatic arthritis

AND

1.1.2 History of failure to a 3 month trial of methotrexate at the maximally indicated dose within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document date, and duration of trial)

AND

1.1.3 Patient is not receiving Cosentyx in combination with ANY of the following:

• Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Taltz (ixekizumab), Orencia (abatacept)] • Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]

• Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

1.1.4 History of failure, contraindication, or intolerance to THREE of the following:

- Humira (adalimumab)
- Enbrel (etanercept)
- Otezla (apremilast)
- Xeljanz (tofacitinib)

AND

1.1.5 Prescribed by or in consultation with ONE of the following:

- Rheumatologist
- Dermatologist

OR

1.2 All of the following:

1.2.1 Patient is currently on Cosentyx therapy as documented by claims history or medical records (document date, and duration of therapy)

AND

1.2.2 Diagnosis of active psoriatic arthritis

AND

1.2.3 Patient is not receiving Cosentyx in combination with ANY of the following:

- Biologic DMARD [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Taltz (ixekizumab), Orencia (abatacept)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

1.2.4 Prescribed by or in consultation with ONE of the following:

- Rheumatologist
- Dermatologist

AND

2 - Patient is 2 years of age or older

| Product Name: Cosentyx | |
|------------------------|---------------------|
| Diagnosis | Psoriatic Arthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Cosentyx therapy

AND

- **2** Patient is not receiving Cosentyx in combination with ANY of the following:
 - Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Taltz (ixekizumab), Orencia (abatacept)]
 - Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
 - Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

- 3 Prescribed by or in consultation with ONE of the following:
 - Rheumatologist

Dermatologist

| Product Name: Cosentyx | |
|------------------------|--|
| Diagnosis | Non-radiographic axial spondyloarthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- 1 One of the following:
- **1.1** All of the following:

1.1.1 Diagnosis of active non-radiographic axial spondyloarthritis

AND

1.1.2 History of failure to two NSAIDs (non-steroidal anti-inflammatory drugs) (e.g., ibuprofen, naproxen) at maximally indicated doses, each used for at least 4 weeks, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trials)*

AND

1.1.3 Patient is not receiving Cosentyx in combination with ANY of the following:

- Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Taltz (ixekizumab), Orencia (abatacept)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]

AND

1.1.4 Prescribed by or in consultation with a rheumatologist

OR

1.2 All of the following:

1.2.1 Patient is currently on Cosentyx therapy as documented by claims history or medical records (document date, and duration of therapy)

AND

1.2.2 Diagnosis of active non-radiographic axial spondyloarthritis

AND

1.2.3 Patient is not receiving Cosentyx in combination with ANY of the following:

- Biologic DMARD [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Taltz (ixekizumab), Orencia (abatacept)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]

AND

1.2.4 Prescribed by or in consultation with a rheumatologist

| *Note: Claims history may be used in conjunction as documentation of |
|--|
| drug, date, and duration of trials |

| Product Name: Cosentyx | |
|------------------------|--|
| Diagnosis | Non-radiographic axial spondyloarthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Cosentyx therapy

AND

2 - Patient is not receiving Cosentyx in combination with ANY of the following:

- Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Taltz (ixekizumab), Orencia (abatacept)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]

AND

3 - Prescribed by or in consultation with a rheumatologist

| Product Name: Cosent | ух |
|----------------------------------|--|
| Diagnosis | Enthesitis-Related Arthritis (ERA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| Approval Criteria | |
| 1 - Diagnosis of active | enthesitis-related arthritis |
| | AND |
| 2 - Patient is 4 years of | f age or older |
| | AND |
| 3 - Prescribed by or in | consultation with a rheumatologist |
| | AND |
| 4 - Paid claims or subn | nission of medical records (e.g., chart notes) confirming trial and failure, |

contraindication, or intolerance to TWO preferred non-steroidal anti-inflammatory drugs (NSAIDs) (e.g., ibuprofen, naproxen)

| Product Name: Cosentyx | |
|------------------------|------------------------------------|
| Diagnosis | Enthesitis-Related Arthritis (ERA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Cosentyx therapy

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

3 - Documentation of a positive clinical response to therapy as evidenced by at least one of the following:

- Reduction in the total active (swollen and tender) joint count from baseline
- Improvement in symptoms (e.g., pain, stiffness, inflammation) from baseline

2. Revision History

| Date | Notes |
|-----------|--|
| 9/26/2022 | Updated criteria: age requirements and indications |

Cough and Cold Products

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-104889 Cough and Cold Products

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 3/28/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

Product Name: Hydromet, generic Tussigon, Z-Tuss AC, Tuzistra XR, Tussicaps, generic Tussionex, M-END PE, Poly-Tussin AC, Capcof, Pro-Red AC, Histex-AC, Maxi-Tuss, generic promethazine w/codeine, generic promethazine-phenylephrine-codeine, Rydex, Mar-Cof BP/Mar-Cof GG, Ninjacof-XG, Coditussin AC/Coditussin DAC, generic guaifenesin-codeine, generic pseudoephedrine w/codeine-guaifenesin, Tuxarin ER

| Diagnosis | Under the Age of 18 Years for Cough and Cold Products |
|-----------------|---|
| Approval Length | 30 Day(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Prescriber attests they are aware of Food and Drug Administration (FDA) labeled

contraindications regarding use of opioid containing cough and cold products in patients less than 18 years of age and feels the treatment with the requested product is medically necessary (Document rationale for use)

AND

2 - Patient does not have a comorbid condition that may impact respiratory depression (e.g., asthma or other chronic lung disease, sleep apnea, body mass index greater than 30)

AND

3 - Patient has tried and failed at least one non-opioid containing cough and cold remedy

Product Name: Hydromet, generic Tussigon, Z-Tuss AC, Tuzistra XR, Tussicaps, generic Tussionex, M-END PE, Poly-Tussin AC, Capcof, Pro-Red AC, Histex-AC, Maxi-Tuss, generic promethazine w/codeine, generic promethazine-phenylephrine-codeine, Rydex, Mar-Cof BP/Mar-Cof GG, Ninjacof-XG, Coditussin AC/Coditussin DAC, generic guaifenesin-codeine, generic pseudoephedrine w/codeine-guaifenesin, Tuxarin ER

| Diagnosis | Quantity Limit |
|-----------------|-----------------|
| Approval Length | 30 Day(s) |
| Guideline Type | Quantity Limit* |

Approval Criteria

1 - Prescriber attests that a larger quantity is medically necessary

AND

2 - The requested dose is within the Food and Drug Administration (FDA) maximum dose per day, where an FDA maximum dose per day exists (See table in background section)

| Notes | *Authorization will be issued for up to 30 days. The authorization shou |
|-------|---|
| | Id be entered for the quantity requested. |

2. Background

| C Recommended Opioid Maximu | Im Morphine Milligram Equivalents per Day* |
|-----------------------------|--|
| Active Ingredient | FDA Label Max Daily Doses |
| | |
| lorphine | None |
| ydromorphone | None |
| ydrocodone | None |
| apentadol | 600mg IR products |
| xymorphone | None |
| xycodone | None |
| odeine | 360mg |
| entazocine | None |
| amadol | 400mg IR products |
| eperidine | 600mg |
| utorphanol nasal | None |
| bium | 4 suppositories/day |
| | Deodorized tincture: 24mg/day Camphorated tincture: 16mg/da |
| cetaminophen | 4g/day |
| spirin | 2080mg/day |
| uprofen | 3200mg/day |
| enzhydrocodone** | None |

3. Revision History

| Date | Notes |
|-----------|---|
| 3/28/2022 | Updated product list, no changes to criteria. |

Coverage of Off-Label Non-FDA Approved Indications

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99735 | Coverage of Off-Label Non-FDA Approved Indications |
|----------|--|
|----------|--|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: A drug (non-anti-cancer chemotherapeutic regimen) used for an off-label indication or non-FDA approved indication | | |
|---|---------------------------------|--|
| Diagnosis | Off-label non-cancer indication | |
| Approval Length | 12 month(s) | |
| Guideline Type | be Administrative | |

Approval Criteria

1 - The use of this drug is supported by information from ONE of the following appropriate compendia of current literature:

• Food and Drug Administration (FDA) approved indications and limits

- Published practice guidelines and treatment protocols
- Comparative data evaluating the efficacy, type and frequency of side effects and potential drug interactions among alternative products as well as the risks, benefits and potential member outcomes
- Drug Facts and Comparisons
- American Hospital Formulary Service Drug Information
- United States Pharmacopeia Drug Information
- DRUGDEX Information System
- UpToDate
- MicroMedex
- Peer-reviewed medical literature, including randomized clinical trials, outcomes, research data and pharmacoeconomic studies
- Other drug reference resources

| Off-label use may be reviewed for medical necessity and denied as su ch if the off-label criteria are not met. Please refer to drug specific PA guideline for off-label criteria if available. |
|--|

| Product Name: A drug or biological in an anti-cancer chemotherapeutic regimen | | |
|---|-----------------------------|--|
| Diagnosis | Off-label cancer indication | |
| Approval Length | 12 month(s) | |
| Guideline Type Administrative | | |

Approval Criteria

- 1 One of the following:
- **1.1** Diagnosis is supported as a use in AHFS DI [2]

OR

1.2 Diagnosis is supported as a use in the National Comprehesive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B (see NCCN Categories of Evidence and Consensus table in Background section) [2, A]

OR

1.3 Diagnosis is supported in the FDA Uses/Non-FDA Uses section in DRUGDEX Evaluation with a Strength of Recommendation rating of Class I, Class IIa, or Class IIb (see DRUGDEX Strength of Recommendation table in Background section) [2] OR

1.4 Diagnosis is supported as an indication in Clinical Pharmacology [2]

OR

1.5 Off-label use is supported in one of the published, peer-reviewed medical literature listed below: [2, B]

- American Journal of Medicine
- Annals of Internal Medicine
- Annals of Oncology
- Annals of Surgical Oncology
- Biology of Blood and Marrow Transplantation
- Blood
- Bone Marrow Transplantation
- British Journal of Cancer
- British Journal of Hematology
- British Medical Journal
- Cancer
- Clinical Cancer Research
- Drugs
- European Journal of Cancer (formerly the European Journal of Cancer and Clinical Oncology)
- Gynecologic Oncology
- International Journal of Radiation, Oncology, Biology, and Physics
- The Journal of the American Medical Association
- Journal of Clinical Oncology
- Journal of the National Cancer Institute
- Journal of the National Comprehensive Cancer Network (NCCN)
- Journal of Urology
- Lancet
- Lancet Oncology
- Leukemia
- The New England Journal of Medicine
- Radiation Oncology

OR

1.6 Diagnosis is supported as a use in Wolters Kluwer Lexi-Drugs rated as "Evidence Level A" with a "Strong" recommendation. (see Lexi-Drugs Strength of Recommendation table in Background section) [2, 4, 5]

| Off-label use may be reviewed for medical necessity and denied as su |
|--|
| ch if the off-label criteria are not met. Please refer to drug specific PA |
| guideline for off-label criteria if available. |

2. Background

| Clinical | Practice | Guidelines |
|----------|----------|------------|
|----------|----------|------------|

| Class | Recommendation | Description |
|------------------------|-------------------------------|---|
| Class I | Recommended | The given test or treatment has been proven useful, and should be performed or administered. |
| Class IIa | Recommended, In Most Cases | The given test or treatment is generally considered to be useful, and is indicated in most cases. |
| Class IIb | Recommended, in Some Cases | The given test or treatment may be useful, and is indicated in some, but not most, cases. |
| Class III | Not Recommended | The given test or treatment is not useful, and should be avoided |
| Class Indeterminate | Evidence Inconclusive | |
| s of Evidence and | d Concerneuro [A] | |

| | 2A | B | ased upon lower-level evidence, there is uniform NCCN |
|--------------------|----------|-----------------------------|---|
| | | | onsensus that the intervention is appropriate. |
| | 2B | | ased upon lower-level evidence, there is NCCN consensus that the intervention is appropriate. |
| | 3 | | ased upon any level of evidence, there is major NCCN sagreement that the intervention is appropriate. |
| | | | dation for Inclusion in Lexi-Drugs for Oncology Off- |
| Label Use and Leve | el of Ev | idence Sca | ale for Oncology Off-Label Use [5] |
| Strength of Recor | nmend | lation for In | clusion |
| Strong (for propo | | The | |
| off-label use) | | evidence | |
| | | persuasivel supports th | |
| | | off-label us | |
| | | (ie, Level of | |
| | | Evidence A |). |
| Equivocal (for | | The | |
| proposed off-labe | | evidence to | |
| use) | | support the off-label us | |
| | | is of | |
| | | uncertain | |
| | | clinical | |
| | | significance | |
| | | (ie, Level of Evidence B | |
| | | C). | |
| | | Ádditional | |
| | | studies may | / |
| | | be | |
| | | necessary further defined | |
| | | the role of | |
| | | this | |
| | | medication | |
| | | for the off- label use. | |
| | | | |
| Against proposed | | The | |
| label use | | evidence either | |
| | | advocates | |
| | | against the | |
| | | | |

| | off-label use or suggests a lack of support for the off-label use (independent of Level of Evidence). Additional studies are necessary to define the role of this medication for the off- label use. | | |
|--------------|---|--|--|
| Level of Evi | dence Scale for Oncology Off-Label Use Consistent evidence from well-performed randomized, controlled trials or overwhelming evidence of some other form (eg, results of the introduction of penicillin treatment) to support off-label use. Further research is unlikely to change confidence in the estimate of benefit. | | |
| В | Evidence from randomized, controlled trials with important limitations (eg, inconsistent results, methodologic flaws, indirect, imprecise); or very strong evidence of some other research design. Further research (if performed) is likely to have an impact on confidence in the estimate of benefit and risk and may change the estimate. | | |
| С | Evidence from observational studies (eg, retrospective case series/reports providing significant impact on patient care); unsystematic clinical experience; or potentially flawed randomized, controlled trials (eg, when limited options exist for condition). Any estimate of effect is uncertain. | | |
| G | Use has been substantiated by inclusion in at least one evidence- based or consensus-based clinical practice guideline. | | |
| | | | |

3. Endnotes

- A. NCCN Categories of Evidence and Consensus. Category 1: The recommendation is based on high-level evidence (i.e., high-powered randomized clinical trials or metaanalyses), and the NCCN Guideline Panel has reached uniform consensus that the recommendation is indicated. In this context, uniform means near unanimous positive support with some possible neutral positions. Category 2A: The recommendation is based on lower level evidence, but despite the absence of higher level studies, there is uniform consensus that the recommendation is appropriate. Lower level evidence is interpreted broadly, and runs the gamut from phase II to large cohort studies to case series to individual practitioner experience. Importantly, in many instances, the retrospective studies are derived from clinical experience of treating large numbers of patients at a member institution, so NCCN Guideline Panel Members have first-hand knowledge of the data. Inevitably, some recommendations must address clinical situations for which limited or no data exist. In these instances the congruence of experience-based judgments provides an informed if not confirmed direction for optimizing patient care. These recommendations carry the implicit recognition that they may be superseded as higher level evidence becomes available or as outcomes-based information becomes more prevalent. Category 2B: The recommendation is based on lower level evidence, and there is nonuniform consensus that the recommendation should be made. In these instances, because the evidence is not conclusive, institutions take different approaches to the management of a particular clinical scenario. This nonuniform consensus does not represent a major disagreement, rather it recognizes that given imperfect information, institutions may adopt different approaches. A Category 2B designation should signal to the user that more than one approach can be inferred from the existing data. Category 3: Including the recommendation has engendered a major disagreement among the NCCN Guideline Panel Members. The level of evidence is not pertinent in this category, because experts can disagree about the significance of high level trials. Several circumstances can cause major disagreements. For example, if substantial data exist about two interventions but they have never been directly compared in a randomized trial, adherents to one set of data may not accept the interpretation of the other side's results. Another situation resulting in a Category 3 designation is when experts disagree about how trial data can be generalized. An example of this is the recommendation for internal mammary node radiation in postmastectomy radiation therapy. One side believed that because the randomized studies included this modality, it must be included in the recommendation. The other side believed, based on the documented additional morbidity and the role of internal mammary radiation therapy in other studies, that this was not necessary. A Category 3 designation alerts users to a major interpretation issue in the data and directs them to the manuscript for an explanation of the controversy. [3]
- B. Abstracts (including meeting abstracts) are excluded from consideration. When evaluating peer-reviewed medical literature, the following (among other things) should be considered: 1) Whether the clinical characteristics of the beneficiary and the cancer are adequately represented in the published evidence 2) Whether the administered chemotherapy regimen is adequately represented in the published evidence. 3) Whether

the reported study outcomes represent clinically meaningful outcomes experienced by patients. 4) Whether the study is appropriate to address the clinical question. The following should be considered: a) Whether the experimental design, in light of the drugs and conditions under investigation, is appropriate to address the investigative question. (For example, in some clinical studies, it may be unnecessary or not feasible to use randomization, double blind trials, placebos, or crossover.); b) That non-randomized clinical trials with a significant number of subjects may be a basis for supportive clinical evidence for determining accepted uses of drugs; and c) That case reports are generally considered uncontrolled and anecdotal information and do not provide adequate supportive clinical evidence for determining accepted uses of drugs. [2]

4. References

- Center for Medicaid & Medicare Services. Medicare Prescription Drug Benefit Manual. Chapter 6 – Part D Drugs and Formulary Requirements. Section 10.6. Available at: https://www.cms.gov/Medicare/Prescription-Drug-Coverage/PrescriptionDrugCovContra/Downloads/Part-D-Benefits-Manual-Chapter-6.pdf. Accessed September 9, 2020.
- Center for Medicaid & Medicare Services. Medicare Benefit Policy Manual. Chapter 15 -Covered Medical and Other Health Services. Section 50.4.5. Available at: https://www.cms.gov/Regulations-and-Guidance/Guidance/Manuals/downloads/bp102c15.pdf. Accessed September 9, 2020.
- National Comprehensive Cancer Network Categories of Evidence and Consensus. Available at: https://www.nccn.org/professionals/physician_gls/categories_of_consensus.aspx. Accessed September 9, 2020.
- 4. Center for Medicaid & Medicare Services. Medicare Benefit Policy Manual. Wolters Kluwer Clinical Drug Information Lexi-Drugs Compendium Revision Request - CAG-00443O. Available at: https://www.cms.gov/medicare-coveragedatabase/details/medicare-coverage-document-details.aspx?MCDId=31#decision. Accessed September 9, 2020.
- Wolters Kluwer Clinical Drug Information's Request for CMS evaluation of Lexi-Drugs as a compendium for use in the determination of medically-accepted indications of drugs/biologicals used off-label in anti-cancer chemotherapeutic regimens. Available at: https://www.cms.gov/Medicare/Coverage/CoverageGenInfo/downloads/covdoc31.pdf. Accessed September 9, 2020.
- Micromedex Healthcare Series. Recommendation, Evidence, and Efficacy Ratings. https://www.micromedexsolutions.com/micromedex2/librarian/ssl/true/CS/6E0ED9/ND_P R/evidencexpert/ND_P/evidencexpert/DUPLICATIONSHIELDSYNC/8B9F5B/ND_PG/evi dencexpert/ND_B/evidencexpert/ND_AppProduct/evidencexpert/ND_T/evidencexpert/P FActionId/evidencexpert.IntermediateToDocumentLink?docId=3198&contentSetId=50. Accessed September 9, 2020.

5. Revision History

| Date | Notes |
|-----------|-------------------------------------|
| 5/18/2021 | Arizona Medicaid 7.1 Implementation |

Coverage of Off-Label Non-FDA Approved Indications

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99530 Coverage of Off-Label Non-FDA Approved Indications

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: A drug (non-anti-cancer chemotherapeutic regimen) used for an off-label indication or non-FDA approved indication | | |
|---|---------------------------------|--|
| Diagnosis | Off-label non-cancer indication | |
| Approval Length | 12 month(s) | |
| Guideline Type | Administrative | |

Approval Criteria

1 - The use of this drug is supported by information from ONE of the following appropriate compendia of current literature:

• Food and Drug Administration (FDA) approved indications and limits

- Published practice guidelines and treatment protocols
- Comparative data evaluating the efficacy, type and frequency of side effects and potential drug interactions among alternative products as well as the risks, benefits and potential member outcomes
- Drug Facts and Comparisons
- American Hospital Formulary Service Drug Information
- United States Pharmacopeia Drug Information
- DRUGDEX Information System
- UpToDate
- MicroMedex
- Peer-reviewed medical literature, including randomized clinical trials, outcomes, research data and pharmacoeconomic studies
- Other drug reference resources

| Off-label use may be reviewed for medical necessity and denied as su ch if the off-label criteria are not met. Please refer to drug specific PA guideline for off-label criteria if available. |
|--|

| Product Name: A drug or biological in an anti-cancer chemotherapeutic regimen | | |
|---|-----------------------------|--|
| Diagnosis | Off-label cancer indication | |
| Approval Length | 12 month(s) | |
| Guideline Type Administrative | | |

Approval Criteria

- 1 One of the following:
- **1.1** Diagnosis is supported as a use in AHFS DI [2]

OR

1.2 Diagnosis is supported as a use in the National Comprehesive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B (see NCCN Categories of Evidence and Consensus table in Background section) [2, A]

OR

1.3 Diagnosis is supported in the FDA Uses/Non-FDA Uses section in DRUGDEX Evaluation with a Strength of Recommendation rating of Class I, Class IIa, or Class IIb (see DRUGDEX Strength of Recommendation table in Background section) [2] OR

1.4 Diagnosis is supported as an indication in Clinical Pharmacology [2]

OR

1.5 Off-label use is supported in one of the published, peer-reviewed medical literature listed below: [2, B]

- American Journal of Medicine
- Annals of Internal Medicine
- Annals of Oncology
- Annals of Surgical Oncology
- Biology of Blood and Marrow Transplantation
- Blood
- Bone Marrow Transplantation
- British Journal of Cancer
- British Journal of Hematology
- British Medical Journal
- Cancer
- Clinical Cancer Research
- Drugs
- European Journal of Cancer (formerly the European Journal of Cancer and Clinical Oncology)
- Gynecologic Oncology
- International Journal of Radiation, Oncology, Biology, and Physics
- The Journal of the American Medical Association
- Journal of Clinical Oncology
- Journal of the National Cancer Institute
- Journal of the National Comprehensive Cancer Network (NCCN)
- Journal of Urology
- Lancet
- Lancet Oncology
- Leukemia
- The New England Journal of Medicine
- Radiation Oncology

OR

1.6 Diagnosis is supported as a use in Wolters Kluwer Lexi-Drugs rated as "Evidence Level A" with a "Strong" recommendation. (see Lexi-Drugs Strength of Recommendation table in Background section) [2, 4, 5]

| Notes | Off-label use may be reviewed for medical necessity and denied as su |
|-------|--|
| | ch if the off-label criteria are not met. Please refer to drug specific PA |
| | guideline for off-label criteria if available. |

2. Background

| | Class I | Recommended | DescriptionThe given test or treatment has been proven useful, and should be performed or |
|---|-----------------------|-------------------------------|---|
| C | <u></u> | | administered. |
| | Class IIa | Recommended, In Most Cases | The given test or treatment is generally considered to be useful, and is indicated in most cases. |
| | Class IIb | Recommended, in Some Cases | The given test or treatment may be useful, and is indicated in some, but not most, cases. |
| C | Class III | Not Recommended | The given test or treatment is not useful, and should be avoided |
| | Class ndeterminate | Evidence Inconclusive | |
| NCCN Categories of Evidence and Consensus [A] | | | |

| | 2A | | Based upon lower-level evidence, there is uniform NCCN onsensus that the intervention is appropriate. |
|--|----|--|---|
| | 2B | | Based upon lower-level evidence, there is NCCN consensus that the intervention is appropriate. |
| | 3 | | Based upon any level of evidence, there is major NCCN lisagreement that the intervention is appropriate. |
| | | lecommen | dation for Inclusion in Lexi-Drugs for Oncology Off- ale for Oncology Off-Label Use [5] |
| Strength of Reco | | | |
| Strong (for propo off-label use) | | The evidence persuasive supports th off-label us (ie, Level o Evidence A | e e f |
| Equivocal (for proposed off-lab use) | | The evidence to support the off-label us is of uncertain clinical significance (ie, Level o Evidence E C). Additional studies ma be necessary further defi the role of this medication for the off- label use. | e e f f, y to ne |
| Against propose label use | | The evidence either advocates against the | |

| | off-label use or suggests a lack of support for the off-label use (independent of Level of Evidence). Additional studies are necessary to define the role of this medication for the off- label use. |
|------------|---|
| Level of E | vidence Scale for Oncology Off-Label Use Consistent evidence from well-performed randomized, controlled trials or overwhelming evidence of some other form (eg, results of the |
| | introduction of penicillin treatment) to support off-label use. Further research is unlikely to change confidence in the estimate of benefit. |
| В | Evidence from randomized, controlled trials with important limitations (eg, inconsistent results, methodologic flaws, indirect, imprecise); or very strong evidence of some other research design. Further research (if performed) is likely to have an impact on confidence in the estimate of benefit and risk and may change the estimate. |
| C | Evidence from observational studies (eg, retrospective case series/reports providing significant impact on patient care); unsystematic clinical experience; or potentially flawed randomized, controlled trials (eg, when limited options exist for condition). Any estimate of effect is uncertain. |
| G | Use has been substantiated by inclusion in at least one evidence- based or consensus-based clinical practice guideline. |
| | |

3. Endnotes

- A. NCCN Categories of Evidence and Consensus. Category 1: The recommendation is based on high-level evidence (i.e., high-powered randomized clinical trials or metaanalyses), and the NCCN Guideline Panel has reached uniform consensus that the recommendation is indicated. In this context, uniform means near unanimous positive support with some possible neutral positions. Category 2A: The recommendation is based on lower level evidence, but despite the absence of higher level studies, there is uniform consensus that the recommendation is appropriate. Lower level evidence is interpreted broadly, and runs the gamut from phase II to large cohort studies to case series to individual practitioner experience. Importantly, in many instances, the retrospective studies are derived from clinical experience of treating large numbers of patients at a member institution, so NCCN Guideline Panel Members have first-hand knowledge of the data. Inevitably, some recommendations must address clinical situations for which limited or no data exist. In these instances the congruence of experience-based judgments provides an informed if not confirmed direction for optimizing patient care. These recommendations carry the implicit recognition that they may be superseded as higher level evidence becomes available or as outcomes-based information becomes more prevalent. Category 2B: The recommendation is based on lower level evidence, and there is nonuniform consensus that the recommendation should be made. In these instances, because the evidence is not conclusive, institutions take different approaches to the management of a particular clinical scenario. This nonuniform consensus does not represent a major disagreement, rather it recognizes that given imperfect information, institutions may adopt different approaches. A Category 2B designation should signal to the user that more than one approach can be inferred from the existing data. Category 3: Including the recommendation has engendered a major disagreement among the NCCN Guideline Panel Members. The level of evidence is not pertinent in this category, because experts can disagree about the significance of high level trials. Several circumstances can cause major disagreements. For example, if substantial data exist about two interventions but they have never been directly compared in a randomized trial, adherents to one set of data may not accept the interpretation of the other side's results. Another situation resulting in a Category 3 designation is when experts disagree about how trial data can be generalized. An example of this is the recommendation for internal mammary node radiation in postmastectomy radiation therapy. One side believed that because the randomized studies included this modality, it must be included in the recommendation. The other side believed, based on the documented additional morbidity and the role of internal mammary radiation therapy in other studies, that this was not necessary. A Category 3 designation alerts users to a major interpretation issue in the data and directs them to the manuscript for an explanation of the controversy. [3]
- B. Abstracts (including meeting abstracts) are excluded from consideration. When evaluating peer-reviewed medical literature, the following (among other things) should be considered: 1) Whether the clinical characteristics of the beneficiary and the cancer are adequately represented in the published evidence 2) Whether the administered chemotherapy regimen is adequately represented in the published evidence. 3) Whether

the reported study outcomes represent clinically meaningful outcomes experienced by patients. 4) Whether the study is appropriate to address the clinical question. The following should be considered: a) Whether the experimental design, in light of the drugs and conditions under investigation, is appropriate to address the investigative question. (For example, in some clinical studies, it may be unnecessary or not feasible to use randomization, double blind trials, placebos, or crossover.); b) That non-randomized clinical trials with a significant number of subjects may be a basis for supportive clinical evidence for determining accepted uses of drugs; and c) That case reports are generally considered uncontrolled and anecdotal information and do not provide adequate supportive clinical evidence for determining accepted uses of drugs. [2]

4. References

- Center for Medicaid & Medicare Services. Medicare Prescription Drug Benefit Manual. Chapter 6 – Part D Drugs and Formulary Requirements. Section 10.6. Available at: https://www.cms.gov/Medicare/Prescription-Drug-Coverage/PrescriptionDrugCovContra/Downloads/Part-D-Benefits-Manual-Chapter-6.pdf. Accessed September 9, 2020.
- Center for Medicaid & Medicare Services. Medicare Benefit Policy Manual. Chapter 15 -Covered Medical and Other Health Services. Section 50.4.5. Available at: https://www.cms.gov/Regulations-and-Guidance/Guidance/Manuals/downloads/bp102c15.pdf. Accessed September 9, 2020.
- National Comprehensive Cancer Network Categories of Evidence and Consensus. Available at: https://www.nccn.org/professionals/physician_gls/categories_of_consensus.aspx. Accessed September 9, 2020.
- 4. Center for Medicaid & Medicare Services. Medicare Benefit Policy Manual. Wolters Kluwer Clinical Drug Information Lexi-Drugs Compendium Revision Request - CAG-00443O. Available at: https://www.cms.gov/medicare-coveragedatabase/details/medicare-coverage-document-details.aspx?MCDId=31#decision. Accessed September 9, 2020.
- Wolters Kluwer Clinical Drug Information's Request for CMS evaluation of Lexi-Drugs as a compendium for use in the determination of medically-accepted indications of drugs/biologicals used off-label in anti-cancer chemotherapeutic regimens. Available at: https://www.cms.gov/Medicare/Coverage/CoverageGenInfo/downloads/covdoc31.pdf. Accessed September 9, 2020.
- Micromedex Healthcare Series. Recommendation, Evidence, and Efficacy Ratings. https://www.micromedexsolutions.com/micromedex2/librarian/ssl/true/CS/6E0ED9/ND_P R/evidencexpert/ND_P/evidencexpert/DUPLICATIONSHIELDSYNC/8B9F5B/ND_PG/evi dencexpert/ND_B/evidencexpert/ND_AppProduct/evidencexpert/ND_T/evidencexpert/P FActionId/evidencexpert.IntermediateToDocumentLink?docId=3198&contentSetId=50. Accessed September 9, 2020.

5. Revision History

| Date | Notes |
|-----------|-------------------------------------|
| 5/18/2021 | Arizona Medicaid 7.1 Implementation |

Cystaran, Cystadrops

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99663 | Cystaran, Cystadrops | |
|----------|----------------------|--|
|----------|----------------------|--|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Cystaran, Cystadrops | | |
|------------------------------------|---------------------|--|
| Diagnosis | Cystinosis | |
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of cystinosis | | |

2. Revision History

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk copy C&S Arizona Medicaid SP to Medicaid Arizona SP for eff 7 /1 |

Daliresp

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99442 Daliresp

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| 12 month(s) |
|-----------------------|
| Initial Authorization |
| Prior Authorization |
| I |

Approval Criteria

1 - Diagnosis of severe to very severe chronic obstructive pulmonary disease (COPD) (i.e., FEV1 less than or equal to 50% of predicted)

| AND |
|--|
| 2 - COPD is associated with chronic bronchitis |
| AND |
| 3 - History of COPD exacerbation(s) |

| Product Name: Daliresp | | |
|---|---------------------|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Documentation of positive clinical response to Daliresp therapy | | |

| Date | Notes |
|-----------|---|
| 3/10/2021 | Bulk Copied C&S Arizona standard to Arizona Medicaid for 7/1 effect ive |

Daraprim

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99605 Daraprim

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Daraprim, generic pyrimethamine | |
|---|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| | |

Approval Criteria

1 - Medical record documentation (e.g. chart notes) of one of the following:

1.1 Treatment of severe acquired toxoplasmosis, including toxoplasmic encephalitis

OR

1.2 Treatment of congenital toxoplasmosis

OR

1.3 Secondary prophylaxis of toxoplasmic encephalitis

OR

1.4 ALL of the following:

1.4.1 Primary Pneumocystis pneumonia (PCP) prophylaxis in human immunodeficiency virus (HIV)-infected patients or as secondary prophylaxis in HIV-infected patients who have been treated for an acute episode of Pneumocystis pneumonia

AND

1.4.2 Patient has experienced intolerance to prior prophylaxis with trimethoprimsulfamethoxazole (TMP-SMX)

AND

1.4.3 ONE of the following:

1.4.3.1 Patient has been re-challenged with trimethoprim-sulfamethoxazole (TMP-SMX) using a desensitization protocol and is still unable to tolerate

OR

1.4.3.2 Evidence of moderately severe or life threatening-reaction to trimethoprimsulfamethoxazole (TMP-SMX) in the past (e.g. toxic epidermal necrolysis (TEN), Stevens-Johnson syndrome) OR

1.5 ALL of the following:

1.5.1 Primary prophylaxis of toxoplasmic encephalitis

AND

1.5.2 Toxoplasma immunoglobulin G (IgG) positive

AND

1.5.3 CD4 (cluster of differentiation 4) less than or equal to 100 cells per mm3 if initiating prophylaxis or CD4 100-200 cells per mm3 if reinstituting prophylaxis

AND

1.5.4 Will be used in combination with dapsone or atovaquone

AND

1.5.5 Patient has experienced intolerance to prior prophylaxis with trimethoprimsulfamethoxazole (TMP-SMX)

AND

1.5.6 ONE of the following:

1.5.6.1 Patient has been re-challenged with trimethoprim-sulfamethoxazole (TMP-SMX) using a desensitization protocol and is still unable to tolerate

OR

1.5.6.2 Evidence of moderately severe or life threatening-reaction to trimethoprim-

| sulfamethoxazole (TMP-SMX) in the past (e.g. toxic epidermal necrolysis (TEN), Stevens- Johnson syndrome) | |
|--|--|
| | *Consider discontinuation of primary prophylaxis if CD4 greater than 2 00 cells/mm3 for greater than 3 months after institution of combination antiretroviral therapy. |

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona SP to Medicaid Arizona SP for 7/1 eff |

DDAVP (desmopressin) tablets - AZM

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-105310 DDAVP (desmopressin) tablets - AZM

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 4/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Brand DDAVP tablets, generic desmopressin acetate tablets | | |
|---|---------------------|--|
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - ONE of the following: | | |
| 1.1 Diagnosis of central diabetes insipidus | | |

OR

1.2 Diagnosis of polyuria and/or polydipsia following head trauma or surgery in the pituitary region

OR

1.3 Diagnosis of primary nocturnal enuresis

AND

2 - For Brand DDAVP ONLY: Trial and failure to generic desmopressin tablets (verified via paid pharmacy claims or submission of medical records)

| Notes | NOTE: Plan setup requires use of generic desmopressin tablets befor |
|-------|---|
| | e Brand DDAVP |

| Date | Notes |
|-----------|---|
| 3/29/2022 | Added step through generic tablets for Brand. |

Declomycin - Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99559 | Declomycin - Arizona |
|----------|----------------------|
|----------|----------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: demeclocycline* | | |
|---|---------------------|--|
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - ONE of the following: | | |
| 1.1 Diagnosis of ONE of the following: | | |
| Rocky Mountain spotted fever, typhus fever and the typhus group, Q fever, rickettsialpox and tick fevers caused by rickettsiae Respiratory tract infections caused by Mycoplasma pneumoniae Lymphogranuloma venereum due to Chlamydia trachomatis | | |

- Psittacosis (Ornithosis) due to Chlamydia psittaci
- Trachoma due to Chlamydia trachomatis
- Inclusion conjunctivitis caused by Chlamydia trachomatis
- Nongonococcal urethritis in adults caused by Ureaplasma urealyticum or Chlamydia trachomatis
- Relapsing fever due to Borrelia recurrentis
- Chancroid caused by Haemophilus ducreyi
- Plague due to Yersinia pestis
- Tularemia due to Francisella tularensis
- Cholera caused by Vibrio cholerae
- Campylobacter fetus infections cause by Campylobacter fetus
- Brucellosis due to Brucella species (in conjunction with streptomycin)
- Bartonellosis due to Bartonella bacilliformis
- Granuloma inguinale caused by Calymmatobacterium granulomatis
- Infection due to Escherichia coli
- Infection due to Enterobacter aerogenes
- Infection due to Shigella species
- Infection due to Acinetobacter species
- Respiratory tract infections caused by Haemophilus influenza
- Respiratory tract and urinary tract infections caused by Klebsiella species
- Upper respiratory infections caused by Streptococcus pneumoniae
- Skin and skin structure infections caused by Staphylococcus aureus.
- Uncomplicated urethritis in men due to Neisseria gonorrhoeae, and for the treatment of other uncomplicated gonococcal infections
- Infections in women caused by Neisseria gonorrhoeae
- Syphilis caused by Treponema pallidum subspecies pallidum
- Yaws caused by Treponema pallidum subspecies pertenue
- Listeriosis due to Listeria monocytogenes
- Anthrax due to Bacillus anthracis
- Vincent's infection caused by Fusobacterium fusiforme
- Actinomycosis caused by Actinomyces israelii
- Clostridial diseases caused by Clostridium species
- Acute intestinal amebiasis, as adjunctive therapy
- Severe acne, as adjunctive therapy

OR

1.2 The medication is being prescribed by or in consultation with an Infectious Disease specialist

Notes *Approval duration: 6 months

| Date | Notes |
|------|-------|
| | |

| 6/23/2021 | update program |
|-----------|----------------|
|-----------|----------------|

Dificid

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99444 Dificid

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Dificid | |
|---|---------------------|
| Approval Length | 10 Day(s) |
| Guideline Type | Prior Authorization |
| Approval Criteria 1 - Diagnosis of Clostridioides difficile-associated diarrhea (CDAD) [previously known as Clostridium difficile- associated diarrhea] | |
| AND | |

2 - ONE of the following:

2.1 History of failure, contraindication, or intolerance to Firvanq (vancomycin) oral solution

OR

2.2 History of failure, contraindication, or intolerance to oral Vancocin (vancomycin) capsules or vancomycin oral solution (NOT Firvanq) if the prescriber provides a reason or special circumstance the patient cannot use Firvanq

OR

2.3 For continuation of prior Dificid therapy

| Date | Notes |
|-----------|---|
| 3/10/2021 | Bulk Copied C&S Arizona standard to Arizona Medicaid for 7/1 effect ive |

Dofetilide - Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99445 Dofe | tilide - Arizona |
|---------------|------------------|
|---------------|------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: : Brand Tikosyn, generic dofetilide | | |
|---|---------------------|--|
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of ONE of the following: | | |
| Atrial fibrillation | | |

Atrial flutter

AND

- **2** Patient requires ONE of the following:
 - Conversion to normal sinus rhythm
 - Maintenance of normal sinus rhythm

AND

3 - Verification that the patient has already started on dofetilide while in the hospital for a minimum of 3 days

AND

4 - Patient does NOT have severe renal impairment [Creatinine Clearance (CrCl) less than 20 milliliters per minute]

AND

5 - Patient does NOT have congenital or acquired long QT syndromes

AND

6 - Patient is NOT concurrently using cimetidine, hydrochlorothiazide, ketoconazole, megestrol, prochlorperazine, trimethoprim, dolutegravir or verapamil

| Date | Notes |
|-----------|---|
| 3/10/2021 | Bulk Copied C&S Arizona standard to Arizona Medicaid for 7/1 effect ive |

Doptelet - Arizona

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Doptelet | |
|------------------------|--|
| Diagnosis | Thrombocytopenia in patients with chronic liver disease who are scheduled to undergo a procedure |
| Approval Length | 1 month(s) |
| Guideline Type | Prior Authorization |
| Approval Criteria | |
| | |

1 - Diagnosis of thrombocytopenia

AND

2 - Patient has chronic liver disease

AND

3 - Patient is scheduled to undergo a procedure

AND

4 - History of failure, contraindication, or intolerance to ALL the preferred alternatives*

• Eltrombopag Tablet (Promacta Tablet)

Romiplostim (Nplate)

| *Prior trials of formulary/PDL alternatives must sufficiently demonstrat e that the formulary/PDL alternatives are either ineffective or inapprop |
|--|
| riate at the time of the request. (Drugs may require PA) |

| Product Name: Doptelet | |
|------------------------|---------------------------------------|
| Diagnosis | Chronic Immune Thrombocytopenia (ITP) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of chronic immune thrombocytopenia (ITP)

AND

2 - ONE of the following:

2.1 BOTH of the following:

2.1.1 History of failure, contraindication, or intolerance to ONE of the following:

- Corticosteroids
- Immunoglobulins

AND

2.1.2 History of failure, contraindication, or intolerance to the preferred alternatives *

- Eltrombopag Tablet (Promacta Tablet)
- Romiplostim (Nplate)

OR

2.2 Patient is currently on Doptelet therapy

| Product Name: Doptelet | |
|------------------------|---------------------------------------|
| Diagnosis | Chronic Immune Thrombocytopenia (ITP) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Doptelet therapy

DPP-4 Inhibitors - Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Tradjenta, Januvia, Onglyza, Kombiglyze XR, Jentadueto, Janumet, Janumet XR | | |
|---|---------------------|--|
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| Approval Criteria 1 - The patient has a diagnosis of type 2 diabetes mellitus | | |
| AND | | |

2 - ONE of the following:

2.1 History of failure to metformin at a minimum dose of 1500 milligrams daily for 90 days

OR

2.2 Contraindication or intolerance to metformin

| Product Name: aloglipt Jentadueto XR | in, Nesina, alogliptin/metformin, Kazano, alogliptin/pioglitazone, Oseni, |
|---|---|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| Approval Criteria 1 - The patient has a di | iagnosis of type 2 diabetes mellitus |
| | AND |
| 2 - ONE of the following | g: |
| 2.1 History of failure to | o metformin at a minimum dose of 1500 milligrams daily for 90 days |
| | OR |
| 2.2 Contraindication c | or intolerance to metformin |
| | AND |
| 3 - ONE of the following | g: |
| 3.1 History of failure for | or 90 days to three of the following: |
| Tradjenta Januvia Onglyza | |

| ne), |
|------|
| - |
| _ |

| Date | Notes |
|-----------|-------------------|
| 6/22/2021 | Updated guideline |

Dry Eye Disease - Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99582 | Dry Eye Disease - Arizona |
|----------|---------------------------|
|----------|---------------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Cequa, Xiidra | |
|-----------------------------|---|
| Diagnosis | Tear deficiency associated with ocular inflammation |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- 1 Tear deficiency associated with ocular inflammation due to ONE of the following:
 - Moderate to severe keratoconjuctivitis sicca

Moderate to severe Dry Eye Disease

AND

2 - Not prescribed to manage dry eyes peri-operative elective eye surgery (e.g.: LASIK)

AND

3 - History of failure to at least three over-the-counter (OTC) artificial tear products (e.g.: Systane Ultra, Akwa Tears, Refresh Optive, Soothe XP, Muro 128 2% Solution, Muro 128 5% Solution, Muro 128 5% Ointment) in the past 60 days as evidenced in the member's claim history.

AND

4 - Prescribed by or in consultation with ONE of the following:

- Ophthalmologist
- Optometrist
- Rheumatologist

AND

5 - The patient has claims history indicating a minimum trial of 60 days of Restasis unless it is contraindicated.

| Tear deficiency associated with ocular inflammation |
|---|
| 12 month(s) |
| Reauthorization |
| Prior Authorization |
| Í |

Approval Criteria

1 - Patient has demonstrated clinically significant improvement with therapy

| Date | Notes |
|-----------|---------------------------------|
| 8/25/2021 | Arizona Medicaid Implementation |

Duexis and Vimovo - Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99563 | Duexis and | Vimovo - Arizona |
|----------|------------|------------------|
|----------|------------|------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Duexis | |
|----------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - ONE of the following risk factors for NSAID (non-steroidal anti-inflammatory drug) induced adverse GI (gastrointestinal) events:

- Patient is greater than or equal to 65 years of age
- Prior history of peptic, gastric, or duodenal ulcer
- History of NSAID-related ulcer

| History of clinically significant GI bleedi | story of clinically signature | gnificant GI bleedin | D |
|---|-------------------------------|----------------------|---|
|---|-------------------------------|----------------------|---|

- Untreated or active H. Pylori gastritis
- Concurrent use of oral corticosteroids (eg, prednisone, prednisolone, dexamethasone)
- Concurrent use of anticoagulants (eg, warfarin, heparin)
- Concurrent use of antiplatelets (eg, aspirin including low-dose, clopidogrel)

AND

2 - Documentation of history of failure, contraindication, or intolerance to THREE combinations of preferred NSAIDS taken with preferred H2 (histamine 2)-receptor antagonists. (Provide name and date preferred products were tried)*

AND

3 - Physician has provided rationale for needing to use fixed-dose combination therapy with Duexis instead of taking individual products in combination.

| Notes *Please reference background section for preferred products table | е |
|---|---|
|---|---|

| Product Name: Brand Vimovo, generic naproxen-esomeprazole | |
|---|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - ONE of the following risk factors for NSAID (non-steroidal anti-inflammatory drug) induced adverse GI (gastrointestinal) events:

- Patient is greater than or equal to 65 years of age
- Prior history of peptic, gastric, or duodenal ulcer
- History of NSAID-related ulcer
- History of clinically significant GI bleeding
- Untreated or active H. Pylori gastritis
- Concurrent use of oral corticosteroids (eg, prednisone, prednisolone, dexamethasone)
- Concurrent use of anticoagulants (eg, warfarin, heparin)
- Concurrent use of antiplatelets (eg, aspirin including low-dose, clopidogrel)

AND

2 - Documentation of history of failure, contraindication, or intolerance to THREE

combinations of preferred NSAIDS taken with preferred proton pump inhibitors (PPIs). (Provide name and date preferred products were tried)*

AND

3 - Physician has provided rationale for needing to use fixed-dose combination therapy with Vimovo instead of taking individual products in combination.

| Notes | *Please reference background section for preferred products table |
|-------|---|
|-------|---|

2. Background

| Benefit/Coverage/Program Information | | |
|--|---|--|
| Preferred Table | | |
| NSAIDS | Proton Pump Inhibitors (PPIs) | H2 (histamine 2)-receptor antagonists |
| Diclofenac DR (Generic Voltaren) | esomeprazole (Generic Nexium) | Famotidine (Generic Pepcid) |
| Diclofenac ER (Generic Voltaren ER) | lansoprazole (Generic Prevacid) | Nizatidine (Generic Axid) |
| Etodolac (Generic Lodine) | omeprazole (Generic Prilosec) | Ranitidine (Generic Zantac) |
| Etodolac ER (Generic Lodine ER) | pantoprazole sodium (Generic Protonix) | |
| Fenoprofen (Generic Nalfon) | | |
| Flurbiprofen (Generic Ansaid) | | |
| Ibuprofen | | |

| Indomethacin (Generic | |
|-------------------------------------|------|
| Indocin) | |
| , | |
| | |
| Ketorolac (Generic Toradol) | |
| | |
| | |
| Mefenamic (Generic | |
| Ponstel) | |
| | |
| | |
| Meloxicam (Generic Mobic) | |
| | |
| | |
| Nabumetone (Generic | |
| Relafen) | |
| | |
| | |
| Nabumetone DS (Generic | |
| Relafen DS) | |
| | |
| | |
| Naproxen (Generic | |
| Anaprox) | |
| | |
| Noprovon DB (Conorio | |
| Naproxen DR (Generic Anaprox DR) | |
| | |
| | |
| Naproxen EC (Generic | |
| Anaprox EC) | |
| , | |
| | |
| Oxaprozin (Generic Daypro) | |
| | |
| | |
| Piroxicam (Generic | |
| Feldene) | |
| | |
| Sulindac (Generic Clinoril) | |
| | |
| | |

Duopa

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99446 Duopa

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Duopa | |
|---|-----------------------|
| Diagnosis | Parkinson's disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of advanced Parkinson's disease | |

AND

2 - Patient is levodopa-responsive

AND

3 - Patient experiences disabling "off" periods for a minimum of 3 hours per day

AND

4 - Disabling "off" periods occur despite therapy with BOTH of the following:

- Oral levodopa-carbidopa
- One drug from a different class of anti-Parkinson's disease therapy (e.g., COMT [catechol-O-methyltransferase] inhibitor [entacapone, tolcapone], MAO-B [monoamine oxidase-B] inhibitor [selegiline, rasagiline], dopamine agonist [pramipexole, ropinirole])

AND

5 - Has undergone or has planned placement of a procedurally-placed tube

AND

6 - Prescribed by or in consultation with a neurologist

| Product Name: Duopa | |
|---------------------|---------------------|
| Diagnosis | Parkinson's disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Documentation of positive clinical response to Duopa therapy

| Date | Notes |
|-----------|---|
| 3/10/2021 | Bulk Copied C&S Arizona standard to Arizona Medicaid for 7/1 effect ive |

Dupixent (dupilumab)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-112924 | Dupixent (dupilumab) |
|-----------|----------------------|
|-----------|----------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 9/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Dupixent | |
|---|-----------------------|
| Diagnosis | Atopic Dermatitis |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Patient is 6 months of age or older | |

AND

2 - Submission of documentation (e.g., chart notes) confirming ONE of the following:

2.1 BOTH of the following:

2.1.1 Diagnosis of moderate to severe chronic atopic dermatitis

AND

2.1.2 History of failure, contraindication, or intolerance to the following topical therapies: (document drug, date of trial, and/or contraindication to medication)*

- One medium to very-high potency topical corticosteroid [e.g., Elocon (mometasone furoate), Synalar (fluocinolone acetonide), Lidex (fluocinonide)] (see Table 1 in Background section)
- One topical calcineurin inhibitor [e.g., Elidel (pimecrolimus), Protopic (tacrolimus)]
- Eucrisa (crisaborole)

OR

2.2 BOTH of the following:

2.2.1 Diagnosis of chronic atopic dermatitis that has been determined to be severe based on physician assessment

AND

2.2.2 History of failure, contraindication, or intolerance to BOTH of the following topical therapies: (document drug, date of trial, and/or contraindication to medication)*

- Medium to very-high potency topical corticosteroid [e.g., Elocon (mometasone furoate), Synalar (fluocinolone acetonide), Lidex (fluocinonide)] (see Table 1 in Background section)
- One topical calcineurin inhibitor [e.g., Elidel (pimecrolimus), Protopic (tacrolimus)]

| Product Name: Dupixent | |
|------------------------|---------------------|
| Diagnosis | Atopic Dermatitis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of documentation (e.g., chart notes) confirming positive clinical response to Dupixent therapy

AND

2 - Patient is NOT receiving Dupixent in combination with another biologic medication [e.g., Xolair (omalizumab), Rituxan (rituximab), Enbrel (etanercept), Remicade/Inflectra (infliximab)]

AND

3 - Prescribed by one of the following:

- Dermatologist
- Allergist
- Immunologist

| Product Name: Dupixent | |
|------------------------|-----------------------|
| Diagnosis | Asthma |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of documentation (e.g., chart notes) confirming diagnosis of moderate to severe asthma

AND

2 - Patient is 6 years of age or older

AND

- **3** ONE of the following:
- **3.1** ALL of the following:

3.1.1 Classification of asthma as uncontrolled or inadequately controlled as defined by at least ONE of the following

- Poor symptom control (e.g., Asthma Control Questionnaire [ACQ] score consistently greater than 1.5 or Asthma Control Test [ACT] score consistently less than 20)
- Two or more bursts of systemic corticosteroids for at least 3 days each in the previous 12 months

• Asthma-related emergency treatment (e.g., emergency room visit, hospital admission, or unscheduled physician's office visit for nebulizer or other urgent treatment)

- Airflow limitation (e.g., after appropriate bronchodilator withhold forced expiratory volume in 1 second [FEV1] less than 80% predicted [in the face of reduced FEV1/forced vital capacity [FVC] defined as less than the lower limit of normal])
- Patient is currently dependent on oral corticosteroids for the treatment of asthma

AND

3.1.2 Dupixent will be used in combination with one of the following:

3.1.2.1 ONE high-dose (appropriately adjusted for age) combination inhaled corticosteroid (ICS)/long-acting beta2 agonist (LABA) [e.g., Advair/AirDuo Respiclick (fluticasone propionate/salmeterol), Symbicort (budesonide/formoterol), Breo Ellipta (fluticasone furoate/vilanterol)] (see Table 2 in Background section)

OR

3.1.2.2 Combination therapy including BOTH of the following:

3.1.2.2.1 ONE high-dose (appropriately adjusted for age) ICS product [e.g., ciclesonide (Alvesco), mometasone furoate (Asmanex), beclomethasone dipropionate (QVAR)] (see Table 2 in Background section)

AND

3.1.2.2.2 ONE additional asthma controller medication [e.g., LABA - olodaterol (Striverdi) or indacaterol (Arcapta); leukotriene receptor antagonist – montelukast (Singulair); theophylline]

AND

3.1.3 ONE of the following:

3.1.3.1 Submission of medical records (e.g., chart notes, laboratory values, etc.) documenting that asthma is an eosinophilic phenotype as defined by a baseline (predupilumab treatment) peripheral blood eosinophil level greater than or equal to 150 cells/microliter within the past 6 weeks

OR

3.1.3.2 Patient is currently dependent on oral corticosteroids for the treatment of asthma

OR

3.2 Patient is currently on Dupixent therapy

AND

4 - Patient is NOT receiving Dupixent in combination with ONE of the following:

- Anti-interleukin-5 therapy [e.g. Nucala (mepolizumab), Cinqair (resilizumab), Fasenra (benralizumab)]
- Anti-IgE (immunoglobulin E) therapy [e.g. Xolair (omalizumab)]

AND

5 - Prescribed by one of the following:

- Pulmonologist
- Allergist
- Immunologist

| Product Name: Dupixent | | |
|------------------------|------------------------------------|--|
| Diagnosis | Asthma | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Guideline Type Prior Authorization | |

Approval Criteria

1 - Submission of documentation (e.g., chart notes) confirming positive clinical response to Dupixent therapy as demonstrated by at least ONE of the following:

| Reduction in the frequency of exacerbations Decreased utilization of rescue medications Increase in percent predicted forced expiratory volume in 1 second (FEV1) from pretreatment baseline Reduction in severity or frequency of asthma-related symptoms (e.g., wheezing, shortness of breath, coughing, etc.) Reduction in oral corticosteroid requirements |
|--|
| AND |
| 2 - Dupixent is being used in combination with an inhaled corticosteroid (ICS)-containing controller medication (see Table 2 in Background section) |
| AND |
| 3 - Patient is NOT receiving Dupixent in combination with ONE of the following: |
| Anti-interleukin-5 therapy [e.g. Nucala (mepolizumab), Cinqair (resilizumab), Fasenra (benralizumab)] Anti-IgE (immunoglobulin E) therapy [e.g. Xolair (omalizumab)] |
| AND |
| 4 - Prescribed by one of the following: |

- Pulmonologist Allergist Immunologist •
- •

•

| Product Name: Dupixent | | |
|------------------------|---|--|
| Diagnosis | Chronic Rhinosinusitis with Nasal Polyposis | |
| Approval Length | 6 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| | | |

Approval Criteria

1 - Patient is 18 years of age or older

AND

2 - Submission of documentation (e.g., chart notes) confirming ONE of the following:

2.1 ALL of the following:

2.1.1 Diagnosis of chronic rhinosinusitis with nasal polyposis (CRSwNP) defined by ALL of the following:

2.1.1.1 TWO or more of the following symptoms for greater than or equal to 12 weeks duration:

- Mucopurulent discharge
- Nasal obstruction and congestion
- Decreased or absent sense of smell
- Facial pressure or pain

AND

2.1.1.2 ONE of the following:

- Evidence of inflammation on paranasal sinus examination or computed tomography (CT)
- Evidence of purulence coming from paranasal sinuses or ostiomeatal complex

AND

2.1.1.3 The presence of nasal polyps

AND

2.1.2 ONE of the following:

- Patient has required prior sino-nasal surgery
- Patient has required systemic corticosteroids in the previous 2 years

2.1.3 Patient has been unable to obtain symptom relief after trial of ALL of the following agents/classes of agents:

- Nasal saline irrigations
- Intranasal corticosteroids (e.g. fluticasone, mometasone, triamcinolone, etc.)
- Antileukotriene agents (e.g. montelukast, zafirlukast, zileuton)

OR

2.2 ALL of the following:

2.2.1 Diagnosis of chronic rhinosinusitis with nasal polyposis (CRSwNP)

AND

2.2.2 Patient is currently on Dupixent therapy

AND

3 - Patient will receive Dupixent as add-on maintenance therapy in combination with intranasal corticosteroids

AND

4 - Patient is NOT receiving Dupixent in combination with another biologic medication [e.g., Xolair (omalizumab), Nucala (mepolizumab), Cinqair (resilizumab), Fasenra (benralizumab)]

AND

5 - Prescribed by one of the following:

- Otolaryngologist
- Allergist
- Immunologist

| Product Name: Dupixent | | |
|---|--|--|
| Chronic Rhinosinusitis with Nasal Polyposis | | |
| 12 month(s) | | |
| Reauthorization | | |
| Prior Authorization | | |
| | | |

Approval Criteria

1 - Submission of documentation (e.g., chart notes) confirming positive clinical response to Dupixent therapy

AND

2 - Patient will continue to receive Dupixent as add-on maintenance therapy in combination with intranasal corticosteroids

AND

3 - Patient is NOT receiving Dupixent in combination with another biologic medication [e.g., Xolair (omalizumab), Nucala (mepolizumab), Cinqair (resilizumab), Fasenra (benralizumab)]

AND

4 - Prescribed by one of the following:

- Otolaryngologist
- Allergist
- Immunologist

| Product Name: Dupixent | |
|-------------------------------------|--------------------------------|
| Diagnosis | Eosinophilic Esophagitis (EoE) |
| Approval Length | 12 month(s) |
| Therapy Stage Initial Authorization | |

| Guideline Type | Prior Authorization | |
|---|---|--|
| | | |
| Approval Criteria | | |
| Submission of documentation (e.g., chart notes) confirming diagnosis of eosinophilic esophagitis (EoE) | | |
| | AND | |
| | ms of esophageal dysfunction (e.g., dysphagia, food impaction, ix disease [GERD]/heartburn symptoms, chest pain, abdominal pain) | |
| | AND | |
| | imentation (e.g., chart notes, lab values) confirming patient has at least ophils per high power field (HPF) | |
| | AND | |
| 4 - Other causes of eso | ophageal eosinophilia have been excluded | |
| | AND | |
| 5 - Both of the following | g: | |
| Patient is at leaPatient weighs | at least 40 kg | |
| | AND | |
| | nission of documentation (e.g., chart notes) confirming trial and failure, plerance to at least an 8-week trial of one of the following: | |
| Proton pump inhibitors (e.g., pantoprazole, omeprazole) Topical (esophageal) corticosteroids (e.g., budesonide, fluticasone) | | |

- 7 Prescribed by one of the following:
 - Gastroenterologist
 - Allergist
 - Immunologist

| Product Name: Dupixent | | |
|------------------------|--------------------------------|--|
| Diagnosis | Eosinophilic Esophagitis (EoE) | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Submission of documentation (e.g., chart notes) confirming positive clinical response to therapy as evidenced by improvement of at least one of the following from baseline:

- Symptoms (e.g., dysphagia, food impaction, heartburn, chest pain)
- Histologic measures (e.g., esophageal intraepithelial eosinophil count)
- Endoscopic measures (e.g., edema, furrows, exudates, rings, strictures)

AND

2 - Prescribed by one of the following:

- Gastroenterologist
- Allergist
- Immunologist

2. Background

Benefit/Coverage/Program Information

| Class | Drug | Dosage Form | Strength (%) |
|----------------------|--------------------------------------|---------------------------------|-----------------|
| Very high potency | Augmented betamethasone dipropionate | Ointment, gel | 0.05 |
| | Clobetasol propionate | Cream, foam, ointment | 0.05 |
| | Diflorasone diacetate | Ointment | 0.05 |
| | Halobetasol propionate | Cream, ointment | 0.05 |
| High Potency | Amcinonide | Cream, lotion, ointment | 0.1 |
| Potency | Augmented betamethasone dipropionate | Cream, lotion | 0.05 |
| | Betamethasone dipropionate | Cream, foam, ointment, solution | 0.05 |
| | Desoximetasone | Cream, ointment | 0.25 |
| | Desoximetasone | Gel | 0.05 |
| | Diflorasone diacetate | Cream | 0.05 |
| | tridifloronide | Cream, gel, ointment, solution | 0.05 |
| | Halcinonide | Cream, ointment | 0.1 |
| | Mometasone furoate | Ointment | 0.1 |
| | Triamcinolone acetonide | Cream, ointment | 0.5 |
| Medium | Betamethasone valerate | Cream, foam, lotion, ointment | 0.1 |
| potency | Clocortolone pivalate | Cream | 0.1 |
| | Desoximetasone | Cream | 0.05 |
| | Fluocinolone acetonide | Cream, ointment | 0.025 |
| | Flurandrenolide | Cream, ointment, lotion | 0.05 |
| | Fluticasone propionate | Cream | 0.05 |
| | Fluticasone propionate | Ointment | 0.005 |
| | Mometasone furoate | Cream, lotion | 0.1 |

| | Triamcinolone acetonide | Cream, ointme | ent, lotion | 0.1 |
|--|---|--|---|--|
| Lower- | Hydrocortisone butyrate | Cream, ointme | ent, solution | 0.1 |
| medium potency | Hydrocortisone probutate | Cream | | 0.1 |
| | Hydrocortisone valerate | Cream, ointme | ent | 0.2 |
| | Prednicarbate | Cream | | 0.1 |
| Low | Alclometasone dipropionate | Cream, ointment | | 0.05 |
| potency | Desonide | Cream, gel, fo | am, ointment | 0.05 |
| | Fluocinolone acetonide | Cream, solution | on | 0.01 |
| Lowest | Dexamethasone | Cream | | 0.1 |
| potency | Hydrocortisone | Cream, lotion, ointment, solution | | 0.25, 0.5, |
| | | | | |
| | Hydrocortisone acetate w, medium and high daily dose (12 years of age and older) | Cream, ointme es of inhaled co | | 0.5-1 dults and |
| | | es of inhaled co | | |
| olescents | w, medium and high daily dos | es of inhaled co | orticosteroids A | |
| Drug | w, medium and high daily dos | es of inhaled co | orticosteroids A aily dose (mcg) | dults and |
| Drug Beclome | w, medium and high daily dose (12 years of age and older) | es of inhaled co Da | aily dose (mcg) | dults and High |
| Drug Beclome | w, medium and high daily dose (12 years of age and older) etasone dipropionate (CFC) etasone dipropionate (HFA) | es of inhaled co Da Low 200-500 | aily dose (mcg) Medium >500-1000 | dults and High >1000 |
| Drug Drug Beclome Beclome Budesor | w, medium and high daily dose (12 years of age and older) etasone dipropionate (CFC) etasone dipropionate (HFA) | es of inhaled co Da Low 200-500 100-200 | aily dose (mcg) Medium >500-1000 >200-400 | dults and High >1000 >400 |
| Drug Drug Beclome Beclome Budesor Ciclesor | w, medium and high daily dose (12 years of age and older) etasone dipropionate (CFC) etasone dipropionate (HFA) hide DPI | es of inhaled co Da Low 200-500 100-200 200-400 | Anticosteroids A aily dose (mcg) Medium >500-1000 >200-400 >400-800 | dults and High >1000 >400 >800 |
| Drug Drug Beclome Beclome Budesor Ciclesor Fluticase | w, medium and high daily dose (12 years of age and older) etasone dipropionate (CFC) etasone dipropionate (HFA) hide DPI | es of inhaled co Da Low 200-500 100-200 200-400 80-160 | Anticosteroids A aily dose (mcg) Medium >500-1000 >200-400 >400-800 >160-320 | High >1000 >400 >800 >320 |
| Drug Drug Beclome Beclome Budesor Ciclesor Fluticaso | w, medium and high daily dose (12 years of age and older) etasone dipropionate (CFC) etasone dipropionate (HFA) nide DPI nide (HFA) one furoate (DPI) | es of inhaled co Da Low 200-500 100-200 200-400 80-160 100 | orticosteroids A aily dose (mcg) Medium >500-1000 >200-400 >400-800 >160-320 N/A | High >1000 >400 >800 >320 200 |
| Drug Drug Beclome Budesor Ciclesor Fluticaso Fluticaso | w, medium and high daily dose (12 years of age and older) etasone dipropionate (CFC) etasone dipropionate (HFA) nide DPI nide (HFA) one furoate (DPI) one propionate (DPI) | es of inhaled co Da Low 200-500 100-200 200-400 80-160 100 100-250 | orticosteroids A aily dose (mcg) Medium >500-1000 >200-400 >400-800 >160-320 N/A >250-500 | High >1000 >400 >800 >320 200 >500 |

| Date | Notes |
|-----------|--|
| 8/29/2022 | Updated age requirements, added criteria for new EoE indication. |

Durezol

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99567 Durezol

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Durezol | |
|-----------------------|---------------------|
| Approval Length | 2 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - History of failure, contraindication, or intolerance to BOTH of the following:

- prednisolone 1%
- dexamethasone ophthalmic drops and/or ointment.

| Date | Notes |
|----------|-------------------------------------|
| 7/8/2021 | Changed approval length to 2 months |

Ecoza (econazole)

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99550 Ecoza (econazole)

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|--------------------|-----------|
| P&T Approval Date: | |
| P&T Revision Date: | |

1. Criteria

| Product Name: Ecoza, Generic econazole | | |
|---|---------------------|--|
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - History of failure, contraindication, or intolerance to ALL of the following: | | |

• butenafine

- •
- ciclopirox clotrimazole •
- clotrimazole w/ betamethasone •
- ketoconazole
- miconazole •
- •
- nystatin terbinafine •
- tolnaftate •

| Date | Notes |
|-----------|------------------|
| 6/10/2021 | Update guideline |

Egrifta

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99606 Egrifta

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Egrifta SV | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of human immunodeficiency virus (HIV)-associated lipodystrophy | |

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona SP to Medicaid Arizona SP for 7/1 eff |

Elaprase - Arizona

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99607 | Elaprase - Arizona |
|----------|--------------------|
|----------|--------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Elaprase | |
|---|---------------------|
| Diagnosis | Hunter syndrome |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of Hunter syndrome (Mucopolysaccharidosis II, MPS II) | |

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona SP to Medicaid Arizona SP for 7/1 eff |

Elidel-Protopic

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99447 Elidel-Protopic

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Elidel, generic pimecrolimus, Brand Protopic 0.03%, generic tacrolimus 0.03% | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| Approval Criteria 1 - The patient is 2 years of age or older | |
| AND | |

2 - ONE of the following:

2.1 History of failure, contraindication, or intolerance to ONE topical corticosteroid in the past 90 days

OR

2.2 Drug is being prescribed for the facial or groin area

| Product Name: Brand I | Protopic 0.1%, generic tacrolimus 0.1% |
|--|--|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - The patient is 16 ye | ears of age or older |
| | AND |
| | AND |
| 2 - ONE of the following | g: |
| 2.1 History of failure, 90 days | contraindication, or intolerance to ONE topical corticosteroid in the past |
| | OR |
| 2.2 Drug is being pres | scribed for the facial or groin area |

| Date | Notes |
|-----------|---|
| 3/10/2021 | Bulk Copied C&S Arizona standard to Arizona Medicaid for 7/1 effect ive |

Elmiron

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99448 Elmiron

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Elmiron | |
|--|--|
| Bladder pain or discomfort associated with interstitial cystitis | |
| 12 month(s) | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Patient has a documented diagnosis of bladder pain or discomfort associated with interstitial cystitis

| Date | Notes |
|-----------|---|
| 3/10/2021 | Bulk Copied C&S Arizona standard to Arizona Medicaid for 7/1 effect ive |

Emflaza - Arizona

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99608 | Emflaza - Arizona |
|----------|-------------------|
|----------|-------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Emflaza | |
|-----------------------|-----------------------------|
| Diagnosis | Duchenne Muscular Dystrophy |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| | |

1 - Diagnosis of Duchenne muscular dystrophy

2 - Patient is 2 years of age or older

AND

3 - History of failure, contraindication, or intolerance to ONE of the following for the treatment of Duchenne muscular dystrophy:

• Prednisone

• Prednisolone

AND

4 - Prescribed by or in consultation with a neurologist

| Product Name: Emflaza | |
|-----------------------|-----------------------------|
| Diagnosis | Duchenne Muscular Dystrophy |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Physician attestestation that the patient has had a positive clinical response to Emflaza therapy

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona SP to Medicaid Arizona SP for 7/1 eff |

Enbrel

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99714 Enbrel

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Enbrel | | |
|----------------------|---|--|
| Diagnosis | Moderately to Severely Active Rheumatoid Arthritis (RA) | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |

Approval Criteria

1 - Diagnosis of moderately to severely active Rheumatoid Arthritis (RA)

2 - History of failure to a 3 month trial of ONE non-biologic disease modifying anti-rheumatic drug (DMARD) [e.g., methotrexate, leflunomide, sulfasalazine, hydroxychloroquine] at maximally indicated doses within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*

AND

3 - Patient is not receiving Enbrel in combination with ONE of the following:

- Biologic DMARD [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

4 - Prescribed by or in consultation with a rheumatologist

| *Note: Claims history may be used in conjunction as documentation of |
|--|
| drug, date, and duration of trial |

| Product Name: Enbrel | | |
|----------------------|---|--|
| Diagnosis | Moderately to Severely Active Rheumatoid Arthritis (RA) | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Documentation of positive clinical response to Enbrel therapy

AND

2 - Patient is not receiving Enbrel in combination with ONE of the following:

- Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with a rheumatologist

| Product Name: Enbrel | | |
|----------------------|---|--|
| Diagnosis | Moderately to Severely Active Polyarticular Juvenile Idiopathic Arthritis | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Diagnosis of moderately to severely active polyarticular juvenile idiopathic arthritis

AND

2 - Patient is not receiving Enbrel in combination with ONE of the following:

- Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with a rheumatologist

Product Name: Enbrel

| Diagnosis | Moderately to Severely Active Polyarticular Juvenile Idiopathic Arthritis |
|-----------------|---|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Enbrel therapy

AND

2 - Patient is not receiving Enbrel in combination with ONE of the following:

- Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with a rheumatologist

| Des dust Newsey Falsed | | |
|------------------------|----------------------------|--|
| Product Name: Enbrel | | |
| Diagnosis | Active Psoriatic Arthritis | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Diagnosis of active psoriatic arthritis

2 - History of failure to a 3 month trial of methotrexate at the maximally indicated dose within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*

AND

3 - Patient is not receiving Enbrel in combination with ONE of the following:

- Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

4 - Prescribed by or in consultation with ONE of the following:

- Rheumatologist
- Dermatologist

| Note: Claims history may be used in conjunction as documentation of drug, date, and duration of trial |
|---|

| Product Name: Enbrel | |
|-------------------------------|----------------------------|
| Diagnosis | Active Psoriatic Arthritis |
| Approval Length | 12 month(s) |
| Therapy Stage Reauthorization | |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Enbrel therapy

AND

2 - Patient is not receiving Enbrel in combination with ONE of the following:

- Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

3 - Prescribed by or in consultation with ONE of the following:

- Rheumatologist
- Dermatologist

| Product Name: Enbrel | | |
|----------------------|---|--|
| Diagnosis | Moderate to Severe Chronic Plaque Psoriasis | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Diagnosis of moderate to severe chronic plaque psoriasis

AND

2 - Greater than or equal to 3% body surface area involvement, palmoplantar, facial, or genital involvement, or severe scalp psoriasis

AND

3 - Both of the following:

3.1 History of failure to one of the following topical therapies, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial):*

• Corticosteroids (e.g., betamethasone, clobetasol, desonide)

| • | Vitamin D | analogs | (e.g., | calcitriol, | calcipotriene) |
|---|-----------|---------|--------|-------------|----------------|
|---|-----------|---------|--------|-------------|----------------|

- Tazarotene
- Calcineurin inhibitors (e.g., tacrolimus, pimecrolimus)
- Anthralin
- Coal tar

3.2 History of failure to a 3 month trial of methotrexate at the maximally indicated dose within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*

AND

4 - Patient is not receiving Enbrel in combination with ONE of the following:

- Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

5 - Prescribed by or in consultation with a dermatologist

| Notes | *Note: Claims history may be used in conjunction as documentation of |
|-------|--|
| | drug, date, and duration of trial |

| Product Name: Enbrel | | |
|----------------------|---|--|
| Diagnosis | Moderate to Severe Chronic Plaque Psoriasis | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Documentation of positive clinical response to Enbrel therapy

2 - Patient is not receiving Enbrel in combination with ONE of the following:

- Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with a dermatologist

| Product Name: Enbrel | |
|----------------------|------------------------|
| Diagnosis | Ankylosing spondylitis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of active ankylosing spondylitis

AND

2 - History of failure to two non-steroidal anti-inflammatory drugs (NSAIDs: e.g., ibuprofen, naproxen) at maximally indicated doses, each used for at least 4 weeks within the last 3 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trials)*

AND

3 - Patient is not receiving Enbrel in combination with ONE of the following:

• Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]

- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

| AND | |
|--|--|
| 4 - Prescribed by or in consultation with a rheumatologist | |
| Notes | *Note: Claims history may be used in conjunction as documentation of drug, date, and duration of trial |

| Product Name: Enbrel | |
|----------------------|------------------------|
| Diagnosis | Ankylosing Spondylitis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Enbrel therapy

AND

- 2 Patient is not receiving Enbrel in combination with ONE of the following:
 - Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
 - Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
 - Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with a rheumatologist

| Date | Notes |
|-----------|-------------------------------------|
| 5/13/2021 | Arizona Medicaid 7.1 Implementation |

Endari

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99450 Endari

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Endari | |
|-----------------------|--|
| Sickle cell disease | |
| 12 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |

Approval Criteria

- **1** BOTH of the following:
 - Diagnosis of sickle cell disease

• Used to reduce acute complications of sickle cell disease

AND

- **2** ONE of the following:
 - Patient is using Endari with concurrent hydroxyurea therapy
 - Patient is unable to take hydroxyurea due to a contraindication or intolerance

AND

3 - Patient has had 2 or more painful sickle cell crises within the past 12 months

| Product Name: Endari | | |
|---|---------------------|--|
| Diagnosis | Sickle cell disease | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Documentation of positive clinical response to Endari therapy | | |

| Date | Notes |
|-----------|---|
| 3/10/2021 | Bulk Copied C&S Arizona standard to Arizona Medicaid for 7/1 effect ive |

Entocort EC

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99451 Entocort EC

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Entocort EC, generic budesonide | |
|---|---------------------|
| Diagnosis | Chrohn's Disease |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Entocort EC is being used for the treatment of Crohn's disease

| Date | Notes |
|-----------|---|
| 3/10/2021 | Bulk Copied C&S Arizona standard to Arizona Medicaid for 7/1 effect ive |

Entresto

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99452 Entresto

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Entresto | |
|------------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - As continuation of therapy initiated during an inpatient stay

OR

2 - Both of the following:

2.1 Diagnosis of pediatric heart failure with systemic left ventricular systolic dysfunction which is symptomatic

AND

2.2 Prescribed by or in consultation with a cardiologist

OR

3 - ALL of the following:

3.1 Diagnosis of heart failure (with or without hypertension)

AND

3.2 Ejection fraction is less than or equal to 40 percent

AND

3.3 Heart failure is classified as ONE of the following:

- New York Heart Association Class II
- New York Heart Association Class III
- New York Heart Association Class IV

AND

3.4 ONE of the following:

3.4.1 Patient is on a stabilized dose and receiving concomitant therapy with ONE of the following beta-blockers:

bisoprolol • carvedilol • metoprolol OR **3.4.2** Patient has a contraindication or intolerance to beta-blocker therapy AND 3.5 Patient does not have a history of angioedema AND **3.6** Patient will discontinue any use of concomitant ACE (angiotensin converting enzyme) Inhibitor or ARB (angiotensin II receptor blocker) before initiating treatment with Entresto* AND **3.7** Patient is not concomitantly on aliskiren therapy AND 3.8 Entresto is prescribed by, or in consultation with, a cardiologist Notes *NOTE: ACE inhibitors must be discontinued at least 36 hours prior to initiation of Entresto

| Product Name: Entresto | |
|------------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - The Entresto dose has been titrated to a dose of 97 mg (milligrams) /103 mg twice daily, or to a maximum dose as tolerated by the patient

AND

2 - Documentation of positive clinical response to therapy

| Date | Notes |
|-----------|---|
| 3/10/2021 | Bulk Copied C&S Arizona standard to Arizona Medicaid for 7/1 effect ive |

Epaned

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99453 Epaned

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Epaned | |
|---|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - ONE of the following: | |
| 1.1 Patient is less than 8 years of age | |

OR

1.2 BOTH of the following:

1.2.1 ONE of the following diagnoses:

- Hypertension
- Heart failure
- Asymptomatic left ventricular dysfunction, defined as left ventricular ejection fraction less than or equal to 35%

AND

1.2.2 ONE of the following:

1.2.2.1 History of failure, contraindication, or intolerance to TWO formulary oral antihypertensives (e.g., angiotensin-converting enzyme (ACE) inhibitor, ACE inhibitor combination, angiotensin-receptor blockers (ARB), ARB combination, thiazide diuretic)

OR

1.2.2.2 Patient is unable to ingest a solid dosage form (e.g. an oral tablet or capsule) due to ONE of the following:

- Oral/motor difficulties
- Dysphagia

| Date | Notes |
|-----------|---|
| 3/10/2021 | Bulk Copied C&S Arizona standard to Arizona Medicaid for 7/1 effect ive |

Epinephrine Pens

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 6/23/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Epinephrine Pens (Non-Mylan Manufacturer) | | |
|--|---|--|
| Approval Length | 6 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - There is a shortage on Epinephrine Pens manufactured by Mylan. | | |
| Notes | *Only approve other rebatable epinephrine autoinjectors if both the br anded EpiPen and authorized generic are on the FDA shortage list. | |

| Product Name: Epinephrine Pens (Mylan Manufacturer) | | |
|---|----------------|--|
| Approval Length | 6 month(s) | |
| Guideline Type | Quantity Limit | |
| | | |
| Approval Criteria | | |
| 1 - Medication has been used or lost or the member is going on vacation.* | | |
| Notes Only approve other rebatable epinephrine autoinjectors if both the bra nded EpiPen and authorized generic are on the FDA shortage list | | |

| Date | Notes |
|-----------|---|
| 6/23/2022 | Updated guideline name as criteria is not specific to only non-mylan products |

Eplerenone- Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99454 Eplerenone- Arizona

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Inspra, generic eplerenone | |
|--|--|
| Approval Length 12 month(s) | |
| Guideline Type Prior Authorization | |
| | |
| Approval Criteria | |

1 - Diagnosis of one of the following:

1.1 Symptomatic heart failure with reduced ejection fraction (HFrEF) after an acute myocardial infarction

| | OR | |
|------------------|----|--|
| 1.2 Hypertension | | |

| Date | Notes |
|-----------|---|
| 3/10/2021 | Bulk Copied C&S Arizona standard to Arizona Medicaid for 7/1 effect ive |

Epsolay (benzoyl peroxide) cream

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-108675 Epsolay (benzoyl peroxide) cream

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 7/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Epsolay | | |
|--------------------------|---------------------|--|
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of rosacea | | |
| | | |
| AND | | |
| | | |

| 2 - Patient has inflammatory lesions | 2 - | Patient | has i | nflammato | ry lesions |
|---|-----|---------|-------|-----------|------------|
|---|-----|---------|-------|-----------|------------|

AND

3 - Trial and failure (of a minimum 30-day supply), contraindication or intolerance to one preferred topical product for rosacea (e.g., metronidazole cream/gel/lotion) (verified via paid pharmacy claims)

| Date | Notes |
|-----------|-------------|
| 6/24/2022 | New Program |

Erivedge

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99679 Erivedge

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Erivedge | |
|------------------------|--|
| Basal cell carcinoma | |
| 12 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - ONE of the following:

1.1 Diagnosis of metastatic basal cell carcinoma

OR

1.2 BOTH of the following:

1.2.1 Diagnosis of locally advanced basal cell carcinoma

AND

1.2.2 ONE of the following:

Cancer has recurred following surgery
Patient is not a candidate for surgery
Patient is not a candidate for radiation

| Product Name: Erivedge | | |
|---|-----------------------|--|
| Diagnosis | Medulloblastoma | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| Approval Criteria 1 - Diagnosis of medulloblastoma | | |
| AND | | |
| 2 - Patient has mutations in the sonic hedgehog pathway | | |
| AND | | |
| 3 - Patient has failed prior chemotherapy | | |

| Product Name: Erivedge | | |
|------------------------|---------------------------------------|--|
| Diagnosis | Basal Cell Carcinoma, Medulloblastoma | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |
| | | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Erivedge therapy

| Product Name: Erivedge | |
|--------------------------|--|
| NCCN Recommended Regimen | |
| 12 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Erivedge will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Erivedge | |
|------------------------|--------------------------|
| Diagnosis | NCCN Recommended Regimen |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Erivedge therapy

| Date | Notes |
|----------|--------------------|
| 4/8/2021 | 7/1 Implementation |

Erythropoietic Agents - AZM

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-107211 Erythropoietic Agents - AZM

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 5/16/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Aranesp, Epogen, Procrit, Mircera, Retacrit | | |
|---|--|--|
| Diagnosis | Anemia Due to Chronic Kidney Disease (CKD) | |
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of chronic kidney disease (CKD) | | |

AND

2 - Hematocrit is less than 30% at initiation of therapy

AND

3 - ONE of the following:

3.1 Patient is on dialysis

OR

3.2 ALL of the following:

3.2.1 Patient is NOT on dialysis

AND

3.2.2 The rate of hematocrit decline indicates the likelihood of requiring a red blood cell (RBC) transfusion

AND

3.2.3 Reducing the risk of alloimmunization and/or other RBC transfusion-related risks is a goal

AND

4 - If the request is for Aranesp, Mircera or Procrit, claims history indicates either Epogen or Retacrit has been tried at maximum doses as indicated by FDA labeling

| Product Name: Epogen, Procrit, Retacrit | |
|---|---|
| Diagnosis | Anemia Associated with Zidovudine Treatment in HIV-Infected Patients |

| Approval Length | 12 month(s) | | |
|--|---|--|--|
| Guideline Type | Prior Authorization | | |
| | | | |
| Approval Criteria | | | |
| 1 - Patient is receiving a week | 1 - Patient is receiving zidovudine administered at less than or equal to 4200 milligrams per week | | |
| | AND | | |
| 2 - Endogenous serum erythropoietin level is less than or equal to 500 milliunits per milliliter | | | |
| AND | | | |
| 3 - Hematocrit is less than 30% at initiation of therapy | | | |
| AND | | | |
| 4 - If the request is for Procrit, claims history indicates either Epogen or Retacrit has been tried at maximum doses as indicated by FDA labeling | | | |

| Product Name: Aranesp, Epogen, Procrit, Retacrit | |
|--|-----------------------------------|
| Diagnosis | Anemia Due to Cancer Chemotherapy |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Hematocrit less than 30% at initiation of therapy

AND

2 - There is a minimum of two additional months of planned chemotherapy

AND

3 - If the request is for Aranesp or Procrit, claims history indicates either Epogen or Retacrit has been tried at maximum doses as indicated by FDA labeling

| Product Name: Epogen, Procrit, Retacrit | | |
|--|---|--|
| Diagnosis | Preoperative Use for Reduction of Allogeneic Blood Transfusions in Surgery Patients | |
| Approval Length | 1 month(s) | |
| Guideline Type | Prior Authorization | |
| Approval Criteria 1 - Perioperative hematocrit is greater than 30% and less than or equal to 39% | | |
| | | |
| | AND | |
| 2 - Patient is at high risk for blood loss during surgery | | |
| AND | | |
| 3 - Patient is unable or unwilling to donate autologous blood | | |
| | AND | |
| 4 - Surgery procedure is elective, non-cardiac, and non-vascular | | |
| AND | | |
| 5 - If the request is for Procrit, claims history indicates either Epogen or Retacrit has been tried at maximum doses as indicated by FDA labeling | | |
| | | |

| Product Name: Aranesp, Epogen, Procrit, or Retacrit | |
|---|--|
| Diagnosis | Anemia Associated with Myelodysplastic Disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of myelodysplastic disease (MDS)

AND

2 - ONE of the following:

- Serum erythropoietin level less than or equal to 500 milliunits per milliliter
- Hematocrit is less than or equal to 30% at the initiation of therapy

AND

3 - If the request is for Aranesp or Procrit, claims history indicates either Epogen or Retacrit has been tried at maximum doses as indicated by FDA labeling

| Product Name: Aranesp, Epogen, Procrit, or Retacrit | | |
|---|--|--|
| Diagnosis | Anemia Associated with Myelodysplastic Disease | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |
| | | |

Approval Criteria

- **1** One of the following:
- **1.1** Hematocrit remains less than 36%

OR

1.2 Patient has demonstrated a response to therapy

AND

2 - If the request is for Aranesp or Procrit, claims history indicates either Epogen or Retacrit has been tried at maximum doses as indicated by FDA labeling

| Product Name: Epogen, Procrit, Retacrit | |
|---|---|
| Diagnosis | Anemia in Patients with Hepatitis C with Ribavirin and Interferon Therapy |
| Approval Length | 3 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of hepatitis C virus (HCV) infection

AND

2 - Patient is receiving ribavirin and interferon therapy

AND

3 - Hematocrit is less than or equal to 30% at initiation of therapy

AND

4 - If the request is for Procrit, claims history indicates either Epogen or Retacrit has been tried at maximum doses as indicated by FDA labeling

| Product Name: Epogen, Procrit, Retacrit* | | | |
|--|--|--|--|
| Diagnosis | Anemia in Patients with Hepatitis C with Ribavirin and Interferon Therapy | | |
| Therapy Stage | Reauthorization | | |
| Guideline Type | Prior Authorization | | |
| Approval Criteria | Approval Criteria | | |
| 1 - One of the following | 1 - One of the following: | | |
| 1.1 Hematocrit remains less than 36% | | | |
| OR | | | |
| 1.2 Patient has demonstrated a response to therapy | | | |
| AND | | | |
| 2 - If the request is for Procrit, claims history indicates either Epogen or Retacrit has been tried at maximum doses as indicated by FDA labeling | | | |
| Notes | *NOTE: Authorization will be issued for 12 months or if patient has de monstrated response to therapy, authorization will be issued for the ful I course of ribavirin therapy. | | |

| Product Name: Aranesp, Epogen, Mircera, Procrit, Retacrit* | |
|--|---|
| Diagnosis | Erythropoietin Stimulating Agents –Off-Label Uses |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Off-label requests will be evaluated on a case-by-case basis by a clinical pharmacist

AND

2 - Requests for coverage in patients with hemoglobin (Hgb) greater than 10 grams per deciliter or hematocrit (Hct) greater than 30% will not be approved

| | AND |
|-------|--|
| | Aranesp, Mircera, or Procrit, claims history indicates either Epogen or at maximum doses as indicated by FDA labeling |
| Notes | *If the request is deemed medically necessary, the authorization will b e issued for requested length of therapy. |

| Date | Notes |
|-----------|--|
| 5/16/2022 | Added Epogen as preferred agent (Retacrit OOS from mfg). |

Esbriet, Ofev

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99609 Esbriet, Ofev

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Esbriet, Ofev | |
|-----------------------------|-------------------------------|
| Diagnosis | Idiopathic Pulmonary Fibrosis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of idiopathic pulmonary fibrosis (IPF) as documented by ALL of the following criteria:

1.1 Exclusion of other known causes of interstitial lung disease (e.g. domestic and occupational environmental exposures, connective tissue disease, and drug toxicity), as documented by the following: ICD-10 Code J84.112 (Idiopathic pulmonary fibrosis) • AND **1.2** ONE of the following: 1.2.1 In patients NOT subjected to surgical lung biopsy, the presence of a usual interstitial pneumonia (UIP) pattern on high-resolution computed tomography (HRCT) revealing IPF or probable IPF OR 1.2.2 In patients subjected to a lung biopsy, both HRCT and surgical lung biopsy pattern reveal IPF or probable IPF AND 2 - The agent is not being used in combination with Esbriet or Ofev AND **3** - The prescriber is a pulmonologist

| Product Name: Esbriet or Ofev | | |
|-------------------------------|-------------------------------|--|
| Diagnosis | Idiopathic Pulmonary Fibrosis | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |

1 - Documentation of positive clinical response to Esbriet or Ofev therapy

AND

2 - The agent is not being used in combination with Esbriet or Ofev

AND

3 - The prescriber is a pulmonologist

| Product Name: Ofev | |
|--------------------|---|
| Diagnosis | Systemic Sclerosis-Associated Interstitial Lung Disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of systemic sclerosis (SSc) - associated interstitial lung disease as documented by ALL of the following:

1.1 ONE of the following:

1.1.1 Skin thickening of the fingers of both hands extending proximal to the metacarpophalangeal joints

OR

1.1.2 TWO of the following:

- Skin thickening of the fingers (e.g., puffy fingers, sclerodactyly of the fingers)
- Fingertip lesions (e.g., digital tip ulcers, fingertip pitting scars)
- Telangiectasia
- Abnormal nailfold capillaries
- Pulmonary arterial hypertension
- Raynaud's phenomenon

 SSc-related autoantibodies (e.g., anticentromere, anti-topoisomerase I, anti-RNA polymerase III)

AND

1.2 Presence of interstitial lung disease as determined by finding evidence of pulmonary fibrosis on high-resolution computed tomography (HRCT), involving at least 10 percent of the lungs

AND

2 - The agent is not being used in combination with Esbriet

AND

3 - The prescriber is a pulmonologist

| Product Name: Ofev | |
|--------------------|--|
| Diagnosis | Chronic fibrosing interstitial lung disease with a progressive phenotype |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of chronic fibrosing interstitial lung disease (ILD) with a progressive phenotype as documented by BOTH of the following criteria:

1.1 Presence of fibrotic ILD as determined by finding evidence of pulmonary fibrosis on HRCT (high-resolution computed tomography), involving at least 10 percent of the lungs

AND

1.2 Patient is presenting with clinical signs of progression as defined by ONE of the following in the previous 24 months:

| 1.2.1 Forced vital capacity (FVC) decline of greater than 10 percent |
|---|
| OR |
| 1.2.2 TWO of the following: |
| FVC decline of greater than or equal to 5 percent, but less than 10 percent Patient is experiencing worsening respiratory symptoms Patient is exhibiting increasing extent of fibrotic changes on chest imaging |
| AND |
| 2 - The agent is not being used in combination with Esbriet |
| AND |
| 3 - The prescriber is a pulmonologist |

| Product Name: Ofev | |
|--------------------|---|
| Diagnosis | Systemic Sclerosis-Associated Interstitial Lung Disease, Chronic fibrosing interstitial lung disease with a progressive phenotype |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Ofev therapy

AND

2 - Ofev is not being used in combination with Esbriet

| | AND | |
|--|-----|--|
| 3 - The prescriber is a pulmonologist | | |

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona SP to Medicaid Arizona SP for 7/1 eff |

Estrogens- Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99455 Estrogens- Arizona

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Femring | |
|---|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of moderate to severe vasomotor symptoms due to menopause | |
| | |
| OR | |
| | |

2 - Diagnosis of moderate to severe vulvar and vaginal atrophy due to menopause

| Product Name: Premarin | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | · |
| Approval Criteria | |
| 1 - Diagnosis of atrophic vaginitis and kraurosis vulvae | |

| Date | Notes |
|-----------|---|
| 3/10/2021 | Bulk Copied C&S Arizona standard to Arizona Medicaid for 7/1 effect ive |

Etoposide- Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99540 | Etoposide- Arizona |
|----------|--------------------|
|----------|--------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: etoposide | |
|---|------------------------|
| Diagnosis | Small cell lung cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of small cell lung cancer | |

AND

2 - Used as first-line therapy with other approved chemotherapeutic agents

| Product Name: etoposide | |
|---------------------------|--|
| NCCN Recommended Regimens | |
| 12 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| i | |

Approval Criteria

1 - Use is supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: etoposide | |
|-------------------------|---|
| Diagnosis | Small cell lung cancer, NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to therapy

| Date | Notes |
|----------|-------------------------------------|
| 6/2/2021 | Arizona Medicaid 7.1 Implementation |

Eucrisa

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99456 Eucrisa

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Eucrisa | | |
|--|--------------|--|
| Approval Length | 12 month(s) | |
| Guideline Type | Step Therapy | |
| | | |
| Approval Criteria | | |
| 1 - BOTH of the following: | | |
| 1.1 History of failure, contraindication, or intolerance to ONE topical corticosteroid [e.g., mometasone furoate, fluocinolone acetonide (generic Synalar), fluocinonide] | | |

AND

1.2 ONE of the following:

1.2.1 Patient is less than 2 years of age

OR

1.2.2 Patient is greater than or equal to 2 years of age and has history of failure, contraindication, or intolerance to ONE topical calcineurin inhibitor [e.g., pimecrolimus (generic Elidel), tacrolimus (generic Protopic)]

| Date | Notes |
|-----------|---|
| 3/10/2021 | Bulk Copied C&S Arizona standard to Arizona Medicaid for 7/1 effect ive |

Evrysdi (risdiplam)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-114470 | Evrysdi (risdiplam) |
|-----------|---------------------|
|-----------|---------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Evrysdi | |
|-----------------------|-------------------------------|
| Diagnosis | Spinal Muscular Atrophy (SMA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| | |

1 - Diagnosis of spinal muscular atrophy (SMA)

AND 2 - Submission of medical records (e.g., chart notes, laboratory values) confirming the mutation or deletion of genes in chromosome 5g resulting in ONE of the following: 2.1 Homozygous gene deletion or mutation of SMN1 gene (e.g., homozygous deletion of exon 7 at locus 5q13) OR 2.2 Compound heterozygous mutation of SMN1 gene [e.g., deletion of SMN1 exon 7 (allele 1) and mutation of SMN1 (allele 2)] AND 3 - Patient is not dependent on invasive ventilation or tracheostomy AND 4 - Patient is not dependent on the use of non-invasive ventilation beyond use for naps and nighttime sleep AND 5 - Patient is not receiving concomitant chronic survival motor neuron (SMN)-modifying therapy [e.g., Spinraza (nusinersen)] AND 6 - Patient has not previously received gene replacement therapy for the treatment of SMA [e.g., Zolgensma (onasemnogene abeparvovec-xioi)] AND

7 - Submission of medical records (e.g., chart notes, laboratory values) documenting the baseline assessment of at least ONE of the following exams (based on patient age and motor ability) to establish baseline motor ability (baseline motor function analysis could include assessments evaluated prior to receipt of previous chronic SMN-modifying therapy if transitioning therapy)*:

- Children's Hospital of Philadelphia Infant Test of Neuromuscular Disorders (CHOP INTEND)
- Hammersmith Infant Neurological Exam Part 2 (HINE-2)
- Hammersmith Functional Motor Scale Expanded (HFMSE)
- Upper Limb Module (ULM) Test
- Motor Function Measure 32 (MFM-32) Scale

AND

8 - Prescribed by a neurologist with expertise in the treatment of SMA

| *Baseline assessments for patients less than 2 months of age request ing Evrysdi are not necessary in order not to delay access to initial the rapy in recently diagnosed infants. Initial assessments shortly post-the rapy can serve as baseline with respect to efficacy reauthorization ass |
|--|
| essment. |

| Product Name: Evrysdi | |
|-----------------------|-------------------------------|
| Diagnosis | Spinal Muscular Atrophy (SMA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, laboratory values) with the most recent results documenting a positive clinical response to Evrysdi compared to pretreatment baseline status [inclusive of baseline assessments prior to receipt of previous chronic survival motor neuron (SMN)-modifying therapy] as demonstrated by at least ONE of the following exams:

1.1 Children's Hospital of Philadelphia Infant Test of Neuromuscular Disorders (CHOP INTEND) with ONE of the following:

1.1.1 Improvement or maintenance of previous improvement of at least a 4-point increase in score from pretreatment baseline

OR

1.1.2 Patient has achieved and maintained any new motor milestone from pretreatment baseline when they would otherwise be unexpected to do so

OR

1.2 Hammersmith Infant Neurological Exam Part 2 (HINE-2) with ONE of the following:

1.2.1 Improvement or maintenance of previous improvement of at least a 2-point (or maximal score) increase in ability to kick

OR

1.2.2 Improvement or maintenance of previous improvement of at least a 1-point increase in any other HINE-2 milestone (e.g., head control, rolling, sitting, crawling, etc.), excluding voluntary grasp

OR

1.2.3 The patient exhibited improvement, or maintenance of previous improvement, in more HINE motor milestones than worsening, from pretreatment baseline (net positive improvement)

OR

1.2.4 Patient has achieved and maintained any new motor milestones when they would otherwise be unexpected to do so

OR

1.3 Hammersmith Functional Motor Scale Expanded (HFMSE) with ONE of the following:

1.3.1 Improvement or maintenance of previous improvement of at least a 3-point increase in score from pretreatment baseline

OR

1.3.2 Patient has achieved and maintained any new motor milestone from pretreatment baseline when they would otherwise be unexpected to do so

OR

1.4 Upper Limb Module (ULM) with ONE of the following:

1.4.1 Improvement or maintenance of previous improvement of at least a 2-point increase in score from pretreatment baseline

OR

1.4.2 Patient has achieved and maintained any new motor milestone from pretreatment baseline when they would otherwise be unexpected to do so

OR

1.5 Motor Function Measure 32 (MFM-32) with ONE of the following:

1.5.1 Improvement or maintenance of previous improvement of at least a 3-point increase in score from pretreatment baseline

OR

1.5.2 Patient has achieved and maintained any new motor milestone from pretreatment baseline when they would otherwise be unexpected to do so

AND

2 - Patient is not dependent on invasive ventilation or tracheostomy

AND

| 3 - Patient is not dependent on the use of non-invasive ventilation beyond use for naps and nighttime sleep |
|--|
| AND |
| 4 - Patient is not receiving concomitant chronic SMN-modifying therapy [e.g., Spinraza (nusinersen)] |
| AND |
| 5 - Patient has not previously received gene replacement therapy for the treatment of spinal muscular atrophy (SMA) [e.g., Zolgensma (onasemnogene abeparvovec-xioi)] |
| AND |
| 6 - Prescribed by a neurologist with expertise in the treatment of SMA |

| Date | Notes |
|-----------|---|
| 9/26/2022 | Updated to remove age requirement per FDA expanded indication |

Exondys- Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99457 Exondys- Arizona

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Exondys | |
|--|-----------------------|
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of Duchenne muscular dystrophy (DMD) | |

| AND |
|--|
| 2 - Documentation of a confirmed mutation of the dystrophin gene amenable to exon 51 skipping |
| AND |
| 3 - Patient is 7 years of age or older |
| AND |
| 4 - Patient is ambulatory |
| AND |
| 5 - Prescribed by or in consultation with a neurologist who has experience treating children |
| AND |
| 6 - Dose will not exceed 30 milligrams per kilogram of body weight once weekly |
| AND |
| 7 - Patient's condition has been evaluated via the 6-minute walk test (6MWT) or North Star ambulatory assessment (NSAA) [documentation of the patient's most recent results must be provided] |

| Product Name: Exondys | |
|-----------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - One of the following:

1.1 Patient has been on therapy for less than 12 months and all of the following:

1.1.1 Patient is maintaining ambulatory status

AND

1.1.2 Patient is tolerating therapy

AND

1.1.3 Dose will not exceed 30 milligrams per kilogram of body weight once weekly

AND

1.1.4 Prescribed by or in consultation with a neurologist who has experience treating children

AND

1.1.5 Patient's condition has been evaluated via the 6-minute walk test (6MWT) or North Star ambulatory assessment (NSAA) [documentation of the patient's most recent results must be provided]

OR

1.2 Patient has been on therapy for 12 months or more and all of the following:

1.2.1 Patient has experienced a benefit from therapy (e.g., disease amelioration compared to untreated patients)

AND

1.2.2 Patient is maintaining ambulatory status

AND

1.2.3 Patient is tolerating therapy

AND

1.2.4 Dose will not exceed 30 milligrams per kilogram of body weight once weekly

AND

1.2.5 Prescribed by or in consultation with a neurologist who has experience treating children

AND

1.2.6 Patient's condition has been evaluated via the 6-minute walk test (6MWT) or North Star ambulatory assessment (NSAA) [documentation of the patient's most recent results must be provided]

| Date | Notes |
|-----------|---|
| 3/10/2021 | Bulk Copied C&S Arizona standard to Arizona Medicaid for 7/1 effect ive |

Fabrazyme- Arizona

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99610 | Fabrazyme- Arizona |
|----------|--------------------|
|----------|--------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Fabrazyme | |
|--------------------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of Fabry disease | |

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona SP to Medicaid Arizona SP for 7/1 eff |

Fasenra

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99716 Fasenra

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Fasenra Pen | |
|---------------------------|-----------------------|
| Diagnosis | Asthma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| | |

1 - Diagnosis of severe asthma

AND

2 - Classification of asthma as uncontrolled or inadequately controlled as defined by ONE of the following:

2.1 Poor symptom control (e.g., Asthma Control Questionnaire [ACQ] score consistently greater than 1.5 or Asthma Control Test [ACT] score consistently less than 20)

OR

2.2 Two or more bursts of systemic corticosteroids for at least 3 days each in the previous 12 months

OR

2.3 Asthma-related emergency treatment (e.g., emergency room visit, hospital admission, or unscheduled physician's office visit for nebulizer or other urgent treatment)

OR

2.4 Airflow limitation (e.g., after appropriate bronchodilator withhold forced expiratory volume in 1 second [FEV1] less than 80 percent predicted [in the face of reduced FEV1-forced vital capacity [FVC] defined as less than the lower limit of normal])

OR

2.5 Patient is currently dependent on oral corticosteroids for the treatment of asthma

AND

3 - Submission of medical records (e.g., chart notes, laboratory values, etc.) documenting ONE of the following:

3.1 Asthma is an eosinophilic phenotype as defined by a baseline (pre-benralizumab treatment) peripheral blood eosinophil level greater than or equal to 150 cells per microliter within the past 6 weeks

OR

3.2 Patient is currently dependent on maintenance therapy with oral corticosteroids for the treatment of asthma

AND

4 - Fasenra will be used in combination with ONE of the following:

4.1 One high dose (appropriately adjusted for age) combination inhaled corticosteroid (ICS)/long-acting beta2 agonist (LABA) [e.g., Advair/AirDuo Respiclick (fluticasone propionate/salmeterol), Symbicort (budesonide/formoterol), Breo Ellipta (fluticasone furoate/vilanterol)]

OR

4.2 Combination therapy including BOTH of the following:

4.2.1 One high-dose (appropriately adjusted for age) ICS product [e.g., ciclesonide (Alvesco), mometasone furoate (Asmanex), beclomethasone dipropionate (QVAR)]

AND

4.2.2 One additional asthma controller medication [e.g., LABA - olodaterol (Striverdi) or indacaterol (Arcapta); leukotriene receptor antagonist – montelukast (Singulair); theophylline]

AND

5 - Patient is not receiving Fasenra in combination with one of the following:

- Anti-interleukin 5 therapy [e.g., Cinqair (resilizumab), Nucala (mepolizumab)]
- Anti-IgE therapy [e.g., Xolair (omalizumab)]
- Anti-interleukin 4 therapy [e.g., Dupixent (dupilumab)]

AND

- **6** Prescribed by one of the following:
 - Pulmonologist
 - Allergist
 - Immunologist

| Product Name: Fasenra Pen | |
|---------------------------|---------------------|
| Diagnosis | Asthma |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response as demonstrated by ONE of the following:

- Reduction in the frequency of exacerbations
- Decreased utilization of rescue medications
- Increase in percent predicted FEV1 (forced expiratory volume in 1 second) from pretreatment baseline
- Reduction in severity or frequency of asthma-related symptoms (e.g., wheezing, shortness of breath, coughing, etc.)
- Reduction in oral corticosteroid requirements

AND

2 - Used in combination with an inhaled corticosteroid (ICS)-containing controller medication

AND

3 - Patient is not receiving Fasenra in combination with one of the following:

- Anti-interleukin 5 therapy [e.g., Cinqair (resilizumab), Nucala (mepolizumab)]
- Anti-IgE therapy [e.g., Xolair (omalizumab)]
- Anti-interleukin 4 therapy [e.g., Dupixent (dupilumab)]

AND 4 - Prescribed by one of the following: • Pulmonologist • Allergist • Immunologist

| Date | Notes |
|----------|-------------------------------------|
| 6/8/2021 | Arizona Medicaid 7.1 Implementation |

Fentanyl IR

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99519 Fentanyl IR

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Fentanyl citrate lozenges (generic Actiq) | | |
|---|--|--|
| 12 month(s) | | |
| Prior Authorization | | |
| | | |
| | | |

Approval Criteria

1 - Submission of medical records demonstrating use is for the management of breakthrough pain associated with a cancer diagnosis (cancer diagnosis must be documented)

AND

2 - Patient must have at least a one week history of ONE of the following medications to demonstrate tolerance to opioids (Document drug and date of trial):

- Morphine sulfate at a doses of greater than or equal to 60 milligrams per day
- Fentanyl transdermal patch at a dose of greater than or equal to 25 micrograms per hour
- Oxycodone at a dose of greater than or equal to 30 milligrams per day
- Oral hydromorphone at a dose of greater than or equal to 8 milligrams per day
- Oral oxymorphone at a dose of greater than or equal to 25 milligrams per day
- An alternative opioid at an equianalgesic dose (e.g., oral methadone greater than or equal to 20 milligrams per day)

AND

3 - The patient is currently taking a long-acting opioid around the clock for cancer pain (Document drug)

AND

4 - ONE of the following:

4.1 The patient is not concurrently receiving an alternative fentanyl transmucosal product

OR

4.2 BOTH of the following:

4.2.1 The patient is currently receiving an alternative transmucosal fentanyl product

AND

4.2.2 The prescriber is requesting the termination of all current authorizations for alternative transmucosal fentanyl products in order to begin treatment with the requested medication (Only one transmucosal fentanyl product will be approved at a time. If previous authorizations cannot be terminated, the PA request will be denied)

Product Name: Abstral, Brand Actiq, Brand Fentora, generic fentanyl citrate buccal tablet, Lazanda, Subsys

| Approval Length | 12 month(s) | |
|--|--|--|
| Guideline Type | Prior Authorization | |
| Approval Criteria 1 - Submission of medical records demonstrating use is for the management of breakthrough pain associated with a cancer diagnosis (cancer diagnosis must be documented) | | |
| | AND | |
| | at least a one week history of ONE of the following medications to to opioids (Document drug and date of trial): | |
| Morphine sulfate at a doses of greater than or equal to 60 milligrams per day Fentanyl transdermal patch at a dose of greater than or equal to 25 micrograms per hour | | |
| Oxycodone at a dose of greater than or equal to 30 milligrams per day Oral hydromorphone at a dose of greater than or equal to 8 milligrams per day Oral oxymorphone at a dose of greater than or equal to 25 milligrams per day An alternative opioid at an equianalgesic dose (e.g., oral methadone greater than or equal to 20 milligrams per day) | | |
| | AND | |
| 3 - The patient is currently taking a long-acting opioid around the clock for cancer pain (Document drug) | | |
| | AND | |
| 4 - ONE of the following: | | |
| 4.1 The patient is not concurrently receiving an alternative fentanyl transmucosal product | | |
| OR | | |
| 4.2 BOTH of the following: | | |
| 4.2.1 The patient is currently receiving an alternative transmucosal fentanyl product | | |
| | | |

AND

4.2.2 The prescriber is requesting the termination of all current authorizations for alternative transmucosal fentanyl products in order to begin treatment with the requested medication (Only one transmucosal fentanyl product will be approved at a time. If previous authorizations cannot be terminated, the PA request will be denied)

AND

5 - History of failure, contraindication, or intolerance to Fentanyl citrate lozenges (generic Actiq) [Document date of trial]

| Date | Notes |
|----------|-------------------------------------|
| 6/8/2021 | Arizona Medicaid 7.1 Implementation |

Fexmid (cyclobenzaprine 7.5mg)- Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99458 Fexmid (cyclobenzaprine 7.5mg)- Arizona

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Fexmid 7.5mg, generic cyclobenzaprine 7.5mg | | |
|---|---------------------|--|
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of muscle spasm associated with acute, painful musculoskeletal conditions | | |
| | | |
| AND | | |
| | | |

2 - Reason or special circumstance the patient cannot use cyclobenzaprine 5 milligram (mg) or 10mg tablet

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk copy C&S Arizona standard to Medicaid Arizona |

Firazyr

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99611 Firazyr

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Firazyr, generic icatibant | |
|--|-----------------------------|
| Diagnosis | Hereditary angioedema (HAE) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of hereditary angioedema (HAE) as confirmed by one of the following:

| 1.1 C1 inhibitor (C1-INH) deficiency or dysfunction (Type I or II HAE) as documented by ONE of the following (per laboratory standard): | |
|---|--|
| C1-INH antigenic level below the lower limit of normal C1-INH functional level below the lower limit of normal | |
| OR | |
| 1.2 HAE with normal C1 inhibitor levels and ONE of the following: | |
| Confirmed presence of a FXII, angiopoietin-1 or plasminogen gene mutation Recurring angioedema attacks that are refractory to high-dose antihistamines with confirmed family history of angioedema | |
| AND | |
| 2 - Prescribed for the acute treatment of HAE attacks | |
| AND | |
| 3 - Not used in combination with other products indicated for the acute treatment of HAE attacks (e.g., Berinert, Ruconest) | |
| AND | |
| 4 - Prescribed by ONE of the following: | |
| ImmunologistAllergist | |

Г

| Product Name: Brand Firazyr, generic icatibant | |
|--|-----------------------------|
| Diagnosis | Hereditary angioedema (HAE) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

| Approval Critoria |
|--|
| Approval Criteria |
| 1 - Documentation of positive clinical response |
| |
| AND |
| 2 - Prescribed for the acute treatment of hereditary angioedema (HAE) attacks |
| AND |
| 3 - Not used in combination with other products indicated for the acute treatment of HAE attacks (e.g., Berinert, Ruconest) |
| AND |
| 4 - Prescribed by ONE of the following: |
| ImmunologistAllergist |

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona SP to Medicaid Arizona SP for 7/1 eff |

Firdapse

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-108661 Firdapse

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 6/23/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Firdapse | |
|------------------------|--|
| Diagnosis | Lambert-Eaton myasthenic syndrome (LEMS) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| | |

Approval Criteria

1 - Patient has a diagnosis of Lambert-Eaton myasthenic syndrome (LEMS)

AND

2 - Patient is not receiving Firdapse in combination with similar potassium channel blockers [e.g.,Ampyra (dalfampridine), Ruzurgi (amiframpridine)]

| Product Name: Firdapse | |
|------------------------|--|
| Diagnosis | Lambert-Eaton myasthenic syndrome (LEMS) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Firdapse therapy

AND

2 - Patient is not receiving Firdapse in combination with similar potassium channel blockers [e.g., Ampyra (dalfampridine), Ruzurgi (amifampridine)]

| Date | Notes |
|-----------|---------------------------------|
| 6/23/2022 | Removed Ruzurgi as prerequisite |

Flucytosine- Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Ancobon, generic flucytosine | | |
|--|---------------------|--|
| Approval Length | 2 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - One of the following: | | |
| 1.1 Diagnosis of septicemia, endocarditis or a urinary system infection caused by Candida species | | |

OR

1.2 Diagnosis of meningitis or a pulmonary infection caused by Cryptococcus species

AND

2 - If the patient is being treated for a systemic infection, flucytosine is being used in combination with amphotericin B

| Product Name: Brand Ancobon, generic flucytosine* | |
|---|---|
| Diagnosis | Infectious Diseases Society of America (IDSA) Recommended Regimens |
| Guideline Type | Prior Authorization |
| Approval Criteria 1 - The medication is being prescribed by or in consultation with an infectious disease specialist. | |
| Notes | *Approval duration based on provider recommended treatment duratio ns, up to 12 months. |

| Date | Notes |
|-----------|-------------------------------------|
| 5/13/2021 | Arizona Medicaid 7.1 Implementation |

Forteo-Arizona

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99790 | Forteo-Arizona |
|----------|----------------|
|----------|----------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Forteo | |
|-------------------------------|--|
| Diagnosis | Patients with Osteoporosis at High Risk for Fracture |
| Approval Length | 24 months * |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of osteoporosis | |

AND

2 - ONE of the following:

2.1 Bone Mineral Density (BMD) T-score less than or equal to -3.5 based on BMD measurements from lumber spine (at least two vertebral bodies), hip (femoral neck, total hip), or radius (one-third radius site). [Provider must submit patient specific BMD T-score]

OR

2.2 BOTH of the following:

2.2.1 BMD T-score between -2.5 and -3.5 (BMD T-score greater than -3.5 and less than or equal to -2.5) based on BMD measurements from lumber spine (at least two vertebral bodies), hip (femoral neck, total hip), or radius (one-third radius site). [Provider must submit patient specific BMD T-score]

AND

2.2.2 ONE of the following:

2.2.2.1 History of ONE of the following resulting from minimal trauma:

- Vertebral compression fracture
- Fracture of the hip
- Fracture of the distal radius
- Fracture of the pelvis
- Fracture of the proximal humerus

OR

2.2.2.2 History of failure, contraindication, or intolerance to ONE conventional osteoporosis therapy [e.g., bisphosphonate or selective estrogen receptor modulator (SERM)] (Document drug, date, and duration of trial)**

OR

2.3 ALL of the following:

2.3.1 BMD T-score between -1 and -2.5 (BMD T-score greater than -2.5 and less than or equal to -1) based on BMD measurements from lumber spine (at least two vertebral bodies), hip (femoral neck, total hip), or radius (one-third radius site). [Provider must submit patient specific BMD T-score]

AND

2.3.2 ONE of the following:

2.3.2.1 History of ONE of the following resulting from minimal trauma:

- Vertebral compression fracture
- Fracture of the hip
- Fracture of the distal radius
- Fracture of the pelvis
- Fracture of the proximal humerus

OR

2.3.2.2 ONE of the following FRAX 10-year probabilities:

- Major osteoporotic fracture at 20% or more
- Hip fracture at 3% or more

AND

2.3.3 History of failure, contraindication, or intolerance to one conventional osteoporosis therapy [e.g., bisphosphonate or selective estrogen receptor modulator (SERM)] (Document drug, date, and duration of trial)**

AND

3 - Treatment duration has not exceeded a total of 24 months of cumulative use of parathyroid hormone analogs (e.g., Forteo, Tymlos) during the patient's lifetime

AND

4 - Prescriber attests to the following: the information provided is true and accurate to the best

| of their knowledge and they understand that OptumRx may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided | |
|--|--|
| Notes | *Authorization will be issued for 24 months. Duration of coverage will be limited to 24 months of cumulative use of parathyroid hormone ana logs (e.g., Forteo, Tymlos) in the member's lifetime. **Note: Claims hi story may be used in conjunction as documentation of drug, date, and duration of trial |

| Product Name: Forteo | |
|----------------------|---|
| Diagnosis | Glucocorticoid-Induced Osteoporosis at High Risk for Fracture |
| Approval Length | 24 months * |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of glucocorticoid-induced osteoporosis

AND

2 - History of prednisone or its equivalent at a dose greater than or equal to 5 milligrams per day for greater than or equal to 3 months

AND

3 - ONE of the following:

3.1 Bone Mineral Density (BMD) T-score less than or equal to -2.0 based on BMD measurements from lumber spine (at least two vertebral bodies), hip (femoral neck, total hip), or radius (one-third radius site). [Provider must submit patient specific BMD T-score]

OR

3.2 BOTH of the following:

3.2.1 BMD T-score between -1.0 and -2.0 (BMD T-score greater than -2.0 and less than or equal to -1.0) based on BMD measurements from lumber spine (at least two vertebral bodies), hip (femoral neck, total hip), or radius (one-third radius site). [Provider must submit patient specific BMD T-score]

3.2.2 ONE of the following:

3.2.2.1 History of ONE of the following resulting from minimal trauma:

- Vertebral compression fracture
- Fracture of the hip
- Fracture of the distal radius
- Fracture of the pelvis
- Fracture of the proximal humerus

OR

3.2.2.2 History of failure, contraindication, or intolerance to ONE conventional osteoporosis therapy [e.g., bisphosphonate or selective estrogen receptor modulator (SERM)] (Document drug, date, and duration of trial)**

OR

3.2.3 BOTH of the following:

3.2.3.1 History of ONE of the following resulting from minimal trauma:

- Vertebral compression fracture
- Fracture of the hip
- Fracture of the distal radius
- Fracture of the pelvis
- Fracture of the proximal humerus

AND

3.2.3.2 History of failure, contraindication, or intolerance to one conventional osteoporosis therapy [e.g., bisphosphonate or selective estrogen receptor modulator (SERM)] (Document drug, date, and duration of trial)**

AND

4 - Treatment duration has not exceeded a total of 24 months of cumulative use of parathyroid hormone analogs (e.g., Forteo, Tymlos) during the patient's lifetime

AND

5 - Prescriber attests to the following: the information provided is true and accurate to the best of their knowledge and they understand that OptumRx may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided

| *Authorization will be issued for 24 months. Duration of coverage will be limited to 24 months of cumulative use of parathyroid hormone ana logs (e.g., Forteo, Tymlos) in the member's lifetime. **Note: Claims hi story may be used in conjunction as documentation of drug, date, and |
|---|
| duration of trial |

| Date | Notes |
|-----------|------------------|
| 6/10/2021 | Update Guideline |

Galafold

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99613 Galafold

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Galafold | |
|------------------------|-----------------------|
| Diagnosis | Fabry disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| | |

1 - Diagnosis of Fabry disease

2 - Patient has an amenable galactosidase alpha gene (GLA) variant based on in vitro assay data

AND

3 - Patient is not receiving Galafold in combination with Fabrazyme (agalsidase beta)

| Product Name: Galafold | |
|------------------------|---------------------|
| Diagnosis | Fabry disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | • |

Approval Criteria

1 - Documentation of positive clinical response to Galafold therapy

AND

2 - Patient is not receiving Galafold in combination with Fabrazyme (agalsidase beta)

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona SP to Medicaid Arizona SP for 7/1 eff |

Gattex

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99614 Gattex

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Gattex | |
|----------------------|----------------------------|
| Diagnosis | Short Bowel Syndrome (SBS) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of Sho | rt Bowel Syndrome (SBS) |

| A | Ν | D |
|---|---|---|
| | | |

2 - Dependent on parenteral support

| Product Name: Gattex | |
|----------------------|----------------------------|
| Diagnosis | Short Bowel Syndrome (SBS) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Documentation of positive clinical response to Gattex therapy

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona SP to Medicaid Arizona SP for 7/1 eff |

Gaucher's Disease Agents- Arizona

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99615 Gaucher's Disease Agents- Arizona

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Cerdelga | | |
|---|--------------------------|--|
| Diagnosis | Type 1 Gaucher's disease | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of Type 1 Gaucher's disease | | |

2 - Patient is one of the following as detected by a Food and Drug Administration (FDA)cleared test:

- CYP2D6 extensive metabolizer,
- CYP2D6 intermediate metabolizer
- CYP2D6 poor metabolizer

| Product Name: Cerezyme | |
|------------------------|--------------------------|
| Diagnosis | Type 1 Gaucher's disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of Type 1 Gaucher's disease that results in one or more of the following conditions:

- Anemia
- Thrombocytopenia
- Bone disease
- Hepatomegaly or splenomegaly

| Product Name: Vpriv, Elelyso | |
|------------------------------|--|
| Type 1 Gaucher's disease | |
| 12 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Diagnosis of Type 1 Gaucher's disease

| Product Name: Brand Zavesca, generic miglustat | |
|--|--------------------------|
| Diagnosis | Type 1 Gaucher's disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of mild to moderate Type 1 Gaucher's disease

AND

2 - If the request is for generic miglustat, there is a reason or special circumstance why the patient cannot use brand Zavesca

| Product Name: Cerdelga, Cerezyme, Elelyso, Vpriv, Brand Zavesca, generic miglustat | |
|--|--------------------------|
| Diagnosis | Type 1 Gaucher's disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to therapy

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona SP to Medicaid Arizona SP for 7/1 eff |

Generic fluticasone-salmeterol diskus, Wixela Inhub (authorized generic of Advair Diskus), Airduo, fluticasone/salmeterol (authorized generic of Airduo)

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99573 Generic fluticasone-salmeterol diskus, Wixela Inhub (authorized generic of Advair Diskus), Airduo, fluticasone/salmeterol (authorized generic of Airduo)

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Generic fluticasone-salmeterol diskus, Wixela Inhub (authorized generic of Advair Diskus), Airduo, fluticasone/salmeterol (authorized generic of Airduo) | | |
|--|---|--|
| Guideline Type | Prior Authorization | |
| Advair Diskus), Airduo, | ic fluticasone-salmeterol diskus, Wixela Inhub (authorized generic of fluticasone/salmeterol (authorized generic of Airduo) should be erred products are brand Advair Diskus, Advair HFA, Dulera, | |

Generic tretinoin cream and gel, generic Avita cream and gel, generic atralin gel

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99577 gel | Generic tretinoin cream and gel, generic Avita cream and gel, generic atralin |
|-----------------|---|
| Formulary | Medicaid - Arizona (AZM, AZMREF, AZMDDD) |
| Formulary No | ote |

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: generic tretinoin cream and gel, generic Avita cream and gel, generic atralin gel | | |
|---|--|--|
| Guideline Type | Prior Authorization | |
| Approval Criteria | | |
| 1 - Requests for generic tretinoin cream and gel, generic Avita cream and gel, generic atralin gel should be denied. The plan's preferred product is Brand Retin-A cream or gel.* | | |
| Notes | *Brand Retin-A cream or gel may require PA | |

| Date | Notes |
|-----------|--------------------|
| 7/21/2021 | 7/1 Implementation |

Gleevec - Arizona

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99805 Gleevec - Arizona

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Generic imatinib | |
|--------------------------------|---|
| Diagnosis | Chronic myelogenous or myeloid leukemia (CML) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of chronic myelogenous or myeloid leukemia (CML)

2 - History of failure, intolerance, or contraindication to Brand Gleevec.

| Product Name: Generic imatinib | |
|--------------------------------|--|
| Diagnosis | Philadelphia chromosome-positive acute lymphoblastic leukemia (Ph+ALL) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of Philadelphia chromosome-positive acute lymphoblastic leukemia (Ph+ALL)

AND

2 - History of failure, intolerance, or contraindication to Brand Gleevec.

| Product Name: Generic imatinib | |
|--------------------------------|---|
| Diagnosis | Myelodysplastic Disease (MDS) or Myeloproliferative Disease (MPD) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of myelodysplastic disease or myeloproliferative disease (MDS/MPD)

AND

2 - ONE of the following:

- •
- Disease is associated with 5q31-33 (gene) translocations Disease is associated with platelet-derived growth factor receptor (PDGRF) beta gene • re-arrangements

3 - History of failure, intolerance, or contraindication to Brand Gleevec.

| Product Name: Generic imatinib | | |
|--|--|--|
| Diagnosis | Aggressive Systemic Mastocytosis (ASM) | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| Approval Criteria 1 - Diagnosis of aggressive systemic mastocytosis (ASM) | | |
| AND | | |
| 2 - ONE of the following: | | |

- Patient is without the D816V c-Kit (gene)mutation •
- c-Kit mutational status unknown

AND

3 - History of failure, intolerance, or contraindication to Brand Gleevec.

| Product Name: Generic imatinib | |
|--------------------------------|--|
| Diagnosis | Hypereosinophilic Syndrome (HES) / Chronic Eosinophilic Leukemia (CEL) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |

| Guideline Type | Prior Authorization | |
|---|---------------------|--|
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of at least ONE of the following: | | |
| Hypereosinophilic syndrome (HES) Chronic eosinophilic leukemia (CEL) | | |
| AND | | |

2 - History of failure, intolerance, or contraindication to Brand Gleevec.

| Product Name: Generic imatinib | |
|--------------------------------|--|
| Diagnosis | Dermatofibrosarcoma Protuberans (DFSP) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of dermatofibrosarcoma protuberans (DFSP)

AND

2 - History of failure, intolerance, or contraindication to Brand Gleevec.

| Product Name: Generic imatinib | |
|--------------------------------|-----------------------|
| Diagnosis | Soft Tissue Sarcoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

- **1** Diagnosis of ONE of the following:
 - Gastrointestinal stromal tumors (GIST)
 - Desmoid tumors / aggressive fibromatosis
 - Pigmented villonodular synovitis (PVNS) or tenosynovial giant cell tumor (TGCT)

2 - History of failure, intolerance, or contraindication to Brand Gleevec

| Product Name: Generic imatinib | |
|--------------------------------|-----------------------|
| Diagnosis | Chordoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of chordoma:

AND

2 - History of failure, intolerance, or contraindication to Brand Gleevec.

| Product Name: Generic imatinib | | |
|--------------------------------|-----------------------|--|
| Diagnosis | Melanoma | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |

1 - Diagnosis of melanoma

2 - Patient has C-KIT (gene) mutation

AND

3 - History of failure, intolerance, or contraindication to Brand Gleevec.

| Product Name: Generic | Product Name: Generic imatinib | | |
|--|--|--|--|
| Diagnosis | AIDS-Related Kaposi Sarcoma | | |
| Approval Length | 12 month(s) | | |
| Therapy Stage | Initial Authorization | | |
| Guideline Type | Prior Authorization | | |
| Approval Criteria 1 - Diagnosis of AIDS (| Approval Criteria 1 - Diagnosis of AIDS (acquired immunodeficiency syndrome)-related Kaposi Sarcoma | | |
| | AND | | |
| 2 - Patient is currently being treated with antiretroviral therapy (ART) | | | |
| AND | | | |
| 3 - Not used as first line therapy | | | |
| AND | | | |
| 4 - History of failure, intolerance, or contraindication to Brand Gleevec. | | | |
| Product Name: Brand (| Gleevec, generic imatinib | | |

Product Name: Brand Gleevec, generic imatinib

| Diagnosis | Steroid-Refractory Chronic Graft-Versus-Host Disease (GVHD) | |
|---|---|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| Approval Criteria 1 - Diagnosis of chronic graft-versus-host disease AND | | |
| 2 - Patient is currently being treated with systemic corticosteroids | | |
| | AND | |
| 3 - Patient had no response to first-line therapy options | | |
| AND | | |
| 4 - If the request is for generic imatinib, there is a reason or special circumstance the patient cannot use brand Gleevec | | |

| Product Name: Brand Gleevec, generic imatinib | |
|---|--|
| Diagnosis | Myeloid/Lymphoid Neoplasms with Eosinophilia |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of lymphoid, myeloid, or mixed lineage neoplasms with eosinophilia

AND

- **2** One of the following:
 - FIP1L1-PDGFRA rearrangement
 - PDGFRB rearrangement
 - ABL1 rearrangement

3 - If the request is for generic imatinib, there is a reason or special circumstance the patient cannot use brand Gleevec

| Product Name: Brand Gleevec, generic imatinib | |
|---|-----------------------------|
| Diagnosis | All Indications except NCCN |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Gleevec therapy

| Product Name: Brand Gleevec, generic imatinib | |
|---|--------------------------|
| Diagnosis | NCCN Recommended Regimen |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Use supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Brand Gleevec, generic imatinib | |
|---|--------------------------|
| Diagnosis | NCCN Recommended Regimen |

| Approval Length | 12 month(s) |
|-----------------|---------------------|
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Gleevec therapy

| Date | Notes |
|-----------|-------------|
| 7/21/2021 | Updated Gpi |

Global Quantity Limits

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99460 | Global Quantity Limits |
|-----------|--|
| Formulary | Medicaid - Arizona (AZM, AZMREF, AZMDDD) |

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Quantity Limit, Prescription Limit | |
|--|---------------------------------|
| Diagnosis | Quantity limit review (General) |
| Approval Length | 12 month(s) |
| Guideline Type | Administrative |
| | |
| Approval Criteria | |
| 1 - ONE of the following: | |
| 1.1 The requested drug must be used for an FDA-approved indication | |

OR

1.2 The use of this drug is supported by information from ONE of the following appropriate compendia of current literature:

- American Hospital Formulary Service Drug Information
- National Comprehensive Cancer Network Drugs and Biologics Compendium
- Thomson Micromedex DrugDex
- Clinical pharmacology
- United States Pharmacopoeia-National Formulary (USP-NF)

AND

2 - The drug is being prescribed within the manufacturer's published dosing guidelines or falls within dosing guidelines found in ONE of the following compendia of current literature:

- American Hospital Formulary Service Drug Information
- National Comprehensive Cancer Network Drugs and Biologics Compendium
- Thomson Micromedex DrugDex
- Clinical pharmacology
- United States Pharmacopoeia-National Formulary (USP-NF)

AND

3 - The requested dosage cannot be achieved using the plan accepted quantity limit of a different dose or formulation.

AND

4 - The drug is being prescribed for a medically accepted indication that is recognized as a covered benefit by the applicable health plans' program.

| Product Name: Quantity Limit, Prescription Limit | |
|--|--|
| Diagnosis | Quantity limit review for the treatment of gender dysphoria* |
| Approval Length | 12 month(s) |
| Guideline Type | Administrative |

Approval Criteria

1 - The use of this drug is supported by information from ONE of the following appropriate compendia of current literature:

- American Hospital Formulary Service Drug Information
- National Comprehensive Cancer Network Drugs and Biologics Compendium
- Thomson Micromedex DrugDex
- Clinical pharmacology
- United States Pharmacopoeia-National Formulary (USP-NF)

AND

2 - The drug is being prescribed for an indication that is recognized as a covered benefit by the applicable health plans' program.

| * If the above criteria are not met, then refer for clinical review by an a ppropriate trained professional (physician or pharmacist) based on the |
|--|
| applicable regulatory requirement. |

| Product Name: Quantity Limit, Prescription Limit | |
|--|--|
| Diagnosis | Monthly prescription limit review for migraine therapy, benzodiazepines, or muscle relaxants |
| Approval Length | 1 month(s) |
| Guideline Type | Administrative |

Approval Criteria

1 - Medical necessity rationale provided for why the member requires 5 or more fills of the same drug or drug class within a month.

| | *If deemed medically necessary, longer authorization duration is perm itted |
|--|---|
|--|---|

| Product Name: Quantity Limit, Prescription Limit | |
|--|--|
| Diagnosis | Topical products exceeding the allowable package size per fill OR the allowable quantity per month |
| Approval Length | 12 month(s) |
| Guideline Type | Administrative |

Approval Criteria

1 - The physician attests that a larger quantity is needed for treatment of a larger surface area.

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk copy C&S Arizona standard to Medicaid Arizona |

GLP-1 Agonists - AZM

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-114508 GLP-1 Agonists - AZM

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Preferred Drugs: Bydureon, Byetta, Trulicity, Victoza | |
|---|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting both of the following:

1.1 Diagnosis of type 2 diabetes mellitus

1.2 History of failure to metformin at a minimum dose of 1500 milligrams (mg) daily for 90 days, or contraindication or intolerance to metformin (verified via paid pharmacy claims or submission of medical records)

| Product Name: Non-Preferred Drugs: Adlyxin, Bydureon BCise, Mounjaro, Ozempic | |
|---|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting both of the following:

1.1 Diagnosis of type 2 diabetes mellitus

AND

1.2 History of failure to metformin at a minimum dose of 1500 milligrams (mg) daily for 90 days, or contraindication or intolerance to metformin (verified via paid pharmacy claims or submission of medical records)

AND

2 - History of a 90 day trial per member's pharmacy claims resulting in a therapeutic failure, contraindication, or intolerance to ALL of the following:

- Byetta
- Victoza
- Trulicity

| Product Name: Non-Preferred: Rybelsus | |
|---------------------------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting both of the following:

1.1 Diagnosis of type 2 diabetes mellitus

AND

1.2 History of failure to metformin at a minimum dose of 1500 milligrams (mg) daily for 90 days, or contraindication or intolerance to metformin (verified via paid pharmacy claims or submission of medical records)

AND

2 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting ONE of the following:

2.1 History of a 90 day trial per member's pharmacy claims resulting in a therapeutic failure, contraindication, or intolerance to ALL of the following:

- Byetta
- Victoza
- Trulicity

OR

2.2 BOTH of the following:

2.2.1 The patient is unable to self-inject due to ONE of the following:

- Physical impairment
- Visual impairment
- Lipohypertrophy
- Documented needle-phobia to the degree that the patient has previously refused any injectable therapy or medical procedure (refer to DSM-5 for specific phobia diagnostic criteria)

AND

2.2.2 History of failure, intolerance, or contraindication to ALL of the following:

- •
- Farxiga Jardiance •
- Invokana •
- Invokamet
- •
- Synjardy Xigduo XR •

| Date | Notes |
|-----------|-----------------------------|
| 9/26/2022 | Added Mounjaro as NP target |

Glycopyrrolate Products

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 7/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Brand Cuvposa oral solution, Dartisla ODT, Brand Robinul, Brand Robinul Forte | |
|---|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting requested drug is being used for a Food and Drug Administration (FDA)-approved indication

2 - Trial and failure or intolerance to generic glycopyrrolate tablets or oral solution (verified via pharmacy paid claims or submission of medical records/chart notes)

| Product Name: Glycopyrrolate injection 0.6mg/3ml | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting requested drug is being used for a Food and Drug Administration (FDA)-approved indication

AND

2 - Trial and failure or intolerance to preferred glycopyrrolate injection strengths (e.g., 0.2 mg/ml, 0.4mg/2ml, 1 mg/5ml, 4mg/20ml) (verified via pharmacy paid claims or submission of medical records/chart notes)

| Date | Notes |
|-----------|---|
| 6/24/2022 | Added NP glycopyrrolate inj as target. Changed guideline name to Gl ycopyrrolate Products. |

Gonadotropin-Releasing Hormone Agonists

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99798 | Gonadotropin-Releasing | Hormone Agonists |
|----------|------------------------|------------------|
| | | |

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: leuprolide acetate, Lupron Depot Ped, Triptodur, Fensolvi | |
|---|----------------------------------|
| Diagnosis | Central Precocious Puberty (CPP) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of central precocious puberty (idiopathic or neurogenic)

2 - Onset of secondary sexual characteristics in one of the following:

2.1 Females less than or equal to 8 years of age

OR

2.2 Males less than or equal to 9 years of age

AND

3 - Confirmation of diagnosis as defined by one of the following:

3.1 Pubertal basal level of luteinizing hormone (based on laboratory reference ranges)

OR

3.2 A pubertal luteinizing hormone response to a gonadotropin releasing hormone (GnRH) stimulation test

OR

3.3 Bone age advanced one year beyond the chronological age

AND

4 - If the request is for Triptodur or Fensolvi, history of failure, contraindication, or intolerance to Lupron-Depot Ped

| Product Name: leuprolide acetate, Lupron Depot Ped, Triptodur, Fensolvi | |
|---|----------------------------------|
| Diagnosis | Central Precocious Puberty (CPP) |
| Approval Length | 12 month(s) |

| Therapy Stage | Reauthorization | |
|--|---------------------|--|
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Patient is currently receiving therapy for central precocious puberty | | |
| | | |
| | AND | |
| 2 - Documentation of positive clinical response to therapy | | |
| AND | | |
| 3 - Patient is ONE of the following (younger than the appropriate time point for the onset of puberty): | | |
| 3.1 Female younger than 11 years of age | | |
| | | |
| OR | | |
| 3.2 Male younger thar | 12 years of age | |

| Product Name: Lupaneta Pack, Lupron Depot 3.75 mg and 3-month 11.25 mg | | |
|--|-----------------------|--|
| Diagnosis | Endometriosis | |
| Approval Length | 6 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Diagnosis of endometriosis or endometriosis is suspected

AND

2 - One of the following:

2.1 History of failure, contraindication, or intolerance to both of the following:

2.1.1 Oral contraceptives or depot medroxyprogesterone (e.g., Depo- Provera)

AND

2.1.2 Non-steroidal anti-inflammatory drugs (NSAIDs)

OR

2.2 Patient has had surgical ablation to prevent recurrence

AND

3 - If the request is for Lupaneta Pack, history of failure, contraindication, or intolerance to Lupron Depot

| Product Name: Lupaneta Pack, Lupron Depot 3.75 mg and 3-month 11.25 mg | |
|--|---------------------|
| Diagnosis | Endometriosis |
| Approval Length | 6 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of endometriosis or endometriosis is suspected

AND

2 - Recurrence of symptoms following an initial course of therapy

3 - Concurrently to be used with add-back therapy (e.g., progestin, estrogen, or bone sparing agents)

| Product Name: Lupron Depot 3.75 mg and 3-month 11.25 mg | |
|---|--------------------------------|
| Diagnosis | Uterine Leiomyomata (Fibroids) |
| Approval Length | 3 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - One of the following:

1.1 All of the following:

1.1.1 For the treatment of uterine leiomyomata-related anemia

AND

1.1.2 Patient did not respond to iron therapy of 1 month duration

AND

1.1.3 For use prior to surgery

OR

1.2 For use prior to surgery to reduce the size of fibroids to facilitate a surgical procedure (e.g., myomectomy, hysterectomy)

| Product Name: Lupron Depot, Lupron Depot-Ped, Lupaneta Pack, leuprolide acetate, Triptodur, Fensolvi | |
|---|---------------------------------|
| Diagnosis | Gender dysphoria in adolescents |

| Approval Length | 12 month(s) |
|-----------------|-----------------------|
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

1 - Diagnosis of gender dysphoria, according to the current Diagnostic and Statistical Manual of Mental Disorders (i.e., DSM-5) criteria, by a mental health professional with expertise in child and adolescent psychiatry

AND

2 - Medication is prescribed by or in consultation with an endocrinologist or a medical provider experienced in gender dysphoria hormone therapy

AND

3 - Patient has experienced puberty development to at least Tanner stage 2

AND

4 - One of the following laboratory tests, based upon the laboratory reference range, confirming:

- Pubertal levels of estradiol in females
- Pubertal levels of testosterone in males
- Pubertal basal level of luteinizing hormone (based on laboratory reference ranges)
- A pubertal luteinizing hormone response to a gonadotropin-releasing hormone (GnRH) stimulation test

AND

5 - A letter from the prescriber and/or formal documentation stating all of the following:

5.1 Patient has experienced pubertal changes that have resulted in an increase of their gender dysphoria that has significantly impaired psychological or social functioning

5.2 Coexisting psychiatric and medical comorbidities or social problems that may interfere with the diagnostic procedures or treatment have been addressed or removed

AND

5.3 Both of the following:

5.3.1 Current enrollment, attendance, and active participation in psychological and social support treatment program

AND

5.3.2 Patient will continue enrollment, attendance and active participation in psychological and social support throughout the course of treatment

AND

5.4 Patient demonstrates knowledge and understanding of the expected outcomes of treatment and related transgender therapies

AND

6 - If the request is for Lupaneta Pack, leuprolide acetate, Triptodur, Fensolvi, history of failure, contraindication, or intolerance to Lupron Depot

| Product Name: Lupron Depot, Lupron Depot-Ped, Lupaneta Pack, leuprolide acetate, Triptodur, Fensolvi | |
|--|---------------------------------|
| Diagnosis | Gender dysphoria in adolescents |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - One of the following:

- Documentation (within the last 6 months) of appropriate luteinizing hormone (LH) suppression
- Change in dosing

AND

2 - Documented diagnosis of gender dysphoria, according to the current Diagnostic and Statistical Manual of Mental Disorders (i.e., DSM-5) criteria, by a mental health professional with expertise in child and adolescent psychiatry

AND

3 - Medication is prescribed by or in consultation with an endocrinologist or a medical provider experienced in gender dysphoria hormone therapy

AND

4 - A letter from the prescriber and/or formal documentation stating all of the following:

4.1 Patient continues to meet their individual goals of therapy for gender dysphoria

AND

4.2 Patient continues to have a strong affinity for the desired (opposite of natal) gender

AND

4.3 Discontinuation of treatment and subsequent pubertal development would interfere with or impair psychological functioning and well-being

AND

4.4 Coexisting psychiatric and medical comorbidities or social problems that may interfere with treatment continue to be addressed or removed

AND

4.5 Both of the following:

4.5.1 Current enrollment, attendance, and active participation in psychological and social support treatment program

AND

4.5.2 Patient will continue enrollment, attendance and active participation in psychological and social support throughout the course of treatment

AND

4.6 Patient demonstrates knowledge and understanding of the expected outcomes of treatment and related transgender therapies

| Product Name: Lupron Depot, Lupron Depot-Ped, Lupaneta Pack, leuprolide acetate, Triptodur, Fensolvi | |
|---|---|
| Diagnosis | Adjunct for Gender-Affirming Hormonal Therapy for Transgender Adults |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of gender dysphoria, according to the current Diagnostic and Statistical Manual of Mental Disorders (i.e., DSM-5) criteria, by a mental health professional

AND

2 - Medication is prescribed by or in consultation with an endocrinologist or a medical provider experienced in transgender hormone therapy
 AND
 3 - Gonads (i.e., testes, ovaries) have not been removed and are functional (e.g., hormone

AND

4 - Patient is currently receiving hormonal therapy (e.g., testosterone, estrogens, progesterones) to achieve the desired (e.g., non-natal) gender

AND

5 - Inability of cross sex hormone therapy to inhibit natal secondary sex characteristics, luteinizing hormone (LH), or gonadotropins (e.g., menses, testosterone)

AND

6 - A letter from the prescriber and/or formal documentation stating all of the following:

6.1 Transgender patient has identified goals of gender-affirming hormone therapy

AND

6.2 Coexisting psychiatric and medical comorbidities or social problems that may interfere with the diagnostic procedures or treatment have been addressed or removed

AND

6.3 Both of the following:

producing)

6.3.1 Current enrollment, attendance, and active participation in psychological and social support treatment program

6.3.2 Patient will continue enrollment, attendance and active participation in psychological and social support throughout the course of treatment

AND

6.4 Patient demonstrates knowledge and understanding of the expected outcomes of treatment and related transgender therapies

AND

7 - If the request is for Lupaneta Pack, leuprolide acetate, Triptodur, Fensolvi, history of failure, contraindication, or intolerance to Lupron Depot

| Product Name: Lupron Depot, Lupron Depot-Ped, Lupaneta Pack, leuprolide acetate, Triptodur, Fensolvi | |
|---|---|
| Diagnosis | Adjunct for Gender-Affirming Hormonal Therapy for Transgender Adults |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - One of the following:

- Documentation (within the last 6 months) of appropriate luteinizing hormone (LH) suppression
- Change in dosing

AND

2 - Documented diagnosis of gender dysphoria, according to the current Diagnostic and Statistical Manual of Mental Disorders (i.e., DSM-5) criteria, by a mental health professional

3 - Medication is prescribed by or in consultation with an endocrinologist or a medical provider experienced in transgender hormone therapy

AND

4 - Gonads (i.e., testes, ovaries) are intact

AND

5 - Patient is currently receiving hormonal therapy (e.g., testosterone, estrogens, progesterones) to achieve the desired (e.g., non-natal) gender

AND

6 - Inability of cross sex hormone therapy to inhibit natal secondary sex characteristics, luteinizing hormone (LH), or gonadotropins (e.g., menses, testosterone)

AND

7 - A letter from the prescriber and/or formal documentation stating all of the following:

7.1 Transgender patient continues to meet goals of gender-affirming hormone therapy

AND

7.2 Coexisting psychiatric and medical comorbidities or social problems that may interfere with the diagnostic procedures or treatment continue to be addressed or removed

AND

7.3 Both of the following:

7.3.1 Current enrollment, attendance, and active participation in psychological and social support treatment program

AND

7.3.2 Patient will continue enrollment, attendance and active participation in psychological and social support throughout the course of treatment

AND

7.4 Patient demonstrates knowledge and understanding of the expected outcomes of treatment and related transgender therapies

| Product Name: Lupron Depot, Lupron Depot Ped, Lupaneta Pack, Triptodur, generic leuprolide acetate solution for injection, Fensolvi | |
|---|------------------------|
| Diagnosis | Fertility Preservation |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - For use in pre-menopausal women

AND

2 - Patient is receiving a cytotoxic agent that is associated with causing primary ovarian insufficiency (premature ovarian failure) [e.g., Cytoxan (cyclophosphamide), procarbazine, vinblastine, cisplatin]

AND

3 - If the request is for Lupaneta Pack, leuprolide acetate, Triptodur, Fensolvi, history of failure, contraindication, or intolerance to Lupron Depot.

| Product Name: Lupron Depot, Lupron Depot Ped, Lupaneta Pack, Triptodur, generic leuprolide acetate solution for injection, Fensolvi | |
|---|------------------------|
| Diagnosis | Fertility Preservation |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Patient is currently receiving gonadotropin-releasing hormone (GnRH) analog therapy for the purpose of fertility preservation

AND

2 - Patient continues to receive a cytotoxic agent that is associated with causing primary ovarian insufficiency (premature ovarian failure) [e.g., Cytoxan (cyclophosphamide), procarbazine, vinblastine, cisplatin]

| Product Name: Lupron Depot 7.5 mg, 22.5 mg, 30 mg and 45 mg, generic leuprolide acetate solution for injection | |
|--|--|
| Diagnosis | Advanced or Metastatic Prostate Cancer |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of advanced or metastatic prostate cancer

2. Revision History

| Date | Notes |
|----------|-------------------|
| 7/1/2021 | Updated Guideline |

Gralise, Horizant - Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Gralise | | |
|---|--|--|
| 12 month(s) | | |
| Prior Authorization | | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of postherpetic neuralgia (PHN) | | |
| | | |

| Product Name: Horizant | |
|------------------------|-------------|
| Approval Length | 12 month(s) |

| Guideline Type | Prior Authorization | |
|---|---------------------|--|
| | | |
| Approval Criteria | | |
| 1 - One of the following: | | |
| 1.1 Diagnosis of postherpetic neuralgia (PHN) | | |
| | OR | |
| | | |
| 1.2 Diagnosis of restless legs syndrome | | |

2. Revision History

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk copy C&S Arizona standard to Medicaid Arizona |

Growth Hormone, Growth Stimulating Agents - AZM Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-112076 | Growth Hormone, | e, Growth Stimulating Agents - AZM | Ν |
|-----------|-----------------|------------------------------------|---|
|-----------|-----------------|------------------------------------|---|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 8/22/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: All products | | | |
|---|--|--|--|
| Diagnosis | Idiopathic Short Stature (ISS) | | |
| Approval Length | N/A - Requests for non-approvable diagnoses should not be approved | | |
| Guideline Type Prior Authorization | | | |
| | | | |
| Approval Criteria | | | |
| 1 - Requests for coverage for diagnosis of Idiopathic Short Stature (ISS) are not authorized and will not be approved | | | |
| Notes | Approval Length: N/A - Requests for Idiopathic Short Stature (ISS) sh ould not be approved. Deny as a benefit exclusion. | | |

| Product Name: Non Preferred* Humatrope, Nutropin AQ NuSpin, Omnitrope, Saizen, Saizen Click Easy, Zomacton, Zorbitive, Serostim* | | | | |
|--|---|--|--|--|
| Approval Length | oval Length N/A - Requests for Non-Preferred Drugs should not be approved | | | |
| Guideline Type Prior Authorization | | | | |
| | | | | |

1 - Patient has tried and failed the 2 preferred products listed below.

- Brand Genotropin/Genotropin Miniquick Brand Norditropin Flexpro •
- •

| Notes | *Requests for coverage of Non-Preferred drugs are not authorized an d will not be approved. Patient must use one of the preferred products listed above. All Non preferred products will be denied for appeals pr ocess. |
|-------|--|
|-------|--|

| Product Name: Preferred (Brand Genotropin/Genotropin Miniquick, Brand Norditropin Flexpro) | | | |
|--|---|--|--|
| Diagnosis | Pediatric Growth Hormone Deficiency (GHD) | | |
| Approval Length | 12 month(s) | | |
| Therapy Stage Initial Authorization | | | |
| Guideline Type Prior Authorization | | | |

Approval Criteria

1 - ONE of the following:

- **1.1** ONE of the following:
- **1.1.1** All of the following:
 - Infant is less than 4 months of age •
 - Infant has growth deficiency
 - Prescribed by an endocrinologist •

1.1.2 BOTH of the following:

- History of neonatal hypoglycemia associated with pituitary disease
- Prescribed by an endocrinologist

OR

1.1.3 BOTH of the following:

- Diagnosis of panhypopituitarism
- Prescribed by an endocrinologist

OR

1.2 ALL of the following:

1.2.1 Diagnosis of pediatric growth hormone (GH) deficiency as confirmed by ONE of the following:

1.2.1.1 Projected height (as determined by extrapolating pre-treatment growth trajectory along current channel to 18-20 year mark) is greater than 2.0 standard deviations [SD] below midparental height utilizing age and gender growth charts related to height

OR

1.2.1.2 Height is greater than 2.25 SD below population mean (below the 1.2 percentile for age and gender) utilizing age and gender growth charts related to height

OR

1.2.1.3 Growth velocity is greater than 2 SD below mean for age and gender

OR

1.2.1.4 Delayed skeletal maturation of greater than 2 SD below mean for age and gender (e.g., delayed greater than 2 years compared with chronological age)

1.2.2 ONE of the following:

1.2.2.1 BOTH of the following:

- Patient is male
- Bone age less than 16 years

OR

1.2.2.2 BOTH of the following:

- Patient is female
- Bone age less than 14 years

AND

1.2.3 Submission of medical records (e.g., chart notes, laboratory values) documenting ONE of the following:

1.2.3.1 BOTH of the following:

1.2.3.1.1 Patient has undergone TWO of the following provocative GH stimulation tests:

- Arginine
- Clonidine
- Glucagon
- Insulin
- Levodopa
- Growth hormone releasing hormone

AND

1.2.3.1.2 BOTH GH response values are less than 10 micrograms per liter

OR

1.2.3.2 BOTH of the following:

1.2.3.2.1 Patient is less than 1 year of age

AND

1.2.3.2.2 ONE of the following is below the age and gender adjusted normal range as provided by the physician's lab:

- Insulin-like Growth Factor 1 (IGF-1/Somatomedin-C)
- Insulin Growth Factor Binding Protein-3 (IGFBP-3)

AND

1.2.4 ONE of the following:

1.2.4.1 Request does not exceed a maximum supply limit of 0.3 milligrams per kilogram per week

OR

1.2.4.2 BOTH of the following:

- Tanner Stage 3 or greater
- Request does not exceed a maximum supply limit of 0.7 milligrams per kilogram per week

AND

1.2.5 Prescribed by an endocrinologist

| Notes | *Includes children who have undergone brain radiation. If patient is a Transition Phase Adolescent or Adult who had childhood onset GH de |
|-------|---|
| | ficiency, utilize criteria for Transition Phase Adolescent or Adult GH D eficiency. |

| Product Name: Preferred (Brand Genotropin/Genotropin Miniquick, Brand Norditropin Flexpro) | |
|--|--|
| Diagnosis Pediatric Growth Hormone Deficiency (GHD) | |

| Approval Length | 12 month(s) | | |
|--|--|--|--|
| Therapy Stage | Reauthorization | | |
| Guideline Type | Prior Authorization | | |
| Approval Criteria | | | |
| BOTH of the followin | of at least 2 centimeters per year over the previous year documented by g:** | | |
| | ght and date obtained ht and date obtained | | |
| | AND | | |
| 2 - BOTH of the follo | wing:** | | |
| | ult height not attained on of expected adult height goal (e.g. genetic potential) | | |
| | AND | | |
| 3 - Calculated height | t (growth) velocity over the past 12 months | | |
| | AND | | |
| 4 - ONE of the follow | <i>v</i> ing: | | |
| 4.1 BOTH of the fol | lowing: | | |
| Patient is maBone age les | le s than 16 years | | |
| | OR | | |
| 4.2 BOTH of the fol | lowing: | | |
| | | | |

• Patient is female

| • | Bone age | less than | 14 years |
|---|----------|-----------|----------|
|---|----------|-----------|----------|

5 - ONE of the following:

5.1 Request does not exceed a maximum supply limit of 0.3 milligrams per kilogram per week

OR

5.2 BOTH of the following:

- Tanner Stage 3 or greater
- Request does not exceed a maximum supply limit of 0.7 milligrams per kilogram per week

AND

| 6 - Prescribed by an endocrinologist | |
|--------------------------------------|---|
| Notes | *Includes children who have undergone brain radiation. If patient is a Transition Phase Adolescent or Adult who had childhood onset GH de ficiency, utilize criteria for Transition Phase Adolescent or Adult GH D eficiency. ** Documentation of previous height, current height and goa I expected adult height will be required for renewal. |

| Product Name: Preferred (Brand Genotropin/Genotropin Miniquick, Brand Norditropin Flexpro) | | |
|--|-----------------------|--|
| Diagnosis | Prader-Willi Syndrome | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Diagnosis of Prader-Willi Syndrome

2 - Prescribed by an endocrinologist

Product Name: Preferred (Brand Genotropin/Genotropin Miniquick, Brand Norditropin
Flexpro)DiagnosisPrader-Willi SyndromeApproval Length12 month(s)Therapy StageReauthorization

Prior Authorization

Approval Criteria

Guideline Type

1 - ONE of the following criteria:

1.1 BOTH of the following:

1.1.1 Evidence of positive response to therapy (e.g., increase in total lean body mass, decrease in fat mass)

AND

1.1.2 Prescribed by an endocrinologist

OR

1.2 ALL of the following:

1.2.1 Height increase of at least 2 centimeters per year over the previous year of treatment as documented by BOTH of the following:

- Previous height and date obtained
- Current height and date obtained

AND

1.2.2 BOTH of the following:

- Expected adult height not attained
- Documentation of expected adult height goal

AND

1.2.3 Prescribed by an endocrinologist

Product Name: Preferred (Brand Genotropin/Genotropin Miniquick, Brand Norditropin Flexpro)

| Diagnosis | Growth Failure in Children Small for Gestational Age (SGA) | |
|-----------------|--|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Diagnosis of small for gestational age (SGA) based on demonstration of catch up growth failure in the first 24 months of life using a 0-36 month growth chart as confirmed by documentation that ONE of the following is below the third percentile for gestational age (more than 2 standard deviations [SD] below population mean):

- Birth weight
- Birth length

AND

2 - Documentation that height remains less than or equal to the third percentile (more than 2 SD below population mean)

AND

3 - Prescribed by an endocrinologist

| Product Name: Preferred (Brand Genotropin/Genotropin Miniquick, Brand Norditropin Flexpro) | | |
|--|--|--|
| Diagnosis | Growth Failure in Children Small for Gestational Age (SGA) | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |

1 - Height increase of at least 2 centimeters per year over the previous year documented by BOTH of the following:*

- Previous height and date obtained
- Current height and date obtained

AND

2 - Documentation of BOTH of the following:*

- Expected adult height not attained
- Expected adult height goal

AND

3 - Prescribed by an endocrinologist

Notes *Documentation of previous height, current height and goal expected adult height will be required for renewal.

| Product Name: Preferred (Brand Genotropin/Genotropin Miniquick, Brand Norditropin Flexpro) | | |
|--|------------------------------------|--|
| Diagnosis | Turner Syndrome or Noonan Syndrome | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |

Approval Criteria

1 - Diagnosis of pediatric growth failure associated with ONE of the following: **1.1** BOTH of the following: **1.1.1** Turner Syndrome (Gonadal Dysgenesis) AND **1.1.2** BOTH of the following: Patient is female • Bone age less than 14 years • OR **1.2** BOTH of the following: 1.2.1 Noonan Syndrome AND 1.2.2 ONE of the following: **1.2.2.1** BOTH of the following: Patient is male Bone age less than 16 years • OR 1.2.2.2 BOTH of the following: Patient is female • Bone age less than 14 years • AND 2 - Height is below the fifth percentile on growth charts for age and gender

3 - Prescribed by an endocrinologist

| Product Name: Preferred (Brand Genotropin/Genotropin Miniquick, Brand Norditropin Flexpro) | | |
|--|------------------------------------|--|
| Diagnosis | Turner Syndrome or Noonan Syndrome | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Height increase of at least 2 centimeters per year over the previous year documented by BOTH of the following:*

- Previous height and date obtained
- Current height and date obtained

AND

- 2 Documentation of BOTH of the following:*
 - Expected adult height not attained
 - Expected adult height goal

AND

3 - Prescribed by an endocrinologist

| Notes | *Documentation of previous height, current height and goal expected |
|-------|---|
| | adult height will be required for renewal. |

Product Name: Preferred (Brand Genotropin/Genotropin Miniquick, Brand Norditropin Flexpro)

| Approval Length12 month(s)Therapy StageInitial AuthorizationGuideline TypePrior Authorization | | |
|--|--------------------------------|---|
| Therapy Stage Initial Authorization Guideline Type Prior Authorization Approval Criteria I - Diagnosis of pediatric growth failure with short-stature homeobox (SHOX) gene deficiency as confirmed by genetic testing AND 2 - ONE of the following: 2.1 BOTH of the following: • Patient is male • Bone age less than 16 years OR 2.2 BOTH of the following: • Patient is female • Bone age less than 14 years | Diagnosis | Short-Stature Homeobox (SHOX) Gene Deficiency |
| Guideline Type Prior Authorization Approval Criteria 1 - Diagnosis of pediatric growth failure with short-stature homeobox (SHOX) gene deficiency as confirmed by genetic testing AND 2 - ONE of the following: 2.1 BOTH of the following: • Patient is male • Bone age less than 16 years OR 2.2 BOTH of the following: • Patient is female • Bone age less than 14 years | Approval Length | 12 month(s) |
| Approval Criteria 1 - Diagnosis of pediatric growth failure with short-stature homeobox (SHOX) gene deficiency as confirmed by genetic testing AND 2 - ONE of the following: 2.1 BOTH of the following: 9 Patient is male 9 Bone age less than 16 years OR 2.2 BOTH of the following: 9 Patient is female 9 Bone age less than 14 years AND | Therapy Stage | Initial Authorization |
| 1 - Diagnosis of pediatric growth failure with short-stature homeobox (SHOX) gene deficiency as confirmed by genetic testing AND 2 - ONE of the following: Patient is male Bone age less than 16 years 2.2 BOTH of the following: Patient is female Bone age less than 14 years | Guideline Type | Prior Authorization |
| 1 - Diagnosis of pediatric growth failure with short-stature homeobox (SHOX) gene deficiency as confirmed by genetic testing AND 2 - ONE of the following: Patient is male Bone age less than 16 years 2.2 BOTH of the following: Patient is female Bone age less than 14 years | | |
| AND 2 - ONE of the following: 2 - ONE of the following: 3 - Patient is male - Bone age less than 16 years OR 2.2 BOTH of the following: - Patient is female - Bone age less than 14 years AND | Approval Criteria | |
| 2 - ONE of the following: 2.1 BOTH of the following: Patient is male Bone age less than 16 years OR 2.2 BOTH of the following: Patient is female Bone age less than 14 years AND | | |
| 2.1 BOTH of the following: Patient is male Bone age less than 16 years OR 2.2 BOTH of the following: Patient is female Bone age less than 14 years | | AND |
| Patient is male Bone age less than 16 years OR 2.2 BOTH of the following: Patient is female Bone age less than 14 years AND | 2 - ONE of the following | g: |
| Bone age less than 16 years OR 2.2 BOTH of the following: Patient is female Bone age less than 14 years AND | 2.1 BOTH of the follow | ving: |
| 2.2 BOTH of the following: Patient is female Bone age less than 14 years | | |
| Patient is female Bone age less than 14 years AND | | OR |
| Bone age less than 14 years AND | 2.2 BOTH of the follow | ving: |
| | | |
| 3 - Prescribed by an endocrinologist | | AND |
| | 3 - Prescribed by an en | docrinologist |

| Product Name: Preferred (Brand Genotropin/Genotropin Miniquick, Brand Norditropin Flexpro) | | |
|--|---|--|
| Diagnosis | Short-Stature Homeobox (SHOX) Gene Deficiency | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |

1 - Height increase of at least 2 centimeters per year over the previous year documented by BOTH of the following:*

- Previous height and date obtained
- Current height and date obtained

AND

2 - Documentation of BOTH of the following:*

- Expected adult height not attained
- Expected adult height goal

AND

3 - Prescribed by an endocrinologist

| *Documentation of previous height, current height and goal expected |
|---|
| adult height will be required for renewal. |

| Product Name: Preferred (Brand Genotropin/Genotropin Miniquick, Brand Norditropin Flexpro) | | |
|--|--|--|
| Diagnosis | Growth Failure associated with Chronic Renal Insufficiency | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Diagnosis of pediatric growth failure associated with chronic renal insufficiency

AND

2 - ONE of the following:

| 2.1 BOTH of the following: | |
|--|---------|
| Patient is maleBone age less than 16 ye | ears |
| | OR |
| 2.2 BOTH of the following: | |
| Patient is femaleBone age less than 14 ye | ears |
| | AND |
| 3 - Prescribed by ONE of the foll | lowing: |

- Endocrinologist Nephrologist •
- •

| Product Name: Preferred (Brand Genotropin/Genotropin Miniquick, Brand Norditropin Flexpro) | |
|--|--|
| Diagnosis | Growth Failure associated with Chronic Renal Insufficiency |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Height increase of at least 2 centimeters per year over the previous year documented by BOTH of the following:*

- Previous height and date obtainedCurrent height and date obtained

| 2 - | Documentation | of BOTH of | f the followina:* |
|-----|----------------|------------|-------------------|
| _ | Doodinontation | 0.001110 | r allo ronoming. |

- Expected adult height not attained
- Expected adult height goal

3 - Prescribed by ONE of the following:

- Endocrinologist
- Nephrologist

| Notes | *Documentation of previous height, current height and goal expected |
|-------|---|
| | adult height will be required for renewal. |

| Product Name: Preferred (Brand Genotropin/Genotropin Miniquick, Brand Norditropin Flexpro) | |
|--|---------------------------------|
| Diagnosis | Adult Growth Hormone Deficiency |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of adult growth hormone deficiency (GHD) as a result of ONE of the following:

1.1 Clinical records supporting a diagnosis of childhood-onset GHD

OR

1.2 BOTH of the following:

1.2.1 Adult-onset GHD

AND

1.2.2 Clinical records documenting that hormone deficiency is a result of hypothalamicpituitary disease from organic or known causes (e.g., damage from surgery, cranial irradiation, head trauma, or subarachnoid hemorrhage)

AND

2 - Submission of medical records (e.g., chart notes, laboratory values) documenting ONE of the following:

2.1 BOTH of the following:

2.1.1 Patient has undergone ONE of the following GH (growth hormone) stimulation tests to confirm adult GH deficiency:

- Insulin tolerance test (ITT)
- ARG (Arginine) and GHRH (growth hormone releasing hormone)
- Glucagon
- ARG

AND

2.1.2 ONE of the following peak GH values:

2.1.2.1 ITT less than or equal to 5 micrograms per liter

OR

2.1.2.2 GHRH and ARG of ONE of the following:

- Less than or equal to 11 micrograms per liter if body mass index [BMI] is less than 25 kilograms per meter squared
- Less than or equal to 8 micrograms per liter if BMI is greater than or equal to 25 and less than 30 kilograms per meter squared
- Less than or equal to 4 micrograms per liter if BMI is greater than or equal to 30 kilograms per meter squared

OR

2.1.2.3 Glucagon less than or equal to 3 micrograms per liter

OR

2.1.2.4 ARG less than or equal to 0.4 micrograms per liter

OR

2.2 BOTH of the following:

2.2.1 Submission of medical records (e.g., chart notes, laboratory values) documenting deficiency of THREE of the following anterior pituitary hormones:

- Prolactin
- ACTH (adrenocorticotropic hormone)
- TSH (thyroid stimulating hormone)
- FSH/LH (follicle-stimulating hormone/luteinizing hormone)

AND

2.2.2 Insulin-like Growth Factor 1 (IGF-1)/Somatomedin-C level is below the age and gender adjusted normal range as provided by the physician's lab

AND

3 - ONE of the following:

3.1 Diagnosis of panhypopituitarism

OR

3.2 Other diagnosis and not used in combination with BOTH of the following:

- Aromatase inhibitors [e.g., Arimidex (anastrazole), Femara (letrazole)]
- Androgens [e.g., Delatestryl (testosterone enanthate), Depo-Testosterone (testosterone cypionate)]

AND

4 - Request does not exceed a maximum supply limit of 0.3 milligrams per kilogram per week

AND

5 - Prescribed by an endocrinologist

| Product Name: Preferred (Brand Genotropin/Genotropin Miniquick, Brand Norditropin Flexpro) | |
|--|---------------------------------|
| Diagnosis | Adult Growth Hormone Deficiency |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of Insulin-like Growth Factor 1 (IGF-1)/Somatomedin C level within the past 12 months

AND

2 - ONE of the following:

2.1 Diagnosis of panhypopituitarism

OR

2.2 Other diagnosis and not used in combination with BOTH of the following:

- Aromatase inhibitors [e.g., Arimidex (anastrazole), Femara (letrazole)]
- Androgens [e.g., Delatestryl (testosterone enanthate), Depo-Testosterone (testosterone cypionate)]

AND

3 - Request does not exceed a maximum supply limit of 0.3 milligrams per kilogram per week

4 - Prescribed by an endocrinologist

| Product Name: Preferred (Brand Genotropin/Genotropin Miniquick, Brand Norditropin Flexpro) | |
|--|--------------------------------------|
| Diagnosis | Transition Phase Adolescent Patients |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Request does not exceed a maximum supply limit of 0.3 milligrams per kilogram per week

AND

2 - Documentation of ONE of the following:

- Attained expected adult height
- Closed epiphyses on bone radiograph

AND

3 - Submission of medical records (e.g., chart notes, laboratory values) documenting ONE of the following:

3.1 BOTH of the following:

3.1.1 Documentation of high risk of growth hormone (GH) deficiency due to GH deficiency in childhood from ONE of the following:

3.1.1.1 Embryopathic/congenital defects

OR

3.1.1.2 Genetic mutations OR 3.1.1.3 Irreversible structural hypothalamic-pituitary disease OR 3.1.1.4 Panhypopituitarism OR **3.1.1.5** Deficiency of THREE of the following anterior pituitary hormones: ACTH (adrenocorticotropic hormone) ٠ TSH (thyroid stimulating hormone) • Prolactin • FSH/LH (follicle-stimulating hormone/luteinizing hormone) • AND 3.1.2 ONE of the following: 3.1.2.1 Insulin-like Growth Factor 1 (IGF-1)/Somatomedin-C level is below the age and gender adjusted normal range as provided by the physician's lab OR 3.1.2.2 ALL of the following: 3.1.2.2.1 Patient does not have a low IGF-1/Somatomedin C level AND **3.1.2.2.2** Discontinued GH therapy for at least 1 month

3.1.2.2.3 Patient has undergone ONE of the following GH stimulation tests after discontinuation of therapy for at least 1 month:

- Insulin tolerance test (ITT)
- ARG (Arginine) and GHRH (growth hormone releasing hormone)
- ARG
- Glucagon

AND

3.1.2.2.4 ONE of the following peak GH values:

3.1.2.2.4.1 ITT less than or equal to 5 micrograms per liter

OR

3.1.2.2.4.2 GHRH and ARG of ONE of the following:

- Less than or equal to 11 micrograms per liter if body mass index [BMI] is less than 25 kilograms per meter squared
- Less than or equal to 8 micrograms per liter if BMI is greater than or equal to 25 and less than 30 kilograms per meter squared
- Less than or equal to 4 micrograms per liter if BMI is greater than or equal to 30 kilograms per meter squared

OR

3.1.2.2.4.3 Glucagon less than or equal to 3 micrograms per liter

OR

3.1.2.2.4.4 ARG less than or equal to 0.4 micrograms per liter

3.2 ALL of the following:

3.2.1 At low risk of severe GH deficiency (e.g., due to isolated and/or idiopathic GH deficiency)

AND

OR

3.2.2 Discontinued GH therapy for at least 1 month

AND

3.2.3 BOTH of the following:

3.2.3.1 Patient has undergone ONE of the following GH stimulation tests after discontinuation of therapy for at least 1 month:

- ITT
- GHRH and ARG
- ARG
- Glucagon

AND

3.2.3.2 ONE of the following peak GH values:

3.2.3.2.1 ITT less than or equal to 5 micrograms per liter

OR

3.2.3.2.2 GHRH and ARG of ONE of the following:

• Less than or equal to 11 micrograms per liter if body mass index [BMI] is less than 25 kilograms per meter squared

- Less than or equal to 8 micrograms per liter if BMI is greater than or equal to 25 and less than 30 kilograms per meter squared
- Less than or equal to 4 micrograms per liter if BMI is greater than or equal to 30 kilograms per meter squared

OR

3.2.3.2.3 Glucagon less than or equal to 3 micrograms per liter

OR

3.2.3.2.4 ARG less than or equal to 0.4 micrograms per liter

AND

4 - Prescribed by an endocrinologist

| Product Name: Preferred (Brand Genotropin/Genotropin Miniquick, Brand Norditropin Flexpro) | |
|--|--------------------------------------|
| Diagnosis | Transition Phase Adolescent Patients |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive response to therapy (e.g., increase in total lean body mass, exercise capacity or IGF-1 [Insulin-like Growth Factor 1] and IGFBP-3 [Insulin-like growth factor binding protein 3] levels)

AND

2 - Request does not exceed a maximum supply limit of 0.3 milligrams per kilogram per week

3 - Prescribed by an endocrinologist

| Product Name: Serosti | m |
|---|--|
| Diagnosis | Human Immunodeficiency Virus (HIV)-associated wasting syndrome or cachexia |
| Approval Length | 3 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| Approval Criteria 1 - Diagnosis of human cachexia | immunodeficiency virus (HIV)-associated wasting syndrome or |
| | AND |
| 2 - Documentation of C | ONE of the following: |
| 2.1 Unintentional weig | ght loss of greater than 10 percent over the last 12 months |
| | OR |
| 2.2 Unintentional weig | ght loss of greater than 7.5 percent over the last 6 months |
| | OR |
| 2.3 Loss of 5 percent | body cell mass (BCM) within 6 months |
| | OR |
| 2.4 Body mass index | (BMI) less than 20 kilograms per meter squared |
| | |

OR

2.5 ONE of the following:

2.5.1 ALL of the following:

- Patient is male
- BCM less than 35 percent of total body weight
- BMI less than 27 kilograms per meter squared

OR

2.5.2 ALL of the following:

- Patient is female
- BCM less than 23 percent of total body weight
- BMI less than 27 kilograms per meter squared

AND

3 - A nutritional evaluation has been completed since onset of wasting first occurred

AND

4 - Patient has not had weight loss as a result of other underlying treatable conditions (e.g., depression, mycobacterium avium complex, chronic infectious diarrhea, or malignancy with the exception of Kaposi's sarcoma limited to skin or mucous membranes)

AND

5 - Patient's anti-retroviral therapy has been optimized to decrease the viral load

| Product Name: Serostim | |
|------------------------|--|
| Diagnosis | Human Immunodeficiency Virus (HIV)-associated wasting syndrome or cachexia |

| Approval Length | 6 month(s) |
|-----------------|---------------------|
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Evidence of positive response to therapy (i.e., greater than or equal to 2 percent increase in body weight and/or body cell mass [BCM])

AND

2 - ONE of the following targets or goals has not been achieved:

- Weight
- ВСЙ
- Body Mass Index (BMI)

| Product Name: Zorbtive* | | | |
|--|--|--|--|
| Diagnosis | Short Bowel Syndrome | | |
| Guideline Type | Prior Authorization | | |
| | | | |
| Approval Criteria | | | |
| 1 - Diagnosis of Short I | Bowel Syndrome | | |
| | | | |
| | AND | | |
| 2 - Patient is currently receiving specialized nutritional support (e.g., intravenous parenteral nutrition, fluid, and micronutrient supplements) | | | |
| AND | | | |
| 3 - Patient has not previously received 4 weeks of treatment with Zorbtive* | | | |
| Notes | *Treatment with Zorbtive will not be authorized beyond 4 weeks. Admi nistration for more than 4 weeks has not been adequately studied. | | |

| Product Name: Increlex | | |
|--|--|--|
| Diagnosis | Severe Primary IGF-1 Deficiency / Growth Hormone Gene Deletion | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - ONE of the following | g criteria: | |
| 1.1 Documentation of | ALL of the following: | |
| 1.1.1 Diagnosis of se | vere primary Insulin-like Growth Factor 1 (IGF-1) deficiency | |
| | AND | |
| 1.1.2 Height standard | deviation score less than or equal to -3.0 | |
| AND | | |
| 1.1.3 Basal IGF-1 standard deviation score less than or equal to -3.0 | | |
| AND | | |
| 1.1.4 Normal or elevated growth hormone levels | | |
| AND | | |
| 1.1.5 Documentation of open epiphyses on last bone radiograph | | |
| AND | | |
| 1.1.6 The patient will | not be treated with concurrent growth hormone therapy | |

AND

1.1.7 Prescribed by an endocrinologist

OR

1.2 ALL of the following:

1.2.1 Diagnosis of growth hormone gene deletion and has developed neutralizing antibodies to growth hormone

AND

1.2.2 Documentation of open epiphyses on last bone radiograph

AND

1.2.3 The patient will not be treated with concurrent growth hormone therapy

AND

1.2.4 Prescribed by an endocrinologist

| Product Name: Increlex | |
|------------------------|--|
| Diagnosis | Severe Primary IGF-1 Deficiency / Growth Hormone Gene Deletion |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Height increase of at least 2 centimeters per year over the previous year of treatment as documented by BOTH of the following:*

| | and date obtained and date obtained | |
|---|--|--|
| | AND | |
| 2 - Documentation of B | OTH of the following:* | |
| Expected adult height not obtainedExpected adult height goal | | |
| AND | | |
| 3 - Patient is not treated with concurrent growth hormone therapy | | |
| AND | | |
| 4 - Prescribed by an endocrinologist | | |
| Notes | *Documentation of previous height, current height and goal expected adult height will be required for renewal. | |

2. Background

| Benefit/Coverage/Program Information | |
|---|--|
| Human Growth Hormone: | |
| | |
| Preferred Agents: | |
| Somatropin (Genotropin [®] , Genotropin Miniquick [®] , or Norditropin Flexpro [®]) | |

Nonpreferred Agents:

Somatropin (Humatrope[®], Nutropin AQ Nuspin[®], Omnitrope[®], Saizen[®], Serostim[®], Zomacton[®], and Zorbtive[®])

Growth Stimulating Products :

Mecasermin (Increlex®)

Preferred and Non-Preferred Agents

| Preferred | | |
|----------------|--------------------------------|-----|
| GPI Number | | MSC |
| 30100020002121 | GENOTROPIN INJ 5MG | N |
| 30100020002134 | GENOTROPIN INJ 12MG | N |
| 30100020002166 | GENOTROPIN MINIQUICK INJ 0.2MG | N |
| 30100020002168 | GENOTROPIN MINIQUICK INJ 0.4MG | N |
| 30100020002170 | GENOTROPIN MINIQUICK INJ 0.6MG | N |
| 30100020002172 | GENOTROPIN MINIQUICK INJ 0.8MG | N |
| 30100020002174 | GENOTROPIN MINIQUICK INJ 1MG | N |
| 30100020002176 | GENOTROPIN MINIQUICK INJ 1.2MG | N |
| 30100020002178 | GENOTROPIN MINIQUICK INJ 1.4MG | N |
| 30100020002180 | GENOTROPIN MINIQUICK INJ 1.6MG | N |
| 30100020002182 | GENOTROPIN MINIQUICK INJ 1.8MG | N |
| 30100020002184 | GENOTROPIN MINIQUICK INJ 2MG | N |
| 3010002000D212 | NORDITROPIN FLEXPRO 5MG | N |
| 3010002000D230 | NORDITROPIN FLEXPRO 10MG | N |
| 3010002000D240 | NORDITROPIN FLEXPRO 15MG | N |

| 3010002000D260 | NORDITROPIN FLEXPR | O 30MG N |
|----------------|--------------------------|---------------------------------|
| Non-Preferred | | |
| GPI Number | Drug Name and Strength | Drug Name |
| 30100020002125 | HUMATROPE INJ 6MG | HUMATROPE |
| 30100020002132 | HUMATROPE INJ 12MG | HUMATROPE |
| 30100020002150 | HUMATROPE INJ 24MG | HUMATROPE |
| 30160045002020 | MECASERMIN INJ 40 MG/4ML | INCRELEX |
| 3010002000D220 | NUTROPIN AQ INJ 10MG/2ML | NUTROPIN AQ NUSPIN 10 |
| 3010002000D250 | NUTROPIN AQ INJ 20MG/2ML | NUTROPIN AQ NUSPIN 20 |
| 3010002000D207 | NUTROPIN AQ INJ NUSPIN 5 | NUTROPIN AQ NUSPIN 5 |
| 30100020002123 | OMNITROPE INJ 5.8MG | OMNITROPE |
| 3010002000E210 | OMNITROPE INJ 5/1.5ML | OMNITROPE |
| 3010002000E213 | OMNITROPE INJ 10/1.5ML | OMNITROPE |
| 30100020102120 | SAIZEN INJ 5MG | SAIZEN |
| 30100020102130 | SAIZEN INJ 8.8MG | SAIZEN |
| 30100020102130 | SAIZENPREP INJ 8.8MG | SAIZENPREP RECONSTITUTIONKIT |
| 30100020102118 | SEROSTIM INJ 4MG | SEROSTIM |
| 30100020102121 | SEROSTIM INJ 5MG | SEROSTIM |
| 30100020102125 | SEROSTIM INJ 6MG | SEROSTIM |
| 30100020002121 | ZOMACTON INJ 5MG | ZOMACTON |
| 30100020002140 | ZOMACTON INJ 10MG | ZOMACTON |
| 30100020102132 | ZORBTIVE INJ 8.8MG | ZORBTIVE |

| Date | Notes |
|-----------|---|
| 8/22/2022 | Updated NP section notes, clarified approval duration of NP drugs is N/A. |

Haegarda - AZ

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99616 Haegarda - AZ

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Haegarda | |
|------------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of hereditary angioedema (HAE) as confirmed by ONE of the following:

1.1 C1 inhibitor (C1-INH) deficiency or dysfunction (Type I or II HAE) as documented by one of the following (per laboratory standard):

| C1-INH antigenic level below the lower limit of norm. | • | C1-INH | antigenic | level | below t | the lower | limit of | norma |
|---|---|--------|-----------|-------|---------|-----------|----------|-------|
|---|---|--------|-----------|-------|---------|-----------|----------|-------|

• C1-INH functional level below the lower limit of normal

OR

1.2 HAE with normal C1 inhibitor levels and one of the following:

- Confirmed presence of a FXII, angiopoietin-1 or plasminogen gene mutation
- Recurring angioedema attacks that are refractory to high-dose antihistamines with confirmed family history of angioedema

AND

2 - For prophylaxis against HAE attacks

AND

3 - Not used in combination with other approved C1 esterase inhibitors indicated for prophylaxis against HAE attacks (e.g. Cinryze)

AND

4 - Prescriber attests that patient has experienced attacks of a severity and/or frequency such that they would clinically benefit from prophylactic therapy with Haegarda

AND

5 - Prescribed by ONE of the following:

- Immunologist
- Allergist

| Product Name: Haegarda | | |
|------------------------|-------------|--|
| Approval Length | 12 month(s) | |

| Therapy Stage | Reauthorization | |
|--|---|--|
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| | ositive clinical response, defined as a clinically significant reduction in of hereditary angioedema (HAE) attacks, while on Haegarda therapy | |
| | AND | |
| | ization of on-demand therapies used for acute attacks (e.g., Berinert, letermined by claims information, while on Haegarda therapy | |
| | AND | |
| 3 - Prescribed for the prophylaxis of HAE attacks | | |
| | AND | |
| 4 - Not used in combina (e.g., Cinryze, Takhzyro | ation with other products indicated for prophylaxis against HAE attacks o) | |
| | AND | |
| 5 - Prescribed by ONE | of the following: | |
| Immunologist Allergist | | |

Allergist

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona SP to Medicaid Arizona SP for 7/1 eff |

HCG

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99463 HCG

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Novarel, Ovidrel, Brand Pregnyl, generic chorionic gonadotropin | | |
|---|------------------------------|--|
| Diagnosis | s Prepubertal Cryptorchidism | |
| Approval Length | 6 Week(s) | |
| Guideline Type | Prior Authorization | |
| | | |

Approval Criteria

1 - Diagnosis of prepubertal cryptorchidism not due to anatomical obstruction

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk copy C&S Arizona standard to Medicaid Arizona |

Hemangeol

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| n | | |
|---|--|--|
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of proliferating infantile hemangioma | | |
| | | |
| AND | | |
| | | |

2 - Prescriber provides a reason or special circumstance the patient cannot use generic propranolol oral solution

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk copy C&S Arizona standard to Medicaid Arizona |

Hemlibra

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99717 Hemlibra

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Hemlibra | |
|-------------------------------|--|
| Diagnosis | Hemophilia A with Factor VIII Inhibitors |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of hemophilia A | |

AND 2 - Patient has factor VIII inhibitors AND

3 - Prescribed for the prevention of bleeding episodes (i.e., routine prophylaxis)

| Product Name: Hemlibra | |
|------------------------|--|
| Diagnosis | Hemophilia A with Factor VIII Inhibitors |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Hemlibra therapy

| Product Name: Hemlibra | |
|------------------------|---|
| Diagnosis | Hemophilia A without Factor VIII Inhibitors |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- 1 ONE of the following:
- **1.1** BOTH of the following:
- 1.1.1 Diagnosis of severe hemophilia A

AND

1.1.2 Documentation of endogenous factor VIII levels less than 1% of normal factor VIII (less than 0.01 international units per milliliter)

OR

1.2 BOTH of the following:

1.2.1 ONE of the following

1.2.1.1 BOTH of the following

1.2.1.1.1 Diagnosis of moderate hemophilia A

AND

1.2.1.1.2 Documentation of endogenous factor VIII level greater than or equal to 1% and less than 5% (greater than or equal to 0.01 international units per milliliter [IU/mL] to less than 0.05 IU/mL)

OR

1.2.1.2 BOTH of the following

1.2.1.2.1 Diagnosis of mild hemophilia A

AND

1.2.1.2.2 Documentation of endogenous factor VIII level greater than 5% (greater than or equal to 0.05 international units per milliliter)

AND

1.2.2 Submission of medical records (e.g. chart notes, laboratory values) documenting a failure to meet clinical goals (e.g., continuation of spontaneous bleeds, inability to achieve

appropriate trough level, previous history of inhibitors) after a trial of prophylactic factor VIII replacement products

OR

1.3 BOTH of the following:

1.3.1 Patient is currently on Hemlibra therapy

AND

1.3.2 Diagnosis of hemophilia A

AND

2 - Hemlibra is prescribed for the prevention of bleeding episodes (i.e., routine prophylaxis)

AND

3 - Physician attestation that the patient is not to receive extended half-life factor VIII replacement products (e.g., Eloctate, Adynovate, Afstyla, Jivi) for the treatment of breakthrough bleeding episodes

| Product Name: Hemlibra | |
|------------------------|---|
| Diagnosis | Hemophilia A without Factor VIII Inhibitors |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Hemlibra therapy

AND

2 - Submission of medical records (e.g., chart notes, laboratory values) documenting that the patient is not receiving Hemlibra in combination with an extended half-life factor VIII replacement product (e.g., Eloctate, Adynovate, Afstyla, Jivi) for the treatment of breakthrough bleeding episodes. [Prescription claim history that does not show any concomitant extended half-life factor VIII replacement product claim within 60 days of reauthorization request may be used as documentation]

| Date | Notes |
|----------|-------------------------------------|
| 6/8/2021 | Arizona Medicaid 7.1 Implementation |

Hemophilia- Arizona

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99617 | Hemophilia- Arizona |
|----------|---------------------|
|----------|---------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Eloctate, Adynovate, Afstyla, Idelvion, Alprolix, Tretten | |
|---|-----------------------|
| Diagnosis | Hemophilia |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of hemophilia

AND

2 - Prescribing physician attests patient is not a suitable candidate for treatment with shorter acting half-life Factor VIII (recombinant) products [Kogenate FS, Kovaltry, Novoeight, or Nuwiq

AND

3 - One of the following:

3.1 Patient is not to receive routine infusions more frequently than 3 times per week

OR

3.2 Both of the following:

3.2.1 Patient is less than 12 years of age

AND

3.2.2 PK (pharmacokinetic) testing results suggest that more frequent than 3 times per week dosing is required

| Product Name: Eloctate, Adynovate, Afstyla, Idelvion, Alprolix, Tretten | | |
|---|---------------------|--|
| Diagnosis | Hemophilia | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Documentation of positive clinical response to therapy

AND

2 - One of the following:

2.1 Patient is not to receive routine infusions more frequently than 3 times per week

OR

2.2 Both of the following:

2.2.1 Patient is less than 12 years of age

AND

2.2.2 PK (pharmacokinetic) testing results suggest that more frequent than 3 times per week dosing is required

| Product Name: Jivi | | |
|--|-----------------------|--|
| Diagnosis | Hemophilia | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of hemophilia A | | |
| | AND | |
| 2 - Patient is 12 years of age or older | | |
| | AND | |

3 - Patient has previously received Factor VIII replacement therapy

AND

4 - Prescribing physician attests patient is not a suitable candidate for treatment with shorter acting half-life Factor VIII (recombinant) products [Kogenate FS, Kovaltry, Novoeight, or Nuwiq]

AND

5 - Patient is not to receive routine infusions more frequently than 2 times per week

| Product Name: Jivi | |
|--------------------|---------------------|
| Diagnosis | Hemophilia |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to therapy

AND

2 - Patient is not to receive routine infusions more frequently than 2 times per week

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona SP to Medicaid Arizona SP for 7/1 eff |

Hetlioz, Hetlioz LQ (tasimelteon)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-108664 | Hetlioz, Hetlioz LQ (tasimelteon) |
|-----------|-----------------------------------|
|-----------|-----------------------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 7/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Hetlioz capsule | |
|-------------------------------|--|
| Diagnosis | Non-24-Hour Sleep-Wake Disorder (Non-24) |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of non-24-hour sleep-wake disorder (also known as free-running disorder, free-

running or non-entrained type circadian rhythm sleep disorder, or hypernychthemeral syndrome)

AND

2 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting patient is totally blind (has no light perception)

AND

3 - Prescribed by or in consultation with one of the following:

- Specialist in sleep disorders
- Neurologist

| Product Name: Hetlioz capsule | |
|-------------------------------|--|
| Diagnosis | Non-24-Hour Sleep-Wake Disorder (Non-24) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting positive clinical response to therapy

| Product Name: Hetlioz capsule | |
|-------------------------------|------------------------------|
| Diagnosis | Smith-Magenis Syndrome (SMS) |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of Smith-Magenis Syndrome (SMS)

AND

2 - Patient is 16 years of age or older

AND

3 - Patient is experiencing nighttime sleep disturbances (i.e., difficulty falling asleep, frequent nighttime waking and early waking)

AND

4 - Prescribed by or in consultation with one of the following:

- Specialist in sleep disorders
- Neurologist

| Product Name: Hetlioz LQ suspension | |
|-------------------------------------|------------------------------|
| Diagnosis | Smith-Magenis Syndrome (SMS) |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of Smith-Magenis Syndrome (SMS)

AND

2 - Patient is 3 through 15 years of age

AND

3 - Patient is experiencing nighttime sleep disturbances (i.e., difficulty falling asleep, frequent nighttime waking and early waking)

AND

4 - Prescribed by or in consultation with one of the following:

- Specialist in sleep disorders
- Neurologist

| Product Name: Hetlioz capsule, Hetlioz LQ suspension | | |
|--|------------------------------|--|
| Diagnosis | Smith-Magenis Syndrome (SMS) | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting positive clinical response to therapy (i.e., improvement in nighttime total sleep time, improvement in nighttime sleep quality)

| Date | Notes |
|-----------|-------------|
| 6/23/2022 | New Program |

HIV (Fuzeon, Selzentry)

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-112911 | HIV (Fuzeon, S | elzentry) |
|-----------|----------------|-----------|
|-----------|----------------|-----------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 9/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Brand Selzentry tablets, generic maraviroc 150mg and 300mg tablets, Selzentry oral solution | |
|---|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| Approval Criteria 1 - One of the following: | |
| 1.1 All of the following: | |

1.1.1 Diagnosis of CCR5-tropic HIV-1 infection as confirmed by a highly sensitive tropism assay

AND

1.1.2 Patient is currently taking or will be prescribed an optimized background antiretroviral therapy regimen

AND

1.1.3 Prescribed by or in consultation with a clinician with HIV expertise

OR

1.2 For continuation of prior therapy

AND

2 - For generic maraviroc tablets and Selzentry oral solution ONLY; history of failure or intolerance to Brand Selzentry tablets

| Product Name: Fuzeon | |
|---------------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - One of the following: | |

1.1 All of the following:

1.1.1 Patient has been diagnosed with multidrug-resistant HIV-1 infection

AND

1.1.2 Patient is currently taking or will be prescribed an optimized background antiretroviral therapy regimen

AND

1.1.3 Prescribed by or in consultation with a clinician with HIV expertise

OR

1.2 For continuation of prior therapy

| Date | Notes |
|-----------|-------------|
| 8/29/2022 | New Program |

Humira

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99816 Humira

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Humira | |
|----------------------|---------------------------|
| Diagnosis | Rheumatoid Arthritis (RA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of moderately to severely active rheumatoid arthritis

AND

2 - Paid claims or submission of medical records (e.g., chart notes) documenting history of failure to a 3 month trial of ONE non-biologic disease modifying anti-rheumatic drug (DMARD) [e.g., methotrexate, leflunomide, sulfasalazine, hydroxychloroquine] at maximally indicated doses within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*

AND

3 - Patient is NOT receiving Humira in combination with ANY of the following:

- Biologic DMARD [e.g., Enbrel (etanercept), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

4 - Prescribed by or in consultation with a rheumatologist

| Notes | *Note: Claims history may be used in conjunction as documentation of |
|-------|--|
| | drug, date, and duration of trial |

| Product Name: Humira | |
|----------------------|---------------------------|
| Diagnosis | Rheumatoid Arthritis (RA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Humira therapy

AND

2 - Patient is NOT receiving Humira in combination with ANY of the following:

- Biologic disease-modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with a rheumatologist

| Product Name: Humira | |
|----------------------|---|
| Diagnosis | Polyarticular Juvenile Idiopathic Arthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of moderately to severely active polyarticular juvenile idiopathic arthritis

AND

- 2 Patient is NOT receiving Humira in combination with ANY of the following:
 - Biologic disease-modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Cimzia (certolizumab), Simponi (golimumab)]
 - Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
 - Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with a rheumatologist

Product Name: Humira

| Diagnosis | Polyarticular Juvenile Idiopathic Arthritis |
|-----------------|---|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Humira therapy

AND

2 - Patient is NOT receiving Humira in combination with ANY of the following:

- Biologic disease-modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with a rheumatologist

| Product Name: Humira | |
|----------------------|-----------------------|
| Diagnosis | Psoriatic Arthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of active psoriatic arthritis

AND

2 - Paid claims or submission of medical records (e.g., chart notes) documenting history of

failure to a 3 month trial of methotrexate at the maximally indicated dose within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*

AND

3 - Patient is NOT receiving Humira in combination with ANY of the following:

- Biologic disease-modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

4 - Prescribed by or in consultation with ONE of the following:

- Rheumatologist
- Dermatologist

| Notes | *Note: Claims history may be used in conjunction as documentation of |
|-------|--|
| | drug, date, and duration of trial |

| Product Name: Humira | |
|----------------------|---------------------|
| Diagnosis | Psoriatic Arthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Humira therapy

AND

2 - Patient is NOT receiving Humira in combination with ANY of the following:

- Biologic disease-modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

3 - Prescribed by or in consultation with ONE of the following:

- Rheumatologist
- Dermatologist

| Product Name: Humira | | |
|----------------------|-----------------------|--|
| Diagnosis | Plaque Psoriasis | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Diagnosis of moderate to severe chronic plaque psoriasis

AND

2 - Greater than or equal to 3% body surface area involvement, palmoplantar, facial, or genital involvement, or severe scalp psoriasis

AND

3 - Both of the following:

3.1 Paid claims or submission of medical records (e.g., chart notes) documenting history of failure to ONE of the following topical therapies, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial):*

• Corticosteroids (e.g., betamethasone, clobetasol, desonide)

| • | Vitamin D | analogs | (e.g., | calcitriol, | calcipotriene) |
|---|-----------|---------|--------|-------------|----------------|
|---|-----------|---------|--------|-------------|----------------|

- Tazarotene
- Calcineurin inhibitors (e.g., tacrolimus, pimecrolimus)
- Anthralin
- Coal tar

3.2 Paid claims or submission of medical records (e.g., chart notes) documenting history of failure to a 3 month trial of methotrexate at the maximally indicated dose within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*

AND

4 - Patient is NOT receiving Humira in combination with ANY of the following:

- Biologic disease-modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

5 - Prescribed by or in consultation with a dermatologist

| Notes | *Note: Claims history may be used in conjunction as documentation of |
|-------|--|
| | drug, date, and duration of trial |

| aque Psoriasis |
|-------------------|
| |
| e month(s) |
| eauthorization |
| ior Authorization |
| ea |

Approval Criteria

1 - Documentation of positive clinical response to Humira therapy

2 - Patient is NOT receiving Humira in combination with ANY of the following:

- Biologic disease-modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with a dermatologist

| Product Name: Humira | |
|----------------------|------------------------|
| Diagnosis | Ankylosing Spondylitis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of active ankylosing spondylitis

AND

2 - Paid claims or submission of medical records (e.g., chart notes) documenting history of failure to TWO NSAIDs (non-steroidal anti-inflammatory drugs) (e.g., ibuprofen, naproxen) at maximally indicated doses, each used for at least 4 weeks within the last 3 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trials)*

AND

3 - Patient is NOT receiving Humira in combination with ANY of the following:

• Biologic disease-modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Cimzia (certolizumab), Simponi (golimumab)]

- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

| AND | | |
|--|--|--|
| 4 - Prescribed by or in consultation with a rheumatologist | | |
| Notes | *Note: Claims history may be used in conjunction as documentation of drug, date, and duration of trial | |

| Product Name: Humira | |
|----------------------|------------------------|
| Diagnosis | Ankylosing Spondylitis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Humira therapy

AND

- 2 Patient is NOT receiving Humira in combination with ANY of the following:
 - Biologic disease-modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Cimzia (certolizumab), Simponi (golimumab)]
 - Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
 - Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with a rheumatologist

Product Name: Humira

| Diagnosis | Adult Crohn's Disease | |
|-----------------|-----------------------|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

1 - Diagnosis of moderately to severely active Crohn's disease

AND

2 - ONE of the following:

2.1 Paid claims or submission of medical records (e.g., chart notes) documenting history of failure to ONE of the following conventional therapies at maximally indicated doses within the last 3 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial):*

- Corticosteroids (e.g., prednisone, methylprednisolone, budesonide)
- Azathioprine (Imuran)
- 6-mercaptopurine (Purinethol)
- Methotrexate (Rheumatrex, Trexall)

OR

2.2 Patient has lost response or intolerant to infliximab (e.g., Remicade, Inflectra, Renflexis)

AND

3 - Patient is NOT receiving Humira in combination with ANY of the following:

- Biologic disease-modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

| 4 - Prescribed by or in consultation with a gastroenterologist | | |
|--|--|--|
| Notes | *Note: Claims history may be used in conjunction as documentation of drug, date, and duration of trial | |

| Product Name: Humira | | |
|----------------------|---------------------------|--|
| Diagnosis | Pediatric Crohn's Disease | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

1 - Diagnosis of moderately to severely active Crohn's disease

AND

2 - Paid claims or submission of medical records (e.g., chart notes) documenting history of failure to ONE of the following conventional therapies at maximally indicated doses within the last 3 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial):*

- Corticosteroids (e.g., prednisone, methylprednisolone, budesonide)
- Azathioprine (Imuran)
- 6-mercaptopurine (Purinethol)
- Methotrexate (Rheumatrex, Trexall)

AND

- **3** Patient is NOT receiving Humira in combination with ANY of the following:
 - Biologic disease-modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Cimzia (certolizumab), Simponi (golimumab)]
 - Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
 - Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

| 4 - Prescribed by or in consultation with a gastroenterologist | | |
|--|--|--|
| Notes | *Note: Claims history may be used in conjunction as documentation of drug, date, and duration of trial | |

| Product Name: Humira | |
|----------------------|-----------------------|
| Diagnosis | Ulcerative Colitis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

1 - Diagnosis of moderately to severely active ulcerative colitis

AND

2 - Paid claims or submission of medical records (e.g., chart notes) documenting history of failure to ONE of the following conventional therapies at maximally indicated doses within the last 3 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial):*

- Corticosteroids (e.g., prednisone, methylprednisolone, budesonide)
- 6-mercaptopurine (Purinethol)
- Azathioprine (Imuran)
- Aminosalicylates (e.g., mesalamine, sulfasalazine)

AND

- **3** Patient is NOT receiving Humira in combination with ANY of the following:
 - Biologic disease-modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Cimzia (certolizumab), Simponi (golimumab)]
 - Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
 - Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

| 4 - Prescribed by or in | consultation with a gastroenterologist |
|-------------------------|--|
| Notes | *Note: Claims history may be used in conjunction as documentation of drug, date, and duration of trial |

| Product Name: Humira | |
|----------------------|--|
| Diagnosis | Adult Crohn's Disease, Pediatric Crohn's Disease, Ulcerative Colitis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Documentation of positive clinical response to Humira therapy

AND

2 - Patient is NOT receiving Humira in combination with ANY of the following:

- Biologic disease-modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with a gastroenterologist

| Product Name: Humira | |
|----------------------|--------------------------|
| Diagnosis | Hidradenitis Suppurativa |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

| 1 - Diagnosis of moderate to severe hidradenitis suppurativa (i.e., Hurley Stage II or III) | | |
|---|--|--|
| | AND | |
| 2 - Paid claims or submission of medical records (e.g., chart notes) documenting history of failure to at least ONE oral antibiotic (e.g., doxycycline, clindamycin, rifampin) at maximally indicated doses within the last 3 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)* | | |
| | AND | |
| 3 - Patient is NOT rece | iving Humira in combination with ANY of the following: | |
| Biologic disease-modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Cimzia (certolizumab), Simponi (golimumab)] Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)] Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)] | | |
| AND | | |
| 4 - Prescribed by or in consultation with a dermatologist | | |
| Notes | *Note: Claims history may be used in conjunction as documentation of drug, date, and duration of trial | |

| Product Name: Humira | |
|----------------------|--------------------------|
| Diagnosis | Hidradenitis Suppurativa |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Г

1 - Documentation of positive clinical response to Humira therapy

2 - Patient is NOT receiving Humira in combination with ANY of the following:

- Biologic disease-modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with a dermatologist

| Product Name: Humira | |
|----------------------|-----------------------|
| Diagnosis | Uveitis (UV) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of non-infectious uveitis

AND

2 - Uveitis is classified as ONE of the following:

- intermediate
- posterior
- panuveitis

AND

3 - Paid claims or submission of medical records (e.g., chart notes) documenting history of failure to at least ONE corticosteroid (e.g., prednisolone, prednisone) at maximally indicated

dose within the last 3 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*

AND

4 - Paid claims or submission of medical records (e.g., chart notes) documenting history of failure to at least ONE systemic non-biologic immunosuppressant (e.g., methotrexate, cyclosporine, azathioprine, mycophenolate) at a maximally indicated dose within the last 3 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*

AND

5 - Patient is NOT receiving Humira in combination with ANY of the following:

- Biologic disease-modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

6 - Prescribed by or in consultation with ONE of the following:

- Rheumatologist
- Ophthalmologist

| *Note: Claims history may be used in conjunction as documentation of drug, date, and duration of trial |
|--|
| |

| Product Name: Humira | |
|----------------------|---------------------|
| Diagnosis | Uveitis (UV) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Documentation of positive clinical response to Humira therapy

AND

2 - Patient is NOT receiving Humira in combination with ANY of the following:

- Biologic disease-modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with ONE of the following:

- Rheumatologist
- Ophthalmologist

| Date | Notes |
|-----------|---|
| 12/6/2021 | Added "history of paid claims or submission of medical records (e.g., chart notes)" to all t/f criteria. Changed concomitant use criteria from ' one' to ALL. |

Hydroxychloroquine

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99465 | Hydroxychloroquine |
|----------|--------------------|
|----------|--------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Plaquenil, generic hydroxychloroquine | | |
|--|----------------|--|
| Guideline Type | Quantity Limit | |
| | | |
| Approval Criteria | | |
| 1 - ONE of the following: | | |
| 1.1 Treatment of chronic discoid lupus erythematosus or systemic lupus erythematosus | | |
| OR | | |

| 1.2 Treatment of rheu | matoid arthritis |
|-----------------------|---|
| | OR |
| 1.3 Prophylaxis of ma | laria in geographic areas where chloroquine resistance is not reported |
| | OR |
| 1.4 Treatment of unco | omplicated malaria |
| Notes | Authorization will be issued for 6 months up to a quantity of 120 tablet s per 30 days. |

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk copy C&S Arizona standard to Medicaid Arizona |

Hyftor (sirolimus) topical gel

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-114463 | Hyftor (sirolimus) topical gel |
|-----------|--------------------------------|
|-----------|--------------------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Hyftor | |
|----------------------|-----------------------|
| Approval Length | 4 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| | |

Approval Criteria

1 - Diagnosis of facial angiofibroma associated with tuberous sclerosis complex

2 - Patient is 6 years of age or older

AND

3 - Patient is not a candidate for laser therapy or surgical treatments

AND

4 - Prescribed by or in consultation with a dermatologist

| Product Name: Hyftor | |
|----------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to therapy (e.g., improvement in size or redness of facial angiofibroma)

| Date | Notes |
|-----------|-------------|
| 9/26/2022 | New program |

Igalmi (dexmedetomidine)

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-110775 | Igalmi (dexmedetomidine) |
|-----------|--------------------------|
|-----------|--------------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 8/15/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Igalmi | |
|--|---------------------|
| Approval Length | 14 Days [A] |
| Guideline Type | Prior Authorization |
| | |
| | |
| Approval Criteria | |
| Approval Criteria 1 - One of the following | diagnoses: |

2 - For the treatment of acute agitation

AND

3 - Trial and failure, contraindication or intolerance to at least two preferred products used in acute agitation (e.g., olanzapine, ziprasidone)

AND

4 - Patient is currently being managed with maintenance medication for their underlying disorder (e.g., aripiprazole, olanzapine, quetiapine, lithium, valproic acid)

| Date | Notes |
|----------|-------------|
| 8/4/2022 | New Program |

Ilaris

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99618 Ilaris

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Ilaris | |
|--|--|
| Cryopyrin-Associated Periodic Syndromes (CAPS) | |
| 12 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |

Approval Criteria

- **1** Diagnosis of ONE of the following:
 - Familial cold autoinflammatory syndrome (FCAS)

• Muckle-Wells Syndrome (MWS)

AND

2 - Diagnosis is made by, or in consultation with, a rheumatologist or immunologist with expertise in the diagnosis of FCAS and MWS

| Product Name: Ilaris | |
|----------------------|--|
| Diagnosis | Cryopyrin-Associated Periodic Syndromes (CAPS) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- 1 Patient is currently on Ilaris therapy for ONE of the following:
 - Familial cold autoinflammatory syndrome (FCAS)
 - Muckle-Wells Syndrome (MWS)

AND

2 - Documentation of positive clinical response to llaris therapy

| Product Name: Ilaris | |
|----------------------|---|
| Diagnosis | Tumor Necrosis Factor (TNF) Receptor-Associated Periodic Syndrome (TRAPS) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of tumor necrosis factor (TNF) receptor-associated periodic syndrome (TRAPS)

2 - Diagnosis is made by, or in consultation with, a rheumatologist or immunologist with expertise in the diagnosis of TRAPS

| Product Name: Ilaris | |
|----------------------|---|
| Diagnosis | Tumor Necrosis Factor (TNF) Receptor-Associated Periodic Syndrome (TRAPS) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient is currently on Ilaris therapy for tumor necrosis factor (TNF) receptor-associated periodic syndrome (TRAPS)

AND

2 - Documentation of positive clinical response to Ilaris therapy, defined as a decrease in frequency or severity of attacks

| Product Name: Ilaris | |
|----------------------|---|
| Diagnosis | Hyperimmunoglobulin D (Hyper-IgD) Syndrome (HIDS)/Mevalonate Kinase Deficiency (MKD) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- **1** Diagnosis of ONE of the following
 - Hyperimmunoglobulin D (Hyper-IgD) Syndrome (HIDS)

• Mevalonate Kinase Deficiency (MKD)

AND

2 - Diagnosis is made by, or in consultation with, a rheumatologist or immunologist with expertise in the diagnosis of HIDS or MKD

| Product Name: Ilaris | |
|----------------------|---|
| Diagnosis | Hyperimmunoglobulin D (Hyper-IgD) Syndrome (HIDS)/Mevalonate Kinase Deficiency (MKD) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- 1 Patient is currently on Ilaris therapy for ONE of the following:
 - Hyperimmunoglobulin D (Hyper-IgD) Syndrome (HIDS)
 - Mevalonate Kinase Deficiency (MKD)

AND

2 - Documentation of positive clinical response to Ilaris therapy, defined as a decrease in frequency or severity of attacks

| Product Name: Ilaris | |
|----------------------|------------------------------------|
| Diagnosis | Familial Mediterranean Fever (FMF) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of Familial Mediterranean Fever (FMF)

2 - Diagnosis is made by, or in consultation with, a rheumatologist or immunologist with expertise in the diagnosis of FMF

AND

3 - History of failure, contraindication, or intolerance to colchicine

| Product Name: Ilaris | |
|----------------------|------------------------------------|
| Diagnosis | Familial Mediterranean Fever (FMF) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient is currently on Ilaris therapy for Familial Mediterranean Fever (FMF)

AND

2 - Documentation of positive clinical response to Ilaris therapy, defined by a decrease in index disease flare or normalization of CRP (C-reactive protein)

| Product Name: Ilaris | |
|----------------------|---|
| Diagnosis | Systemic Juvenile Idiopathic Arthritis (SJIA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of systemic juvenile idiopathic arthritis (SJIA)

AND

2 - Diagnosis is made by, or in consultation with, a rheumatologist or immunologist with expertise in the diagnosis of SJIA

AND

3 - Patient is not receiving Ilaris in combination with another biologic (e.g., Actemra)

| Product Name: Ilaris | |
|----------------------|---|
| Diagnosis | Systemic Juvenile Idiopathic Arthritis (SJIA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient is currently on Ilaris therapy for systemic juvenile idiopathic arthritis (SJIA)

AND

2 - Documentation of positive clinical response to Ilaris therapy

AND

3 - Patient is not receiving Ilaris in combination with another biologic (e.g., Actemra)

| Product Name: Ilaris | |
|----------------------|--|
| Diagnosis | Still's Disease [Adult-Onset Still's Disease (AOSD)] |
| Approval Length | 12 month(s) |

| Therapy Stage | Initial Authorization | | |
|---|---|--|--|
| Guideline Type | Prior Authorization | | |
| | | | |
| Approval Criteria | | | |
| 1 - Diagnosis of Adult C | 1 - Diagnosis of Adult Onset Still's Disease (AOSD) | | |
| | AND | | |
| | AND | | |
| 2 - Diagnosis is made by, or in consultation with, a rheumatologist or immunologist with expertise in the diagnosis of Still's Disease | | | |
| | AND | | |
| 3 - Patient is not receiv | ing Ilaris in combination with another biologic (e.g., Actemra) | | |

| Product Name: Ilaris | |
|----------------------|--|
| Diagnosis | Still's Disease [Adult-Onset Still's Disease (AOSD)] |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Patient is currently on Ilaris therapy for Adult Onset Still's Disease (AOSD)

AND

2 - Documentation of positive clinical response to Ilaris therapy

AND

3 - Patient is not receiving Ilaris in combination with another biologic (e.g., Actemra)

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Ilumya - Arizona

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99718 | Ilumya - Arizona |
|----------|------------------|
|----------|------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Ilumya | |
|----------------------|---|
| Diagnosis | Chronic Moderate to Severe Plaque Psoriasis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| | |

Approval Criteria

1 - ONE of the following:

1.1 ALL of the following:

1.1.1 Diagnosis of chronic moderate to severe plaque psoriasis

AND

1.1.2 Greater than or equal to 3 percent body surface area involvement, palmoplantar, facial, genital involvement, or severe scalp psoriasis

AND

1.1.3 History of failure, to ONE of the following topical therapies, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial):*

- Corticosteroids (e.g., betamethasone, clobetasol, desonide)
- Vitamin D analogs (e.g., calcitriol, calcipotriene)
- Tazarotene
- Calcineurin inhibitors (e.g., tacrolimus, pimecrolimus)
- Anthralin
- Coal tar

AND

1.1.4 History of failure to a 3 month trial of methotrexate at the maximally indicated dose within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*

AND

1.1.5 History of failure, contraindication, or intolerance to ALL of the following preferred biologic products (document drug, date, and duration of trial):*

- Humira (adalimumab)
- Enbrel (etanercept)
- Otezla (apremilast)

AND

1.1.6 Patient is NOT receiving Ilumya in combination with ONE of the following:

| Cimzia (certolizum (abatacept)] • Janus kinase inhib | (disease modifying anti-rheumatic drug) [e.g., Humira (adalimumab), hab), Simponi (golimumab), Cosentyx (secukinumab), Orencia bitor [e.g., Xeljanz (tofacitinib)] e 4 (PDE4) inhibitor [e.g. Otezla (apremilast)] |
|--|---|
| | AND |
| 1.1.7 Prescribed by or ir | n consultation with a dermatologist |
| | OR |
| 1.2 ALL of the following: | |
| , | y on Ilumya therapy as documented by claims history or medical date, and duration of therapy) |
| | AND |
| 1.2.2 Diagnosis of chror | nic moderate to severe plaque psoriasis |
| | AND |
| 1.2.3 Patient is NOT rec | ceiving Ilumya in combination with ONE of the following: |
| Biologic DMARD [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Cosentyx (secukinumab), Orencia (abatacept)] Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)] Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)] | |
| AND | |
| 1.2.4 Prescribed by or ir | n consultation with a dermatologist |
| | Note: Claims history may be used in conjunction as documentation of Irug, date, and duration of trial |

| Product Name: Ilumya | |
|----------------------|---|
| Diagnosis | Chronic Moderate to Severe Plaque Psoriasis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Documentation of positive clinical response to Ilumya therapy

AND

- **2** Patient is NOT receiving Ilumya in combination with ONE of the following:
 - Biologic DMARD (disease modifying anti-rheumatic drug) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Cosentyx (secukinumab), Orencia (abatacept)]
 - Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
 - Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with a dermatologist

| Date | Notes |
|-----------|-------------------------------------|
| 5/13/2021 | Arizona Medicaid 7.1 Implementation |

Imbruvica

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99748 Imbruvica

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Imbruvica | |
|-------------------------|-----------------------|
| Diagnosis | B-Cell Lymphoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | · |

Approval Criteria

1 - ONE of the following:

1.1 BOTH of the following:

1.1.1 Diagnosis of mantle cell lymphoma (MCL) AND **1.1.2** ONE of the following: Patient has received at least one prior therapy for MCL • Used in pre-treatment therapy in combination with Rituxan (rituximab) to limit the • number of cycles with RHyperCVAD (cyclophosphamide, vincristine, doxorubicin, and dexamethasone) regimen OR **1.2** Diagnosis of ONE of the following: Chronic Lymphocytic Leukemia (CLL) • Small Lymphocytic Lymphoma (SLL) OR **1.3** BOTH of the following: **1.3.1** Diagnosis of ONE of the following: Follicular lymphoma (grade 1-2) • Diffuse large B-cell lymphoma [non-GCB DLBCL (non-germinal center B-cell diffuse • large B-cell) and non-candidate for transplant] Acquired immune deficiency syndrome (AIDS)-related B-cell lymphoma ٠ Post-transplant lymphoproliferative disorders • Histologic transformation to diffuse large B-cell lymphoma • Hairy cell leukemia • Nodal or splenic marginal zone lymphoma (MZL) • Gastric MALT (mucosa-associated lymphoid tissue) lymphoma ٠ Nongastric MALT lymphoma • High grade B-cell lymphoma • AND **1.3.2** Used as second-line or a subsequent therapy

| Product Name: Imbruvica | | |
|-------------------------|--|--|
| Diagnosis | Naldenström's Macroglobulinemia/Lymphoplasmacytic Lymphoma | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |

1 - Diagnosis of Waldenström's Macroglobulinemia/Lymphoplasmacytic Lymphoma

| Product Name: Imbruvica | |
|-------------------------|-----------------------|
| Diagnosis | Primary CNS Lymphoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of primary central nervous system (CNS) lymphoma

AND

2 - ONE of the following:

2.1 Used as second-line or a subsequent therapy

OR

2.2 Used as induction therapy if the patient is unsuitable or intolerant to high-dose methotrexate

| Product Name: Imbruvica | |
|-------------------------|---|
| Diagnosis | B-Cell Lymphoma, Waldenström's Macroglobulinemia/Lymphoplasmacytic Lymphoma, Primary CNS Lymphoma |

| Approval Length | 12 month(s) |
|-----------------|---------------------|
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Patient does not show evidence of progressive disease while on Imbruvica therapy

| Product Name: Imbruvica | |
|-------------------------|-----------------------------------|
| Diagnosis | Chronic Graft Versus Host Disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of chronic graft versus host disease

AND

2 - History of failure of at least one other systemic therapy [e.g. corticosteroids, mycophenolate, etc.]

| Product Name: Imbruvica | |
|-------------------------|-----------------------------------|
| Diagnosis | Chronic Graft Versus Host Disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient shows evidence of positive clinical response while on Imbruvica therapy

| Product Name: Imbruvica | |
|-------------------------|--------------------------|
| Diagnosis | NCCN Recommended Regimen |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

1 - Imbruvica will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Imbruvica | |
|-------------------------|--------------------------|
| Diagnosis | NCCN Recommended Regimen |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Imbruvica therapy

| Date | Notes |
|----------|-------------------------------------|
| 6/2/2021 | Arizona Medicaid 7.1 Implementation |

Immune Globulin- Arizona

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99783 | Immune Globulin- Arizona |
|----------|--------------------------|
|----------|--------------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan
S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif,
Gammaplex, Octagam, Panzyga, Gammaked, XembifyDiagnosisAsthma (severe, persistent, high-dose steroid-dependent)Approval Length12 month(s)Therapy StageInitial AuthorizationGuideline TypePrior Authorization

Approval Criteria

1 - One of the following diagnoses:

| Severe asthma |
|-----------------------------------|
|-----------------------------------|

• Persistent asthma

• High-dose steroid-dependent asthma

AND

2 - Medical records documenting BOTH of the following:

- History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable
- Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested

AND

3 - Patient is receiving optimal conventional asthma therapy (e.g., high-dose inhaled glucocorticoids, short- and long-acting inhaled β agonists)

AND

4 - History of failure, contraindication, or intolerance to at least TWO of the following:

- Anti-IgE therapy [e.g., Xolair (omalizumab)]
- Anti-interleukin 4 therapy [e.g., Dupixent (dupilumab)]
- Anti-interleukin 5 therapy [e.g., Nucala (mepolizumab), Cinqair (resilizumab), Fasenra (benralizumab)]

AND

5 - Patient has required continuous oral glucocorticoid therapy for a minimum of 2 months prior to the decision to initiate immune globulin therapy

AND

6 - For long term treatment, documentation of titration to the minimum dose and frequency needed to maintain a sustained clinical effect

7 - Prescribed by or in consultation with a pulmonologist or allergist or immunologist

AND

8 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to ALL the following products. (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products)

- Flebogamma
- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Gamunex-C
- Hizentra Vial & Syringe
- Privigen

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Asthma (severe, persistent, high-dose steroid-dependent) |
|-----------------|--|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to immune globulin therapy

AND

2 - Statement of expected frequency and duration of proposed immune globulin treatment

3 - For long term treatment, documentation of titration to the minimum effective dose and frequency needed to maintain a sustained clinical response

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Autoimmune Bullous Disease [pemphigus vulgaris, pemphigus foliaceus, bullous pemphigoid, mucous membrane (cicatricial) pemphigoid, epidermolysis bullosa acquisita, pemphigoid gestationis, linear IgA bullous dermatosis] |
|-----------------|---|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of Autoimmune Bullous Disease [pemphigus vulgaris, pemphigus foliaceus, bullous pemphigoid, mucous membrane (cicatricial) pemphigoid, epidermolysis bullosa acquisita, pemphigoid gestationis, linear IgA bullous dermatosis]

AND

2 - Medical records documenting BOTH of the following:

- History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable
- Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested

AND

3 - Extensive and debilitating disease

4 - History of failure, contraindication, or intolerance to systemic corticosteroids with concurrent immunosuppressive treatment (e.g., azathioprine, cyclophosphamide, mycophenolate mofetil)

AND

5 - Intravenous immunoglobulin (IVIG) dose does not exceed 1,000 to 2,000 milligrams (mg) per kilogram (kg) per month divided into 3 equal doses, each given over 3 consecutive days or 400 mg per kg per day given over 5 consecutive days per month. IVIG administration may be repeated monthly as needed for patients requiring maintenance therapy. Dosing interval may need to be adjusted in patients with severe comorbidities

AND

6 - For long term treatment, documentation of titration to the minimum dose and frequency needed to maintain a sustained clinical effect

AND

7 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to ALL the following products. (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products)

- Flebogamma
- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Gamunex-C
- Hizentra Vial & Syringe
- Privigen

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Autoimmune Bullous Disease [pemphigus vulgaris, pemphigus foliaceus, bullous pemphigoid, mucous membrane (cicatricial) pemphigoid, epidermolysis bullosa acquisita, pemphigoid gestationis, linear IgA bullous dermatosis] |
|-----------------|---|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to immune globulin therapy

AND

2 - Statement of expected frequency and duration of proposed immune globulin treatment

AND

3 - For long term treatment, documentation of titration to the minimum effective dose and frequency needed to maintain a sustained clinical response

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Bone Marrow Transplant (BMT) |
|-----------------|------------------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - ONE of the following uses:

- Prevention of acute graft vs. host disease (GVHD)
- Prevention of infection

2 - Medical records documenting BOTH of the following:

- History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable
- Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested

AND

3 - Confirmed allogeneic bone marrow transplant within the last 100 days

AND

4 - Documented severe hypogammaglobulinemia [Immunoglobulin (IgG) less than 400 milligrams (mg) per deciliter (dL)]

AND

5 - Intravenous immunoglobulin (IVIG) dose does not exceed 500 mg per kilogram (kg) once weekly for the first 90 days of therapy, then monthly up to 360 days after transplantation

AND

6 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to ALL the following products. (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products)

- Flebogamma
- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Gamunex-C
- Hizentra Vial & Syringe
- Privigen

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Bone Marrow Transplant (BMT) |
|-----------------|------------------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to immune globulin therapy

AND

2 - Statement of expected frequency and duration of proposed immune globulin treatment

AND

3 - For long term treatment, documentation of titration to the minimum effective dose and frequency needed to maintain a sustained clinical response

| Product Name: : HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify | |
|--|---|
| Diagnosis | Chronic Inflammatory Demyelinating Polyneuropathy |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of chronic inflammatory demyelinating polyneuropathy as confirmed by ALL of the following:

1.1 Progressive symptoms present for at least 2 months

1.2 Symptomatic polyradiculoneuropathy as indicated by progressive or relapsing motor or sensory impairment of more than one limb

AND

1.3 Electrodiagnostic findings [consistent with European Federation of Neurological Societies/Peripheral Nerve Society (EFNS/PNS) guidelines for definite chronic inflammatory demyelinating polyradiculoneuropathy (CIDP)] indicating at least ONE of the following criteria are present:

- Motor distal latency prolongation in 2 nerves
- Reduction of motor conduction velocity in 2 nerves
- Prolongation of F-wave latency in 2 nerves
- Absence of F-waves in at least 1 nerve
- Partial motor conduction block of at least 1 motor nerve
- Abnormal temporal dispersion in at least 2 nerves
- Distal compound muscle action potential (CMAP) duration increase in at least 1 nerve

AND

- 2 Medical records documenting BOTH of the following:
 - History and physical examination documenting the severity of the condition, including frequency and severity of infections where
 - Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested

AND

3 - Prescribed by or in consultation with a neurologist

AND

4 - Intravenous immunoglobulin (IVIG) dose does not exceed 2,000 milligrams (mg) per kilogram (kg) per month given over 2 to 5 consecutive days administered in up to six monthly infusions. Dosing interval may need to be adjusted in patients with severe comorbidities.

5 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to ALL the following products. (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products)

- Flebogamma
- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Gamunex-C
- Hizentra Vial & Syringe
- Privigen

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Chronic Inflammatory Demyelinating Polyneuropathy |
|-----------------|---|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to therapy as measured by an objective scale [e.g., Rankin, Modified Rankin, Medical Research Council (MRC) scale]

AND

2 - For long term treatment, documentation of titration to the minimum effective dose and frequency needed to maintain a sustained clinical response

AND

3 - Prescribed by or in consultation with a neurologist

4 - Intravenous immunoglobulin (IVIG) dose does not exceed 2,000 milligrams (mg) per kilogram (kg) per month given over 2 to 5 consecutive days. IVIG administration may be repeated monthly as needed to prevent exacerbation. Dosing interval may need to be adjusted in patients with severe comorbidities.

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Prevention of infection in B-cell Chronic Lymphocytic Leukemia (CLL) |
|-----------------|--|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of B-cell chronic lymphocytic leukemia (CLL)

AND

- 2 Medical records documenting BOTH of the following:
 - History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable
 - Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested

AND

3 - ONE of the following:

- Documented hypogammaglobulinemia [Immunoglobulin (IgG) less than 500 milligrams (mg) per deciliter (dL)]
- History of bacterial infection(s) associated with B-cell CLL

4 - Intravenous immunoglobulin (IVIG) dose does not exceed 400 milligrams (mg) per kilogram (kg) every 3 to 4 weeks

AND

5 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to the following products.* (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products)

- Flebogamma
- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Gamunex-C
- Hizentra Vial & Syringe
- Privigen

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Prevention of infection in B-cell Chronic Lymphocytic Leukemia (CLL) |
|-----------------|--|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to immune globulin therapy

AND

2 - Statement of expected frequency and duration of proposed immune globulin treatment

3 - For long term treatment, documentation of titration to the minimum effective dose and frequency needed to maintain a sustained clinical response

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Dermatomyositis or polymyositis |
|-----------------|---------------------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of dermatomyositis or polymyositis

AND

- 2 Medical records documenting BOTH of the following:
 - History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable
 - Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested

AND

3 - History of failure, contraindication, or intolerance to immunosuppressive therapy (e.g., azathioprine, corticosteroids, cyclophosphamide, methotrexate)

AND

4 - Intravenous immunoglobulin (IVIG) dose does not exceed 2,000 milligrams (mg) per

kilogram (kg) per month given over 2 to 5 consecutive days administered as monthly infusions. Dosing interval may need to be adjusted in patients with severe comorbidities

AND

5 - For long term treatment, documentation of titration to the minimum dose and frequency needed to maintain a sustained clinical effect

AND

6 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to the following products.* (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products)

- Flebogamma
- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Gamunex-C
- Hizentra Vial & Syringe
- Privigen

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Dermatomyositis or polymyositis |
|-----------------|---------------------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to immune globulin therapy

2 - Statement of expected frequency and duration of proposed immune globulin treatment

AND

3 - For long term treatment, documentation of titration to the minimum effective dose and frequency needed to maintain a sustained clinical response

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Diabetes Mellitus |
|-----------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient is newly diagnosed with insulin dependent (type 1) diabetes mellitus

AND

- 2 Medical records documenting BOTH of the following:
 - History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable
 - Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested

AND

3 - Patient is not a candidate for or is refractory to insulin therapy

AND

4 - If the request is for a non-preferred product, there must be a history of failure,

contraindication or intolerance to ALL the following products.(Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products)

- Flebogamma
- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Gamunex-C
- Hizentra Vial & Syringe
- Privigen

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Diabetes Mellitus |
|-----------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to immune globulin therapy

AND

2 - Statement of expected frequency and duration of proposed immune globulin treatment

AND

3 - For long term treatment, documentation of titration to the minimum effective dose and frequency needed to maintain a sustained clinical response

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Feto-neonatal Alloimmune Thrombocytopenia (AIT) |
|-----------|---|
|-----------|---|

| Approval Length | 12 month(s) | | |
|--|--|--|--|
| Therapy Stage | Initial Authorization | | |
| Guideline Type | Prior Authorization | | |
| | | | |
| Approval Criteria | | | |
| 1 - For pregnant wome | n all of the following: | | |
| 1.1 Diagnosis of feto-r | neonatal alloimmune thrombocytopenia (AIT) | | |
| | AND | | |
| 1.2 ONE of the followi | ng: | | |
| Family history or | | | |
| | AND | | |
| 1.3 ONE of the followi | ng: | | |
| 1.3.1 Intravenous immunoglobulin (IVIG) dose does not exceed 1,000 milligrams (mg) per kilogram (kg) once weekly until delivery | | | |
| | OR | | |
| 1.3.2 BOTH of the fol | lowing: | | |
| hemorrhage or | rn is considered to be at high risk for developing intracranial other severe complication of AIT not exceed 2,000 mg/kg once weekly until delivery | | |
| | AND | | |
| 2 - For newborns all of | the following: | | |
| 2.1 Diagnosis of feto-r | neonatal alloimmune thrombocytopenia | | |

2.2 Thrombocytopenia that persists after transfusion of antigen-negative compatible platelets

AND

3 - Medical records documenting BOTH of the following:

- History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable
- Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested

AND

4 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to ALL the following products. (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products)

- Flebogamma
- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Gamunex-C
- Hizentra Vial & Syringe
- Privigen

| Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify | |
|--|---|
| Diagnosis | Feto-neonatal Alloimmune Thrombocytopenia (AIT) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to immune globulin therapy

AND

2 - Statement of expected frequency and duration of proposed immune globulin treatment

AND

3 - For long term treatment, documentation of titration to the minimum effective dose and frequency needed to maintain a sustained clinical response

| Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan |
|--|
| S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, |
| Gammaplex, Octagam, Panzyga, Gammaked, Xembify |

| Diagnosis | Graves' ophthalmopathy Guillain-Barré syndrome (GBS) |
|-----------------|--|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of Guillain-Barré Syndrome

AND

- 2 Medical records documenting BOTH of the following:
 - History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable
 - Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested

3 - Severe disease requiring aid to walk

AND

4 - Onset of neuropathic symptoms within the last four weeks

AND

5 - Prescribed by or in consultation with a neurologist

AND

6 - Intravenous immunoglobulin (IVIG) dose does not exceed 2,000 milligrams (mg) per kilogram (kg) per month given over 2 to 5 consecutive days. IVIG administration may be repeated in up to three monthly infusions. Dosing interval may need to be adjusted in patients with severe comorbidities

AND

7 - For long term treatment, documentation of titration to the minimum dose and frequency needed to maintain a sustained clinical effect

AND

8 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to ALL the following. (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products)

- Flebogamma
- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Gamunex-C
- Hizentra Vial & Syringe
- Privigen

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Graves' ophthalmopathy Guillain-Barré syndrome (GBS) |
|-----------------|--|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to immune globulin therapy

AND

2 - Statement of expected frequency and duration of proposed immune globulin treatment

AND

3 - For long term treatment, documentation of titration to the minimum effective dose and frequency needed to maintain a sustained clinical response

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Prevention of bacterial infection in pediatric HIV |
|-----------------|--|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of HIV disease

- 2 Medical records documenting BOTH of the following:
 - History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable
 - Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested

3 - Patient age less than or equal to 13 years of age

AND

- **4** ONE of the following:
 - Documented hypogammaglobulinemia [Immunoglobulin (IgG) less than 400 milligrams (mg) per deciliter (dL)]
 - Functional antibody deficiency as demonstrated by either poor specific antibody titers or recurrent bacterial infections

AND

5 - Intravenous immunoglobulin (IVIG) dose does not exceed 400 mg per kilogram (kg) every 28 days

AND

6 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to ALL the following products. (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products

- Flebogamma
- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Hizentra Vial & Syringe
- Privigen

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Prevention of bacterial infection in pediatric HIV |
|-----------------|--|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to immune globulin therapy

AND

2 - Statement of expected frequency and duration of proposed immune globulin treatment

AND

3 - For long term treatment, documentation of titration to the minimum effective dose and frequency needed to maintain a sustained clinical response

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Immune thrombocytopenia [Idiopathic thrombocytopenic purpura (ITP)] |
|-----------------|---|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - One of the following:

1.1 ALL of the following:

• Diagnosis of acute thrombocytopenic purpura (ITP)

| • | Documented platelet count less than 50 x 10^9 per Liter (L) (obtained within the past |
|---|---|
| | 30 days) |

• Intravenous immunoglobulin (IVIG) dose does not exceed 1,000 milligrams (mg) per kilogram(kg) per day for 1 to 2 days

OR

1.2 All of the following:

1.2.1 Diagnosis of chronic thrombocytopenic purpura (ITP)

AND

1.2.2 History of failure, contraindication, or intolerance to at least ONE of the following:

- Corticosteroids
- Splenectomy

AND

1.2.3 IVIG dose does not exceed 2,000 mg per kg per month given over 2 to 5 consecutive days. IVIG administration may be repeated monthly as needed to prevent exacerbation. Dosing interval should be adjusted depending upon response and titrated to the minimum effective dose that can be given at maximum intervals to maintain safe platelet levels.

AND

2 - Medical records documenting BOTH of the following:

- History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable
- Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested

AND

3 - If the request is for a non-preferred product, there is a history of failure, contraindication or intolerance to 3 preferred products.* (Note: In instances where there are fewer than three

preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to all of the preferred products

AND

4 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to ALL the following products. (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products)

- Flebogamma
- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Gamunex-C
- Hizentra Vial & Syringe
- Privigen

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Immune thrombocytopenia [Idiopathic thrombocytopenic purpura (ITP)] |
|-----------------|---|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to immune globulin therapy

AND

2 - Statement of expected frequency and duration of proposed immune globulin treatment

AND

3 - For long term treatment, documentation of titration to the minimum effective dose and frequency needed to maintain a sustained clinical response

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Kawasaki Disease |
|-----------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of Kawasaki disease

AND

2 - Medical records documenting BOTH of the following:

- History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable
- Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested

AND

3 - Intravenous immunoglobulin (IVIG) dose does not exceed 4,000 milligrams (mg) per kilograms (kg) for five consecutive days or a single dose of 2,000 mg per kg

AND

4 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to ALL the following products. (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products)

• Flebogamma

- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Gamunex-C
- Hizentra Vial & Syringe
- Privigen

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Kawasaki Disease |
|-----------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to immune globulin therapy

AND

2 - Statement of expected frequency and duration of proposed immune globulin treatment

AND

3 - For long term treatment, documentation of titration to the minimum effective dose and frequency needed to maintain a sustained clinical response

| S-D, Gamunex-C, Cari | Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify | |
|----------------------|--|--|
| Diagnosis | Lambert-Eaton Myasthenic Syndrome (LEMS) | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria 1 - Diagnosis of Lambert-Eaton myasthenic syndrome (LEMS) AND 2 - Medical records documenting BOTH of the following: History and physical examination documenting the severity of the condition, including • frequency and severity of infections where applicable Laboratory results or diagnostic evidence supporting the indication for which immune • alobulin is requested AND **3** - History of failure, contraindication, or intolerance to immunomodulator monotherapy (e.g., azathioprine, corticosteroids) AND 4 - Concomitant immunomodulator therapy (e.g., azathioprine, corticosteroids), unless contraindicated, will be used for long-term management of LEMS AND 5 - Prescribed by or in consultation with a neurologist AND 6 - Intravenous Immunoglobulin (IVIG) dose does not exceed 2,000 milligrams (mg) per kilogram (kg) per month given over 2 to 5 consecutive days. IVIG administration may be repeated monthly as needed to prevent exacerbation. Dosing interval may need to be adjusted in patients with severe comorbidities AND

7 - For long term treatment, documentation of titration to the minimum dose and frequency needed to maintain a sustained clinical effect

AND

8 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to ALL the following products.* (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products

- Flebogamma
- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Gamunex-C
- Hizentra Vial & Syringe
- Privigen

| Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Ga S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamr Gammaplex, Octagam, Panzyga, Gammaked, Xembify | |
|--|--|
| Diagnosis | Lambert-Eaton Myasthenic Syndrome (LEMS) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to immune globulin therapy

AND

2 - Statement of expected frequency and duration of proposed immune globulin treatment

AND

3 - For long term treatment, documentation of titration to the minimum effective dose and frequency needed to maintain a sustained clinical response

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Lennox Gastaut Syndrome |
|-----------------|-------------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - History of failure, contraindication or intolerance to initial treatment with traditional antiepileptic pharmacotherapy (e.g., lamotrigine, phenytoin, valproic acid)

AND

2 - Medical records documenting BOTH of the following:

- History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable
- Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested

AND

3 - Prescribed by or in consultation with a neurologist

AND

4 - Intravenous immunoglobulin (IVIG) dose does not exceed 400 milligrams (mg) per kilogram (kg) per day given for 4 to 5 consecutive days. IVIG administration may be repeated monthly as needed in patients requiring maintenance therapy. Dosing interval may need to be adjusted in patients with severe comorbidities

5 - For long term treatment, documentation of titration to the minimum dose and frequency needed to maintain a sustained clinical effect

AND

6 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to ALL the following products.* (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products)

- Flebogamma
- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Gamunex-C
- Hizentra Vial & Syringe
- Privigen

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| · · · · · · · · · · · · · · · · · · · | |
|---------------------------------------|-------------------------|
| Diagnosis | Lennox Gastaut Syndrome |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to immune globulin therapy

AND

2 - Statement of expected frequency and duration of proposed immune globulin treatment

3 - For long term treatment, documentation of titration to the minimum effective dose and frequency needed to maintain a sustained clinical response

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Multifocal Motor Neuropathy (MMN) |
|-----------------|-----------------------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- 1 Diagnosis of multifocal motor neuropathy as confirmed by ALL of the following:
 - Weakness with slowly progressive or stepwise progressive course over at least one month
 - Asymmetric involvement of two or more nerves
 - Absence of motor neuron signs and bulbar signs

AND

- 2 Medical records documenting BOTH of the following:
 - History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable
 - Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested

AND

3 - Prescribed by or in consultation with a neurologist

4 - Intravenous immunoglobulin (IVIG) dose does not exceed 2,400 milligram (mg) per kilogram (kg) per month given over 2 to 5 consecutive days. IVIG administration may be repeated monthly as needed to prevent exacerbation. Dosing interval may need to be adjusted in patients with severe comorbidities.

AND

5 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to ALL the following products.* (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products)

- Flebogamma
- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Gamunex-C
- Hizentra Vial & Syringe
- Privigen

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Multifocal Motor Neuropathy (MMN) |
|-----------------|-----------------------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to therapy as measured by an objective scale [e.g., Rankin, Modified Rankin, Medical Research Council (MRC) scale]

2 - Prescribed by or in consultation with a neurologist

AND

3 - Intravenous immunoglobulin (IVIG) dose does not exceed 2,400 milligram (mg) per kilogram (kg) per month given over 2 to 5 consecutive days. Dosing interval may need to be adjusted in patients with severe comorbidities

AND

4 - For long term treatment, documentation of titration to the minimum dose and frequency needed to maintain a sustained clinical effect

| Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan |
|--|
| S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, |
| Gammaplex, Octagam, Panzyga, Gammaked, Xembify |

| Diagnosis | Prevention of infection in Multiple Myeloma |
|-----------------|---|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of multiple myeloma

AND

- 2 Medical records documenting BOTH of the following:
 - History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable
 - Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested

3 - ONE of the following:

- Documented hypogammaglobulinemia [immunoglobulin (IgG) less than 500 milligrams (mg) per deciliter (dL)]
- History of bacterial infection(s) associated with multiple myeloma

AND

4 - Intravenous immunoglobulin (IVIG) dose does not exceed 400 mg per kilogram (kg) every 3 to 4 weeks

AND

5 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to ALL the following products. (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products)

- Flebogamma
- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Gamunex-C
- Hizentra Vial & Syringe
- Privigen

| Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan |
|--|
| S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, |
| Gammaplex, Octagam, Panzyga, Gammaked, Xembify |
| |

| Approval Length12 month(s)Therapy StageReauthorizationGuideline TypePrior Authorization | Diagnosis | Prevention of infection in Multiple Myeloma |
|---|-----------------|---|
| | Approval Length | 12 month(s) |
| Guideline Type Prior Authorization | Therapy Stage | Reauthorization |
| | Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to immune globulin therapy

2 - Statement of expected frequency and duration of proposed immune globulin treatment

AND

3 - For long term treatment, documentation of titration to the minimum effective dose and frequency needed to maintain a sustained clinical response

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Relapsing Multiple Sclerosis |
|-----------------|------------------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of relapsing forms of multiple sclerosis (MS) (e.g., relapsing-remitting MS, secondary- progressive MS with relapses, progressive-relapsing MS with relapses)

AND

2 - Medical records documenting BOTH of the following:

- History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable
- Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested

AND

3 - Documentation of an MS exacerbation or progression (worsening) of the patient's clinical status from the visit prior to the one prompting the decision to initiate immune globulin therapy

AND 4 - History of failure, contraindication, or intolerance to at least TWO of the following agents: Aubagio (teriflunomide) • Avonex (interferon beta-1a) • Betaseron (interferon beta-1b) • Copaxone/Glatopa (glatiramer acetate) • Extavia (interferon beta-1b) • Gilenya (fingolimod) • Lemtrada (alemtuzumab) • Mavenclad (cladribine) • Mayzent (siponimod) • Ocrevus (ocrelizumab) • Plegridy (peginterferon beta-1a) • Rebif (interferon beta-1a) • Tecfidera (dimethyl fumarate) • Tysabri (natalizumab) • AND 5 - Prescribed by or in consultation with a neurologist AND 6 - Induction, when indicated, does not exceed a dose of 400 milligrams (mg) per kilogram (kg) daily for up to five days AND 7 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to ALL the following products. (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products) Flebogamma •

- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Gamunex-C

- Hizentra Vial & Syringe
- Privigen

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Relapsing Multiple Sclerosis |
|-----------------|------------------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Medical records, including findings of interval examination including neurological deficits incurred and assessment of disability [e.g., Expanded Disability Status Scale (EDSS), Functional Systems Score (FSS), Multiple Sclerosis Functional Composite (MSFC), Disease Steps (DS)]

AND

2 - Stable or improved disability score (e.g., EDSS, FSS, MSFC, DS)

AND

3 - Documentation of decreased number of relapses since starting immune globulin therapy

AND

4 - Diagnosis continues to be the relapsing forms of multiple sclerosis (MS)

AND

5 - Prescribed by or in consultation with a neurologist

6 - Intravenous immunoglobulin (IVIG) dose does not exceed 1,000 milligram (mg) per kilogram (kg) monthly

AND

7 - For long term treatment, documentation of titration to the minimum dose and frequency needed to maintain a sustained clinical effect

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Myasthenia Gravis - Exacerbation |
|-----------------|----------------------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of generalized myasthenia gravis

AND

2 - Medical records documenting BOTH of the following:

- History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable
- Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested

AND

3 - Evidence of myasthenia exacerbation, defined by at least ONE of the following symptoms in the last month

| Difficulty swallowing Acute respiratory failure Major functional disability responsible for the discontinuation of physical activity Recent immunotherapy treatment with a checkpoint inhibitor [e.g., Keytruda (pembrolizumab), Opdivo (nivolumab), Tecentriq (atezolizumab)] |
|--|
| AND |
| 4 - ONE of the following: |
| History of failure, contraindication, or intolerance to immunomodulator therapy (e.g., azathioprine, mycophenolate mofetil, cyclosporine) for long-term management of myasthenia gravis Currently receiving immunomodulator therapy (e.g., azathioprine, mycophenolate mofetil, cyclosporine) for long-term management of myasthenia gravis |
| AND |
| 5 - Prescribed by or in consultation with a neurologist |
| AND |
| 6 - Intravenous immunoglobulin (IVIG) dose does not exceed 2,000 milligrams (mg) per kilogram (kg) per month given over 2 to 5 days administered in up to three monthly infusions. Dosing interval may need to be adjusted in patients with severe comorbidities. |
| AND |
| 7 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to ALL the following products. (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products) |
| Flebogamma Gammagard Liquid Gammagard S-D Gammaked Gamunex-C Hizentra Vial & Syringe |
| |

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Refractory Myasthenia Gravis |
|-----------------|------------------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of refractory generalized myasthenia gravis by or in consultation with a physician or center with expertise in management of myasthenia gravis

AND

2 - Medical records documenting BOTH of the following:

- History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable
- Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested

AND

3 - Documentation that the disease status is unchanged or worsening (persistent or worsening symptoms that limit functioning) despite failure, contraindication, or intolerance to BOTH of the following (used in adequate doses and duration):

- Corticosteroids
- Two immunomodulator therapies (e.g., azathioprine, mycophenolate mofetil, cyclosporine, methotrexate, tacrolimus)

AND

4 - Currently receiving immunomodulator therapy (e.g., corticosteroids, azathioprine,

mycophenolate mofetil, cyclosporine, methotrexate, tacrolimus), used in adequate doses, for long-term management of myasthenia gravis

AND

5 - Prescribed by or in consultation with a neurologist

AND

6 - Intravenous immunoglobulin (IVIG) dose does not exceed 2,000 milligrams (mg) per kilogram (kg) per month given over 2 to 5 days administered in up to three monthly infusions. Dosing interval may need to be adjusted in patients with severe comorbidities.

AND

7 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to ALL the following products. (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products)

- Flebogamma
- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Gamunex-C
- Hizentra Vial & Syringe
- Privigen

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan
S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif,
Gammaplex, Octagam, Panzyga, Gammaked, XembifyDiagnosisMyasthenia Gravis –Exacerbation and Refractory Myasthenia GravisApproval Length12 month(s)Therapy StageReauthorizationGuideline TypePrior Authorization

Approval Criteria

1 - Documentation of positive clinical response to immune globulin therapy

AND

2 - Statement of expected frequency and duration of proposed immune globulin treatment

AND

3 - For long term treatment, documentation of titration to the minimum effective dose and frequency needed to maintain a sustained clinical response

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Neuromyelitis Optica |
|-----------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, laboratory values, etc.) to support the diagnosis of neuromyelitis optica spectrum disorder (NMOSD) by a neurologist confirming ALL of the following:

1.1 Serologic testing for anti-aquaporin-4 immunoglobulin G (AQP4-IgG) or Neuromyelitis optica immunoglobulin G (NMO-IgG) antibodies has been performed

AND

1.2 ONE of the following:

1.2.1 If AQP4-IgG/NMO-IgG positive, past medical history of ONE of the following:

- Optic neuritis
- Acute myelitis

| Area postrema syndrome: episode of otherwise unexplained hiccups or nausea and vomiting |
|---|
| vomiting Symptomatic narcolepsy or acute diencephalic clinical syndrome with NMOSD-typical |
| diencephalic MRI lesions Symptomatic cerebral syndrome with NMOSD-typical brain lesions |
| |
| OR |
| 1.2.2 If AQP4-IgG/NMO-IgG negative, past medical history of TWO of the following: |
| Optic neuritis |
| Acute myelitis Area postrema syndrome: episode of otherwise unexplained hiccups or nausea and vomiting |
| Acute brainstem syndrome Symptomatic narcolepsy or acute diencephalic clinical syndrome with NMOSD-typical diencephalic MRI lesions |
| Symptomatic cerebral syndrome with NMOSD-typical brain lesions |
| AND |
| |
| 1.3 Diagnosis of multiple sclerosis or other diagnoses have been ruled out |
| AND |
| 2 - Medical records documenting BOTH of the following: |
| History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested |
| AND |
| 3 - History of failure, contraindication, or intolerance to at least TWO of the following: |
| Azathioprine |
| CorticosteroidsMycophenolate mofetil |
| Rituximab |
| |

• Soliris (eculizumab)

AND

4 - Patient is not receiving immune globulin in combination with either of the following:

Rituximab

• Soliris (eculizumab)

AND

5 - Prescribed by or in consultation with a neurologist

AND

6 - Intravenous immunoglobulin (IVIG) dose does not exceed 2,000 milligram (mg) per kilogram (kg) per month given over 2 to 5 days administered in up to six monthly infusions. Dosing interval may need to be adjusted in patients with severe comorbidities.

AND

7 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to ALL the following products. (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products)

- Flebogamma
- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Gamunex-C
- Hizentra Vial & Syringe
- Privigen

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis Neuromyelitis Optica |
|--------------------------------|
|--------------------------------|

| Approval Length | 12 month(s) | |
|---|--|--|
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Patient has previously been treated with immune globulin | | |
| | AND | |
| 2 - Submission of medical records (e.g., chart notes, laboratory tests) to demonstrate a positive clinical response from baseline as demonstrated by BOTH of the following: | | |
| | 2.1 Reduction in the number and or severity of relapses or signs and symptoms of neuromyelitis optica spectrum disorder (NMOSD) | |
| | AND | |
| 2.2 Maintenance, reduction, or discontinuation of dose(s) of any baseline immunosuppressive therapy (IST) prior to starting immune globulin. (NOTE: Add on, dose escalation of IST, or additional rescue therapy from baseline to treat NMOSD or exacerbation of symptoms while on immune globulin therapy will be considered as treatment failure.) | | |
| | AND | |
| 3 - Patient is not rec | eiving immune globulin in combination with either of the following: | |
| RituximabSoliris (eculizumab) | | |
| | AND | |
| 4 - Prescribed by or in consultation with a neurologist | | |
| | AND | |

5 - Intravenous immunoglobulin (IVIG) dose does not exceed 2,000 milligrams (mg) per kilogram (kg) per month given over 2 to 5 days administered in up to six monthly infusions. Dosing interval may need to be adjusted in patients with severe comorbidities

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Posttransfusion Purpura |
|-----------------|-------------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of posttransfusion purpura

AND

2 - Medical records documenting BOTH of the following:

- History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable
- Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested

AND

3 - Intravenous immunoglobulin (IVIG) dose does not exceed 1,000 milligrams (mg) per kilogram (kg) for 2 days

AND

4 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to ALL the following products. (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products)

- Flebogamma
- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Gamunex-C
- Hizentra Vial & Syringe
- Privigen

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Posttransfusion Purpura |
|-----------------|-------------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to immune globulin therapy

AND

2 - Statement of expected frequency and duration of proposed immune globulin treatment

AND

3 - For long term treatment, documentation of titration to the minimum effective dose and frequency needed to maintain a sustained clinical response

| Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify | |
|--|--------------------------------|
| Diagnosis | Post B-Cell Targeted Therapies |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |

| Guideline Type | Prior Authorization | |
|---|--|--|
| | | |
| Approval Criteria | | |
| 1 - Documentation confirming previous treatment of B-cell targeted therapy within the last 100 days [e.g., CAR-T (e.g., Kymriah), Rituxan (rituximab), Besponsa (inotuzumab ozogamicin)] | | |
| AND | | |
| 2 - Medical records documenting BOTH of the following: | | |
| History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested | | |
| | AND | |
| 3 - BOTH of the following: | | |
| Documented hypogammaglobulinemia [immunoglobulin (IgG) less than 500 milligrams (mg) per deciliter (dL)] History of bacterial infection(s) associated with B-cell depletion | | |
| | AND | |
| 4 - Intravenous immunoglobulin (IVIG) dose does not exceed 400 mg per kilogram (kg) every 4 weeks, up to 360 days after discontinuation of B-cell depleting therapy | | |
| AND | | |
| contraindication or into are fewer than three pr | 5 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to ALL the following products. (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products) | |
| Flebogamma Gammagard Liquid Gammagard S-D | | |

- Gammaked
- Gamunex-C
- Hizentra Vial & Syringe
- Privigen

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Post B-Cell Targeted Therapies |
|-----------------|--------------------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to immune globulin therapy

AND

2 - Statement of expected frequency and duration of proposed immune globulin treatment

AND

3 - For long term treatment, documentation of titration to the minimum effective dose and frequency needed to maintain a sustained clinical response

| S-D, Gamunex-C, Cari | roduct Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamasta -D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, ammaplex, Octagam, Panzyga, Gammaked, Xembify | |
|----------------------|--|--|
| Diagnosis | Primary Immunodeficiency Syndromes | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |

Approval Criteria

1 - Diagnosis of primary immunodeficiency

AND

2 - Medical records documenting BOTH of the following:

- History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable
- Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested

AND

3 - Clinically significant functional deficiency of humoral immunity as evidenced by ONE of the following:

- Documented failure to produce antibodies to specific antigens
- History of significant recurrent infections

AND

4 - Initial intravenous immunoglobulin (IVIG) dose is 200 to 800 milligrams (mg) per kilogram (kg) every 3 to 4 weeks, based on product prescribing information, and titrated based upon patient response (For subcutaneous immune globulin (SCIG) products, FDA-labeled dosing and conversion guidelines will used to determine benefit coverage.)

AND

5 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to ALL the following products. (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products)

- Flebogamma
- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Gamunex-C
- Hizentra Vial & Syringe

Privigen

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Primary Immunodeficiency Syndromes |
|-----------------|------------------------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to immune globulin therapy

AND

2 - Statement of expected frequency and duration of proposed immune globulin treatment

AND

3 - For long term treatment, documentation of titration to the minimum effective dose and frequency needed to maintain a sustained clinical response

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Rasmussen Syndrome |
|-----------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of ONE of the following demonstrating that:

- Short term amelioration of encephalitis is needed prior to definitive surgical therapy
- Disease symptoms (e.g., seizures) persist despite surgical treatment
- The patient is not a candidate for surgical treatment

2 - Medical records documenting BOTH of the following:

- History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable
- Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested

AND

3 - Intravenous immunoglobulin (IVIG) dose does not exceed 2,000 milligrams (mg) per kilogram (kg) per month given over 2 to 5 days

AND

4 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to ALL the following products.* (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products)

- Flebogamma
- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Gamunex-C
- Hizentra Vial & Syringe
- Privigen

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan
S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif,
Gammaplex, Octagam, Panzyga, Gammaked, XembifyDiagnosisRasmussen SyndromeApproval Length12 month(s)

| Therapy Stage | Reauthorization |
|----------------------|---|
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Documentation of | of positive clinical response to immune globulin therapy |
| | |
| | AND |
| 2 - Statement of exp | pected frequency and duration of proposed immune globulin treatment |
| | AND |

3 - For long term treatment, documentation of titration to the minimum effective dose and frequency needed to maintain a sustained clinical response

| Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan |
|--|
| S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, |
| Gammaplex, Octagam, Panzyga, Gammaked, Xembify |

| Diagnosis | Stiff-Person Syndrome |
|-----------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of stiff-person syndrome

AND

- 2 Medical records documenting BOTH of the following:
 - History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable

 Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested

AND

3 - History of failure, contraindication or intolerance to GABAergic (gamma-aminobutyric acid analogs) medication (e.g., baclofen, benzodiazepines)

AND

4 - Prescribed by or in consultation with a neurologist

AND

5 - Intravenous immunoglobulin (IVIG) dose does not exceed 2,000 milligrams (mg) per kilogram (kg) per month given over 2 to 5 days. IVIG administration may be repeated monthly as needed for patients requiring maintenance therapy. Dosing interval may need to be adjusted in patients with severe comorbidities

AND

6 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to ALL the following products. (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products)

- Flebogamma
- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Gamunex-C
- Hizentra Vial & Syringe
- Privigen

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Stiff-Person Syndrome |
|-----------|-----------------------|
|-----------|-----------------------|

| Approval Length | 12 month(s) |
|--|---|
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Documentation of a | a positive clinical improvement from baseline |
| | AND |
| 2 - Prescribed by or in | consultation with a neurologist |
| | AND |
| kilogram (kg) per mont as needed for patients | oglobulin (IVIG) dose does not exceed 2,000 milligrams (mg) per th given over 2 to 5 days. IVIG administration may be repeated monthly requiring maintenance therapy. Dosing interval may need to be th severe comorbidities |
| | AND |
| • | ment, documentation of titration to the minimum dose and frequency |

| needed to maintain a sustained clinical effect | | |
|--|--|--|
| | | |
| Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Fleboga | | |

| Gammaplex, Octagam, Panzyga, Gammaked, Xembify | |
|--|---|
| Diagnosis | Thrombocytopenia, secondary to Hepatitis C Virus (HCV), Human Immunodeficiency Virus (HIV), or pregnancy |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - One of the following:

| 1.1 Both of the following: | | |
|---|--|--|
| Diagnosis of thrombocytopenia secondary to Hepatitis C Virus (HCV) infection Patient is receiving concurrent antiviral therapy, unless contraindicated | | |
| OR | | |
| 1.2 Both of the following: | | |
| Diagnosis of thrombocytopenia secondary Human Immunodeficiency Virus (HIV) infection Patient is receiving concurrent antiviral therapy, unless contraindicated | | |
| OR | | |
| 1.3 Diagnosis of thrombocytopenia secondary to pregnancy | | |
| AND | | |
| 2 - Medical records documenting BOTH of the following: | | |
| History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested | | |
| AND | | |
| 3 - Documented platelet count less than 50 x 10^9 per liter (L) (obtained within the past 30 days) | | |
| AND | | |
| 4 - Intravenous immunoglobulin (IVIG) dose does not exceed 1,000 milligrams (mg) per kilogram (kg) per day for 1 to 2 days | | |
| | | |

5 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to ALL the following products. (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products)

- Flebogamma
- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Gamunex-C
- Hizentra Vial & Syringe
- Privigen

Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, Gammaplex, Octagam, Panzyga, Gammaked, Xembify

| Diagnosis | Thrombocytopenia, secondary to Hepatitis C Virus (HCV), Human Immunodeficiency Virus (HIV), or pregnancy |
|-----------------|---|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - One of the following:

1.1 Both of the following:

- Diagnosis of thrombocytopenia secondary to Hepatitis C Virus (HCV) infection
- Patient is receiving concurrent antiviral therapy, unless contraindicated

OR

1.2 Both of the following:

- Diagnosis of thrombocytopenia secondary Human Immunodeficiency Virus (HIV) infection
- Patient is receiving concurrent antiviral therapy, unless contraindicated

OR

1.3 Diagnosis of thrombocytopenia secondary to pregnancy

AND

2 - Intravenous immunoglobulin (IVIG) dose does not exceed 2,000 milligram (mg) per kilogram (kg) per month given over 2 to 5 consecutive days. IVIG administration may be repeated monthly as needed to prevent exacerbation. Dosing interval should be adjusted depending upon response and titrated to the minimum effective dose that can be given at maximum intervals to maintain safe platelet levels.

| Product Name: HyQvia, Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan |
|--|
| S-D, Gamunex-C, Carimune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, |
| Gammaplex, Octagam, Panzyga, Gammaked, Xembify |

| Diagnosis | All other indications |
|-----------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- **1** One of the following diagnoses:
 - Autoimmune Uveitis
 - Cytomegalovirus (CMV) induced pneumonitis in solid organ transplants
 - Enteroviral Meningoencephalitis
 - IgM antimyelin-associated glycoprotein paraprotein-associated peripheral neuropathy
 - Lymphoproliferative disease (treatment of bacterial infections)
 - Monoclonal gammopathy
 - Paraproteinemic neuropathy
 - Renal transplantation (prevention or treatment of acute humoral rejection)
 - Severe Rheumatoid arthritis
 - Rotaviral enterocolitis
 - Staphylococcal toxic shock

- Toxic epidermal necrolysis or Stevens-Johnson syndrome
- Urticaria (delayed pressure)

- 2 Medical records documenting BOTH of the following:
 - History and physical examination documenting the severity of the condition, including frequency and severity of infections where applicable
 - Laboratory results or diagnostic evidence supporting the indication for which immune globulin is requested

AND

3 - If the request is for a non-preferred product, there must be a history of failure, contraindication or intolerance to ALL the following products. (Note: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products)

- Flebogamma
- Gammagard Liquid
- Gammagard S-D
- Gammaked
- Gamunex-C
- Hizentra Vial & Syringe
- Privigen

| S-D, Gamunex-C, Cari | , Gammagard S-D, Gammagard liquid, Cuvitru, Gamastan, Gamastan mune NF Nanofiltered, Privigen, Hizentra, Bivigam, Flebogamma Dif, , Panzyga, Gammaked, Xembify |
|---------------------------------|--|
| Diagnosis All other indications | |

| Diagnosis | All other indications |
|-----------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to immune globulin therapy

2 - Statement of expected frequency and duration of proposed immune globulin treatment

AND

3 - For long term treatment, documentation of titration to the minimum effective dose and frequency needed to maintain a sustained clinical response

2. Revision History

| Date | Notes |
|-----------|-------------------------------------|
| 6/28/2021 | Arizona Medicaid 7.1 Implementation |

Inbrija

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99466 Inbrija

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Inbrija | |
|--------------------------------------|-----------------------|
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of Parkinson's disease | |

2 - Inbrija will be used as intermittent treatment for OFF episodes

AND

3 - Prescribed by, or in consultation with, a neurologist or specialist in the treatment of Parkinson's disease

AND

4 - Patient is currently on a stable dose of a carbidopa/levodopa-containing medication and will continue receiving treatment with a carbidopa/levodopa-containing medication while on therapy

AND

5 - Patient continues to experience greater than or equal to 2 hours of OFF time per day despite optimal management of carbidopa/levodopa therapy including BOTH of the following:

- Taking carbidopa/levodopa on an empty stomach or at least one half-hour or more before or one hour after a meal or avoidance of high protein diet
- Dose and dosing interval optimization

AND

6 - History of failure, contraindication, or intolerance to TWO anti-Parkinson's disease therapies from the following adjunctive pharmacotherapy classes (trial must be from two different classes):

- Dopamine agonists (e.g., pramipexole, ropinirole)
- Catechol-O-methyl transferase (COMT) inhibitors (e.g., entacapone)
- Monoamine oxidase (MAO) B inhibitors (e.g., selegiline)

Product Name: Inbrija

| Approval Length | 12 month(s) |
|-----------------|---------------------|
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Inbrija therapy

AND

2 - Patient will continue to receive treatment with a carbidopa/levodopa-containing medication

2. Revision History

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk copy C&S Arizona standard to Medicaid Arizona |

Ingrezza (valbenazine)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-114379 | Ingrezza (valbenazine) |
|-----------|------------------------|
|-----------|------------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Ingrezza | |
|------------------------|---------------------------------------|
| Diagnosis | Moderate to Severe Tardive Dyskinesia |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of moderate to severe tardive dyskinesia (TD) secondary to a centrally acting dopamine receptor blocking agent (DRBA)

AND 2 - Prescribed by or in consultation with a psychiatrist or neurologist AND 3 - Patient is 18 years of age or older AND 4 - Patient has an Abnormal Involuntary Movement Scale (AIMS) score of 3 or 4 on any one of the AIMS items 1 through 9 AND 5 - Ingrezza is not prescribed concurrently with Austedo® or tetrabenazine AND

| Product Name: Ingrezza | |
|------------------------|---------------------------------------|
| Diagnosis | Moderate to Severe Tardive Dyskinesia |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

6 - Dose does not exceed 80 mg (1 capsule) per day

Approval Criteria

1 - Patient is responding positively to therapy as evidenced by a reduction in the baseline AIMS score in any one of the AIMS items 1 through 9

2 - Ingrezza is not prescribed concurrently with Austedo or tetrabenazine

AND

3 - Dose does not exceed 80 mg (1 capsule) per day.

2. Revision History

| Date | Notes |
|-----------|------------------------------------|
| 9/22/2022 | Removed step through tetrabenazine |

Inhaled Corticosteroids - AZM

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-105180 Inhaled Corticosteroids - AZM

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: |
|-----------------|
|-----------------|

1. Criteria

| Product Name: Alvesco, Arnuity Ellipta, Asmanex HFA, Qvar Redihaler | | | |
|---|---------------------|--|--|
| Approval Length | 12 month(s) | | |
| Guideline Type | Prior Authorization | | |
| | | | |
| Approval Criteria | Approval Criteria | | |
| 1 - Diagnosis of asthma | | | |
| | | | |
| AND | | | |
| | | | |

2 - History of failure, contraindication, intolerance to a majority (not more than 3) of the preferred inhaled corticosteroids:

- Asmanex Twisthaler (mometasone) •
- Flovent Diskus (fluticasone)Flovent HFA (fluticasone)
- Pulmicort Flexhaler (budesonide) •
- budesonide respule (generic) •

2. Revision History

| Date | Notes |
|-----------|--|
| 3/24/2022 | Removed Pulmicort and budesonide respules as targets |

Injectable Oncology Agents

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-113514 Injectable Oncology Agents

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 9/8/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

Product Name: Non-Preferred Injectable Oncology Drugs: Brand Alimta, generic pemetrexed, Brand Bicnu, generic carmustine, Brand Bortezomib, Breyanzi, Brand Carmustine, Carvykti, Enhertu, Brand Hycamtin injection, generic topotecan injection, Kymriah, Kyprolis, Opdivo, Brand Pemetrexed, Brand Pemfexy, Synribo, Brand Velcade, generic bortezomib, Brand Vidaza, generic azacitidine, Yervoy, Yescarta

| Diagnosis | Cancer Indications |
|-------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - The drug is being used as indicated by National Comprehensive Cancer Network (NCCN) guidelines with a Category of Evidence and Consensus of 1, 2A, or 2B

2. Revision History

| Date | Notes |
|----------|------------------------------------|
| 9/8/2022 | Updated with additional NP targets |

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-113514 Injectable Oncology Agents

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 9/8/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

Product Name: Non-Preferred Injectable Oncology Drugs: Brand Alimta, generic pemetrexed, Brand Bicnu, generic carmustine, Brand Bortezomib, Breyanzi, Brand Carmustine, Carvykti, Enhertu, Brand Hycamtin injection, generic topotecan injection, Kymriah, Kyprolis, Opdivo, Brand Pemetrexed, Brand Pemfexy, Synribo, Brand Velcade, generic bortezomib, Brand Vidaza, generic azacitidine, Yervoy, Yescarta

| Diagnosis | Cancer Indications |
|-----------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - The drug is being used as indicated by National Comprehensive Cancer Network (NCCN) guidelines with a Category of Evidence and Consensus of 1, 2A, or 2B

2. Revision History

| Date | Notes |
|----------|------------------------------------|
| 9/8/2022 | Updated with additional NP targets |

Inlyta

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99682 Inlyta

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|--------------------|-----------|
| P&T Approval Date: | |
| P&T Revision Date: | |

1. Criteria

| Product Name: Inlyta | |
|----------------------|-------------------------------|
| Diagnosis | Advanced Renal Cell Carcinoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

| 1 - Diagnosis of renal cell cancer |
|------------------------------------|
| |
| AND |
| |
| 2 - One of the following: |
| 2.1 Disease has relapsed |
| |
| OR |
| |
| 2.2 Diagnosis of Stage IV disease |

| Product Name: Inlyta | | |
|---|-----------------------|--|
| Diagnosis | Thyroid Carcinoma | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| Approval Criteria 1 - ONE of the following diagnosis: Follicular Carcinoma Hürthle Cell Carcinoma Papillary Carcinoma | | |
| | AND | |
| 2 - ONE of the following: Unresectable recurrent Persistent locoregional disease Metastatic disease | | |

3 - Disease is refractory to radioactive iodine treatment

| Product Name: Inlyta | |
|----------------------|--|
| Diagnosis | Advanced Renal Cell Carcinoma, Thyroid Carcinoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Inlyta therapy

| Product Name: Inlyta | |
|----------------------|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Inlyta will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Inlyta | |
|----------------------|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Inlyta therapy

| Date | Notes |
|----------|--------------------|
| 4/8/2021 | 7/1 Implementation |

Insulins, Concentrated- Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99580 | Insulins, Concentrated- Arizona |
|----------|---------------------------------|
|----------|---------------------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Humulin R U-500 vial and kwikpen | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - History of failure, intolerance, or contraindication to ALL of the following:

- Novolog or Humalog
- Lantus
- Levemir

| OR | |
|--|---|
| 2 - There is a reason or special circumstance the patient needs to use a concentrated insuling product | n |

2. Revision History

Г

| Date | Notes |
|----------|------------------|
| 8/4/2021 | Update guideline |

Iressa

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99683 Iressa

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|--------------------|-----------|
| P&T Approval Date: | |
| P&T Revision Date: | |

1. Criteria

| Product Name: Iressa | |
|----------------------|------------------------------------|
| Diagnosis | Non-Small Cell Lung Cancer (NSCLC) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of metastatic or recurrent non-small cell lung cancer (NSCLC)

AND

- **2** ONE of the following:
 - Tumors are positive for epidermal growth factor receptor (EGFR) exon 19 deletions
 - Tumors are positive for exon 21 (L858R) substitution mutations
 - Tumors are positive for a known sensitizing EGFR mutation (e.g., in-frame exon 20 insertions, exon 18 G719 mutation, exon 21 L861Q mutation)

| Product Name: Iressa | |
|----------------------|------------------------------------|
| Diagnosis | Non-Small Cell Lung Cancer (NSCLC) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Iressa therapy

| Product Name: Iressa | |
|----------------------|--------------------------------------|
| Diagnosis | Central Nervous System (CNS) Cancers |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of central nervous system (CNS) cancer with metastatic lesions

AND

2 - Iressa is active against primary (NSCLC) tumor with a known epidermal growth factor receptor (EGFR) sensitizing mutation

| Product Name: Iressa | |
|--------------------------------------|--|
| Central Nervous System (CNS) Cancers | |
| 12 month(s) | |
| Reauthorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Iressa therapy

| Product Name: Iressa | |
|----------------------|---|
| Diagnosis | National Comprehensive Cancer Network (NCCN) Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Iressa will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Iressa | |
|----------------------|---|
| Diagnosis | National Comprehensive Cancer Network (NCCN) Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| | |

1 - Documentation of positive clinical response to Iressa therapy

| Date | Notes |
|----------|--------------------|
| 4/8/2021 | 7/1 Implementation |

Iron Chelators

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-104870 Iron Chelators

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 3/17/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Exjade, Brand Jadenu, generic deferasirox | |
|---|--|
| Diagnosis | Chronic Iron Overload due to Blood Transfusion |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of chronic iron overload (e.g., sickle cell anemia, thalassemia, etc.) due to blood transfusion

| Product Name: Brand Exjade, Brand Jadenu, generic deferasirox | |
|---|--|
| Diagnosis | Chronic Iron Overload due to Blood Transfusion |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| | 1 |

1 - Documentation of positive clinical response to therapy

| Product Name: Brand Ferriprox, generic deferiprone | |
|--|--|
| Diagnosis | Chronic Iron Overload due to Blood Transfusion |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - BOTH of the following

1.1 Diagnosis of transfusional iron overload due to thalassemia syndromes

AND

1.2 Current chelation therapy is inadequate [e.g., Desferal (deferoxamine), Exjade (deferasirox)]

| Product Name: Brand Ferriprox, generic deferiprone | |
|--|--|
| Diagnosis | Chronic Iron Overload due to Blood Transfusion |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to therapy

| Product Name: Brand Exjade, Brand Jadenu, generic deferasirox | |
|---|--|
| Diagnosis | Chronic Iron Overload in Non-Transfusion Dependent Thalassemia Syndrome |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - ALL of the following:

1.1 Diagnosis of chronic iron overload in non-transfusion dependent thalassemia syndrome

AND

1.2 Patient has liver iron (Fe) concentration (LIC) levels consistently greater than or equal to 5 mg Fe per gram of dry weight prior to initiation of treatment with Exjade or Jadenu

AND

1.3 Patient has serum ferritin levels consistently greater than 300 micrograms per liter prior to initiation of treatment with Exjade or Jadenu

| Product Name: Brand Exjade, Brand Jadenu, generic deferasirox | |
|---|--|
| Diagnosis | Chronic Iron Overload in Non-Transfusion Dependent Thalassemia Syndrome |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Documentation of positive clinical response to therapy

| Date | Notes |
|-----------|------------------------------------|
| 3/16/2022 | Added new generic deferiprone tabs |

Irritable Bowel Syndrome-Diarrhea

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99468 | Irritable Bowel Syndrome-Diarrhea |
|----------|-----------------------------------|
|----------|-----------------------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Lotronex, generic alosetron | |
|---|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of severe diarrhea-predominant irritable bowel syndrome (IBS)

| | AND |
|--|-----------------------------------|
| 2 - Symptoms for at least 6 months | |
| | AND |
| 3 - Patient was female at birth | |
| | AND |
| 4 - Age greater than or equal to 18 years | |
| | AND |
| 5 - History of failure, contraindication, or int | olerance to TWO of the following: |
| Antispasmodic agent (e.g. dicyclom Antidiarrheal agents (e.g. loperamid | |

• Tricyclic antidepressant (e.g. amitriptyline)

| Product Name: Brand Lotronex, generic alosetron | |
|---|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | · |

Approval Criteria

1 - Documentation of positive clinical response to Lotronex therapy

| Product Name: Viberzi | |
|-----------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |

| Guideline Type | Prior Authorization |
|---|--|
| | |
| Approval Criteria | |
| 1 - Diagnosis of irritable bowel syndrome with diarrhea (IBS-D) | |
| | |
| | AND |
| 2 History of failure of | $\mathbf{T}_{\mathbf{A}}$ |
| | ontraindication, or intolerance to TWO of the following: |

| Product Name: Viberzi | | |
|-----------------------|---------------------|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |
| | | |
| | | |

1 - Documentation of positive clinical response to Viberzi therapy

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk copy C&S Arizona standard to Medicaid Arizona |

Isotretinoin

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99562 Isotretinoin

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Absorica, Absorica LD, generic Amnesteem, generic Claravis, generic isotretinoin, generic Myorisan, generic Zenatane | | | |
|--|--|--|--|
| Diagnosis Oncology Uses (Off Label) | | | |
| Approval Length 12 month(s) | | | |
| Guideline Type Prior Authorization | | | |

Approval Criteria

1 - Used for oncology indication meeting National Comprehensive Cancer Network (NCCN) with a Category of Evidence and Consensus of 1, 2A, or 2B. or from ONE of the following appropriate compendia of current literature: American Hospital Formulary Service Drug Information, Thomson Micromedex DrugDex, or Clinical Pharmacology

| Product Name: Absorica, Absorica LD, generic Amnesteem, generic Claravis, generic isotretinoin, generic Myorisan, generic Zenatane | | |
|--|-----------------------|--|
| Approval Length 5 month(s) | | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

1 - ONE of the following:

1.1 Diagnosis of severe recalcitrant nodular acne unresponsive to conventional therapy

OR

1.2 Diagnosis of treatment resistant acne

AND

2 - History of failure, contraindication, or intolerance to an adequate trial on TWO of the following conventional therapy regimens:

- Topical retinoid or retinoid-like agent [eg,Retin-A/Retin-A Micro (tretinoin)]
- Oral antibiotic [eg, Ery-Tab (erythromycin), Biaxin (clarithromycin), Minocin (minocycline)]
- Topical antibiotic with or without benzoyl peroxide [eg, Cleocin-T (clindamycin), erythromycin, BenzaClin (benzoyl peroxide/clindamycin), Benzamycin (benzoyl peroxide/erythromycin)]

AND

3 - If the request is for a non-preferred medication, there must be a reason or special circumstance that the patient must be treated with a non-preferred medication (see table in Background section)

Product Name: Absorica, Absorica LD, generic Amnesteem, generic Claravis, generic isotretinoin, generic Myorisan, generic Zenatane

| Diagnosis | Persistent or Recurring Acne After 2 Months Off Therapy |
|-----------------|---|
| Approval Length | 5 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - After greater than or equal to 2 months OFF therapy, persistent or recurring severe recalcitrant nodular acne is still present

| Notes | Authorization will be given only by clinical pharmacist review for up to |
|-------|--|
| | 5 months. |

| Product Name: Absorica, Absorica LD, generic Amnesteem, generic Claravis, generic |
|---|
| isotretinoin, generic Myorisan, generic Zenatane |

| Diagnosis | Dose Titration |
|-----------------|---------------------|
| Approval Length | 1 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Confirmation that the cumulative dose is less than 150 mg/kg (there is little therapeutic benefit to be gained by increasing the cumulative dose beyond 150 mg/kg)*

| Authorization will be given only by clinical pharmacist review for 1 mo nth to allow for titration up to the target dose *See background section |
|---|
| for dosing regimens |

2. Background

| Benefit/Coverage/Program Information |
|--------------------------------------|
| Formulary |
| Preferred Agents: |

Myorisan (isotretinoin), Claravis (isotretinoin), Amnesteem (isotretinoin), Zenatane (isotretinoin), generic isotretinoin

Non-Preferred Agents:

Absorica (isotretinoin)

Absorica LD (isotretinoin)

Dosing by Body Weight (based on administration with food):

| Body | Weigh | ntDaily Dose | Daily Dose | | |
|------|-------|---------------|-------------|-------------|--|
| Kg | Lbs | 0.5 mg/kg/day | 1 mg/kg/day | 2 mg/kg/day | |
| 40 | 88 | 20 | 40 | 80 | |
| 50 | 110 | 25 | 50 | 100 | |
| 60 | 132 | 30 | 60 | 120 | |
| 70 | 154 | 35 | 70 | 140 | |
| 80 | 176 | 40 | 80 | 160 | |
| 90 | 198 | 45 | 90 | 180 | |
| 100 | 220 | 50 | 100 | 200 | |

| Date | Notes |
|-----------|-----------------|
| 6/25/2021 | Updated program |

Isturisa

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99684 Isturisa

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|--------------------|-----------|
| P&T Approval Date: | |
| P&T Revision Date: | |

1. Criteria

| Product Name: Isturisa | |
|------------------------|-----------------------|
| Diagnosis | Cushing's Disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

- **1** Both of the following:
- **1.1** Diagnosis of Cushing's disease

AND

1.2 ONE of the following:

- Patient is not a candidate for pituitary surgery
- Pituitary surgery has not been curative

| Cushing's Disease |
|---------------------|
| 12 month(s) |
| Reauthorization |
| Prior Authorization |
| |
| |

Approval Criteria

1 - Documentation of positive response to Isturisa therapy

| Product Name: Isturisa | |
|------------------------|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Isturisa will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

Product Name: Isturisa

| Diagnosis | NCCN Recommended Regimens |
|-----------------|---------------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| | |

1 - Documentation of positive clinical response to Isturisa therapy

| Date | Notes |
|----------|--------------------|
| 4/8/2021 | 7/1 Implementation |

Jakafi

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99749 Jakafi

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Jakafi | |
|----------------------|-----------------------|
| Diagnosis | Myelofibrosis |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - ONE of the following diagnoses:

1.1 Primary myelofibrosis

OR 1.2 Post-polycythemia vera myelofibrosis OR

1.3 Post-essential thrombocythemia myelofibrosis

| Product Name: Jakafi | |
|----------------------|---------------------|
| Diagnosis | Myelofibrosis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation that patient has evidence of symptom improvement or reduction in spleen volume while on Jakafi*

| C V | * NOTE: If documentation does not provide evidence of symptom impr ovement or reduction in spleen volume while on Jakafi, authorization will be issued for 2 months to allow for dose titration with discontinuati on of therapy. |
|--------|--|
|--------|--|

| Product Name: Jakafi | |
|----------------------|-----------------------|
| Diagnosis | Polycythemia Vera |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of polycythemia vera

AND

2 - History of failure, inadequate response, contraindication, or intolerance to ONE of the following:

2.1 Hydroxyurea

OR

2.2 Interferon therapy (e.g., Intron A, Pegasys, PegIntron)

| Product Name: Jakafi | |
|----------------------|---------------------|
| Diagnosis | Polycythemia Vera |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation that patient has evidence of symptom improvement or reduction in spleen volume while on Jakafi*

| Notes | * NOTE: If documentation does not provide evidence of symptom impr ovement or reduction in spleen volume while on Jakafi, authorization |
|-------|--|
| | will be issued for 2 months to allow for dose titration with discontinuati on of therapy. |

| Product Name: Jakafi | |
|----------------------|----------------------------------|
| Diagnosis | Graft versus host disease (GVHD) |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of graft versus host disease (GVHD)

AND

2 - Disease is steroid refractory

| Product Name: Jakafi | |
|----------------------|----------------------------------|
| Diagnosis | Graft versus host disease (GVHD) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation that patient has symptom improvement while on Jakafi

| Product Name: Jakafi | |
|----------------------|---|
| Diagnosis | National Comprehensive Cancer Network (NCCN) Recommended Regimens |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Jakafi will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Jakafi | |
|----------------------|---|
| Diagnosis | National Comprehensive Cancer Network (NCCN) Recommended Regimens |
| Approval Length | 12 month(s) |

| Therapy Stage | Reauthorization |
|---|---------------------|
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Documentation of positive clinical response to Jakafi therapy | |

| Date | Notes |
|----------|-------------------------------------|
| 6/2/2021 | Arizona Medicaid 7.1 Implementation |

Juxtapid - AZ

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99791 Juxtapid - AZ

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Juxtapid | |
|------------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of homozygous familial hypercholesterolemia (HoFH) as confirmed by BOTH of the following:*

1.1 ONE of the following:

| Pre-treatment low density lipoprotein cholesterol (LDL-C) greater than 500 milligrams per deciliter Treated LDL-C greater than 300 milligrams per deciliter |
|--|
| AND |
| 1.2 ONE of the following: |
| Xanthoma before 10 years of age Evidence of heterozygous familial hypercholesterolemia (HeFH) in both parents |
| AND |
| 2 - Used as an adjunct to a low-fat diet and exercise |
| AND |
| 3 - Patient is receiving other lipid-lowering therapy (e.g., statin, ezetimibe, LDL apheresis) |
| AND |
| 4 - Prescribed by ONE of the following: |
| Cardiologist Endocrinologist Lipid specialist |
| AND |
| 5 - Patient has tried, failed or intolerant to Repatha and Praluent |
| AND |
| 6 - Not used in combination with a proprotein convertase subtilisin/kexin type 9 (PCSK9) inhibitor |

| Results of prior genetic testing can be submitted as confirmation of di agnosis of HoFH. |
|--|
| |

| Product Name: Juxtapi | Product Name: Juxtapid | | |
|---|-------------------------------------|--|--|
| Approval Length | 12 month(s) | | |
| Therapy Stage | Reauthorization | | |
| Guideline Type | Prior Authorization | | |
| Approval Criteria 1 - Patient is continuing | a low-fat diet and exercise regimen | | |
| AND | | | |
| 2 - Patient continues to receive other lipid-lowering therapy (e.g., statin, low density lipoprotein [LDL] apheresis) | | | |
| | AND | | |
| 3 - Submission of medical records (e.g. chart notes, laboratory values) documenting low density lipoprotein cholesterol (LDL-C) reduction while on Juxtapid therapy | | | |
| | AND | | |
| 4 - Prescribed by ONE | of the following: | | |
| CardiologistEndocrinologistLipid specialist | | | |
| AND | | | |
| 5 - Not used in combination with a proprotein convertase subtilisin/kexin type 9 (PCSK9) inhibitor | | | |

| Date | Notes |
|-----------|-------------------------------------|
| 7/13/2021 | Arizona Medicaid 7.1 Implementation |

Jynarque

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-100644 Jynarque

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|--------------------|-----------|
| P&T Approval Date: | |
| P&T Revision Date: | |

1. Criteria

| Product Name: Jynarque, Jynarque Pak | | |
|--------------------------------------|-----------------------|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |

1 - Diagnosis of autosomal dominant polycystic kidney disease (ADPKD)

| Product Name: Jynarque, Jynarque Pak | | |
|--------------------------------------|---------------------|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |

1 - Documentation of positive clinical response to Jynarque therapy

| Date | Notes |
|------------|-------------------------|
| 12/16/2021 | Added new Jynarque GPIs |

Kalydeco

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99620 Kalydeco

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Kalydeco, Kalydeco packet | | |
|---|-----------------------|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of cystic fibrosis (CF) | | |

2 - Submission of laboratory results confirming that patient has ONE of the mutations in the cystic fibrosis transmembrane conductance regulator (CFTR) gene listed in the table in the Background section:

AND

3 - Prescribed by, or in consultation with, a specialist affiliated with a CF care center

| Product Name: Kalydeco, Kalydeco packet | | |
|---|---------------------|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |
| | | |

Approval Criteria

1 - Provider attests that the patient has achieved a clinically meaningful response while on Kalydeco therapy to ONE of the following:

- Lung function as demonstrated by percent predicted expiratory volume in 1 second (ppFEV1)
- Body mass index (BMI)
- Pulmonary exacerbations
- Quality of life as demonstrated by Cystic Fibrosis Questionnaire-Revised (CFQ-R) respiratory domain score

AND

2 - Prescribed by, or in consultation with, specialist affiliated with a cystic fibrosis (CF) care center

2. Background

Benefit/Coverage/Program Information

CFTR Gene Mutations

| CFTR Gene Mutations | | | | | |
|---------------------|--------|--------|--------|----------|---------------------|
| A1067T | E56K | G1244E | R1070Q | S1251N | 2789+5G→A |
| A455E | E193K | G1349D | R1070W | S1255P | 3272 - 26A→G |
| D110E | E831X | G178R | R117C | S549N | 3849+10kbC→T |
| D110H | F1052V | G551S | R117H | S549R | |
| D1152H | F1074L | K1060T | R347H | S945L | |
| D1270N | G1069R | L206W | R352Q | S977F | |
| D579G | G551D | P67L | R74W | 711+3A→G | |

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Kerendia (finerenone)

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-102888 | Kerendia (finerenone) | |
|-----------|--|--|
| Formulary | Medicaid - Arizona (AZM, AZMREF, AZMDDD) | |

Formulary Note

Guideline Note:

| Effective Date: | 2/4/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Kerendia | | |
|------------------------|-----------------------|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |

Approval Criteria

1 - Diagnosis of chronic kidney disease (CKD) associated with type 2 diabetes (T2D) defined by one of the following:

1.1 Both of the following:

• Urinary albumin-to-creatinine ratio (UACR) of 30 to 300 mg/g

• Estimated glomerular filtration rate (eGFR) 25 to 60 mL/min/1.73 m2

OR

1.2 Both of the following:

- UACR of greater than or equal to 300 mg/g
- eGFR of 25 to 75 mL/min/1.73 m2

AND

2 - One of the following:

2.1 Minimum 30-day supply trial of a maximally tolerated dose and will continue therapy with one of the following [2]:

- Generic angiotensin-converting enzyme (ACE) inhibitor (e.g., benazepril, lisinopril)
- Generic angiotensin II receptor blocker (ARB) (e.g., losartan, valsartan)

OR

2.2 Patient has a contraindication or intolerance to ACE inhibitors and ARBs

| Product Name: Kerendia | | |
|------------------------|---------------------|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Documentation of positive clinical response to therapy

2 - One of the following:

2.1 Patient continues to be on a maximally tolerated dose of ACE inhibitor or ARB

OR

2.2 Patient has a contraindication or intolerance to ACE inhibitors and ARBs

| Date | Notes |
|----------|-------------|
| 2/3/2022 | New Program |

Keveyis

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99621 Keveyis

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Keveyis | | |
|---|-----------------------|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - ONE of the following: | | |
| 1.1 Diagnosis of primary hyperkalemic periodic paralysis or related variant | | |

OR

1.2 Diagnosis of primary hypokalemic periodic paralysis or related variant

| Product Name: Keveyis | |
|-----------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Keveyis therapy

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Kevzara- Arizona

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99720 Kevzara- Arizona

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Kevzara | |
|-----------------------|---|
| Diagnosis | Moderately to Severely Active Rheumatoid Arthritis (RA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - One of the following:

1.1 All of the following:

1.1.1 Diagnosis of moderately to severely active rheumatoid arthritis (RA)

AND

1.1.2 History of failure to a 3 month trial of ONE non-biologic disease modifying antirheumatic drug (DMARD) [e.g., methotrexate, leflunomide, sulfasalazine, hydroxychloroquine] at maximally indicated doses within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*

AND

1.1.3 History of failure, contraindication, or intolerance to ALL of the following:

- Humira (adalimumab)
- Enbrel (etanercept)
- Xeljanz (tofacitinib)

AND

1.1.4 Patient is NOT receiving Kevzara in combination with ONE of the following:

- Biologic DMARD [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

1.1.5 Prescribed by or in consultation with a rheumatologist

OR

1.2 All of the following:

1.2.1 Patient is currently on Kevzara therapy as documented by claims history or medical records (document drug, date, and duration of therapy)

1.2.2 Diagnosis of moderately to severely active RA

AND

1.2.3 Patient is not receiving Kevzara in combination with ONE of the following:

- Biologic DMARD [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

1.2.4 Prescribed by or in consultation with a rheumatologist

| Notes | *Note: Claims history may be used in conjunction as documentation of |
|-------|--|
| | drug, date, and duration of trial |

| Product Name: Kevzara | |
|-----------------------|---|
| Diagnosis | Moderately to Severely Active Rheumatoid Arthritis (RA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Kevzara therapy

AND

2 - Patient is not receiving Kevzara in combination with ONE of the following:

- Biologic diease modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

3 - Prescribed by or in consultation with a rheumatologist

| Date | Notes |
|-----------|-------------------------------------|
| 5/13/2021 | Arizona Medicaid 7.1 Implementation |

Kimmtrak (tebentafusp-tebn)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-104978 | Kimmtrak (tebentafusp-tebn) |
|-----------|-----------------------------|
|-----------|-----------------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 4/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Kimmtrak | |
|------------------------|--|
| 12 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of uveal melanoma

2 - Disease is unresectable or metastatic

AND

3 - Patient is HLA-A*02:01 genotype positive as determined by a high-resolution genotyping test [2]

AND

4 - Prescribed by or in consultation with an oncologist

| Product Name: Kimmtrak | |
|------------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting patient does not show evidence of progressive disease while on therapy

| Date | Notes |
|-----------|---|
| 3/22/2022 | New Program mirrors ORx with Submission of Records added to initi al and reauth |

Kineret (anakinra)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-114542 Kineret (anakinra)

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Kineret | |
|-----------------------|---------------------------|
| Diagnosis | Rheumatoid Arthritis (RA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes) confirming the diagnosis of moderately to severely active rheumatoid arthritis (RA)

2 - Prescribed by or in consultation with a rheumatologist

AND

3 - Paid claims or submission of medical records (e.g., chart notes) confirming trial and failure, contraindication, or intolerance to ONE nonbiologic disease-modifying antirheumatic drug (DMARD) (e.g., Rheumatrex/Trexall [methotrexate], Arava [leflunomide], Azulfidine [sulfasalazine])

AND

4 - One of the following:

4.1 Both of the following:

4.1.1 Paid claims or submission of medical records (e.g., chart notes) confirming trial and failure, contraindication, or intolerance to ALL of the following, or attestation demonstrating a trial may be inappropriate*

- Enbrel (etanercept)
- Humira (adalimumab)
- Xeljanz (tofacitinib)

AND

4.1.2 Paid claims or submission of medical records (e.g., chart notes) confirming trial and failure, contraindication, or intolerance to Orencia (abatacept)

OR

4.2 Paid claims or submission of medical records (e.g., chart notes) confirming continuation of prior Kineret therapy

| *Includes attestation that a total of two TNF inhibitors have already be en tried in the past, and the patient should not be made to try a third T |
|---|
| NF inhibitor. |

| Product Name: Kineret | |
|-----------------------|---------------------------|
| Diagnosis | Rheumatoid Arthritis (RA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to therapy

| Product Name: Kineret | | |
|-----------------------|---|--|
| Diagnosis | Neonatal-Onset Multisystem Inflammatory Disease (NOMID) | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes) confirming the diagnosis of neonatalonset multisystem inflammatory disease (NOMID)

AND

2 - Diagnosis of NOMID has been confirmed by one of the following:

2.1 NLRP-3 (nucleotide-binding domain, leucine rich family (NLR), pyrin domain containing 3-gene (also known as Cold-Induced Auto-inflammatory Syndrome-1 [CIAS1]) mutation

OR

2.2 Both of the following:

2.2.1 Two of the following clinical symptoms:

• Urticaria-like rash

- Cold/stress triggered episodes
- Sensorineural hearing loss
- Musculoskeletal symptoms (e.g., arthralgia, arthritis, myalgia)
- Chronic aseptic meningitis
- Skeletal abnormalities (e.g., epiphyseal overgrowth, frontal bossing)

2.2.2 Elevated acute phase reactants (e.g., erythrocyte sedimentation rate [ESR], C-reactive protein [CRP], serum amyloid A [SAA])

AND

3 - Prescribed by or in consultation with one of the following

- Allergist/Immunologist
- Rheumatologist
- Pediatrician

| Product Name: Kineret | |
|-----------------------|---|
| Diagnosis | Neonatal-Onset Multisystem Inflammatory Disease (NOMID) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to therapy

| Product Name: Kineret | |
|-----------------------|--|
| Diagnosis | Deficiency of Interleukin-1 Receptor Antagonist (DIRA) |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Submission of medical records (e.g., chart notes) confirming the diagnosis of deficiency of interleukin-1 receptor antagonist (DIRA)

| Product Name: Kineret | |
|-----------------------|---|
| Diagnosis | Systemic Juvenile Idiopathic Arthritis (SJIA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes) confirming the diagnosis of active systemic juvenile idiopathic arthritis

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

3 - Paid claims or submission of medical records (e.g., chart notes) confirming trial and failure, contraindication, or intolerance to ONE of the following:

- Nonsteroidal anti-inflammatory drug (NSAID) (e.g., Motrin [ibuprofen], Naprosyn [naproxen])
- Systemic glucocorticoid (e.g., prednisone)

| Product Name: Kineret | |
|-----------------------|---|
| Diagnosis | Systemic Juvenile Idiopathic Arthritis (SJIA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| | |

Approval Criteria

1 - Documentation of positive clinical response to therapy

| Date | Notes |
|-----------|---|
| 9/26/2022 | Updated criteria, created new criteria for DIRA |

Korlym

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99622 Korlym

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Korlym | |
|---------------------------|-------------------------------|
| Diagnosis | Endogenous Cushing's Syndrome |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - ALL of the following: | |

1.1 Diagnosis of Endogenous Cushing's Syndrome (i.e., hypercortisolism is not a result of chronic administration of high dose glucocorticoids)

AND

1.2 ONE of the following:

- Diagnosis of type 2 diabetes mellitus
- Diagnosis of glucose intolerance

AND

1.3 ONE of the following:

- Patient has failed surgery
- Patient is not a candidate for surgery

| Product Name: Korlym | |
|----------------------|-------------------------------|
| Diagnosis | Endogenous Cushing's Syndrome |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- **1** Documentation of ONE of the following:
 - Patient has improved glucose tolerance while on Korlym therapy
 - Patient has stable glucose tolerance while on Korlym therapy

| Date | Notes |
|------|-------|
|------|-------|

| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |
|-----------|--|
| | |

Korsuva (difelikefalin)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-107424 | Korsuva | (difelikefalin) | |
|-----------|---------|-----------------|--|
| | | | |

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 6/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Korsuva | |
|-----------------------|-----------------------|
| Approval Length | 3 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting all of the following:

1.1 Diagnosis of chronic kidney disease (CKD)

1.2 Patient is currently undergoing hemodialysis (HD) at an optimal dialysis dose (e.g., Kt/V greater than or equal to 1.2)

AND

1.3 Patient is experiencing moderate to severe pruritus associated with CKD (CKD-aP)

AND

1.4 Exclusion of other causes of pruritus (e. g., eczema, infections, drug-induced skin dryness)

AND

1.5 Trial and failure, contraindication, or intolerance to ONE topical anti-pruritic treatment:

- emollient cream
- analgesics (e.g., pramoxine lotion, capsaicin)
- corticosteroids (e.g., hydrocortisone, triamcinolone)

AND

1.6 Trial and failure, contraindication, or intolerance to ONE oral treatment*:

- antihistamine (e.g., diphenhydramine, hydroxyzine, loratadine)
- gabapentin
- pregabalin

AND

- 2 Prescribed by or in consultation with one of the following:
 - Nephrologist

| Dermatologist | |
|---------------|---------------------|
| Notes | *PA may be required |

| Product Name: Korsuva | |
|-----------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting both of the following:

1.1 Patient is currently undergoing hemodialysis

AND

1.2 Documentation of positive clinical response to therapy (e.g., improved quality of life, improved worst itching intensity numerical rating score from baseline)

| Date | Notes |
|-----------|-------------|
| 5/24/2022 | New Program |

Kuvan

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99623 Kuvan

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Kuvan | |
|--|-----------------------|
| Diagnosis | Phenylketonuria (PKU) |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of phenylketonuria (PKU) | |

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

LAMA, LABA - AZM

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-112081 LAMA, LABA - AZM

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 9/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Anoro, Bevespi, Stiolto | | | | |
|---|---------------------|--|--|--|
| Approval Length | 12 month(s) | | | |
| Guideline Type | Prior Authorization | | | |
| | | | | |
| Approval Criteria | | | | |
| 1 - Diagnosis of chronic obstructive pulmonary disease (COPD) | | | | |
| | | | | |
| AND | | | | |
| | | | | |

2 - One of the following:

2.1 History of failure, contraindication, or intolerance to treatment with a 30 day trial of a long-acting beta-agonist (e.g. Foradil, Serevent, Striverdi, Arcapta)

OR

2.2 History of failure, contraindication, or intolerance to treatment with a 30 day trial of an orally inhaled anticholinergic agent (e.g. Spiriva, Atrovent, Combivent, Tudorza)

AND

3 - For Bevespi requests ONLY: history of failure, contraindication, or intolerance to treatment with a 30 day trial of both of the following Preferred drugs:

- Anoro Ellipta
- Stiolto Respimat

| Date | Notes |
|-----------|--|
| 8/22/2022 | Added Anoro Ellipta as target. Added criteria for Bevespi. |

Lansoprazole/Amoxicillin/Clarithromycin

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99556 Lansoprazole/Amoxicillin/Clarithromycin

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Lansoprazole/Amoxicillin/Clarithromycin | | | |
|--|---------------------|--|--|
| Guideline Type | Prior Authorization | | |
| | | | |
| Approval Criteria | | | |
| 1 - Requests for Lansoprazole/Amoxicillin/Clarithromycin should be denied. These medications are available as individual prescriptions (lansoprazole, amoxicillin, and clarithromycin). | | | |

2. Revision History

| Date | Notes |
|-----------|--------------------|
| 6/29/2021 | 7/1 Implementation |

Leqvio (inclisiran)

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-104974 Leqvio (inclisiran)

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 4/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Leqvio | |
|----------------------|---|
| Diagnosis | Heterozygous Familial Hypercholesterolemia (HeFH), Atherosclerotic Cardiovascular Disease (ASCVD) |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting one of the following diagnoses:

1.1 Heterozygous familial hypercholesterolemia (HeFH) as confirmed by one of the following: **1.1.1** Both of the following: [5] 1.1.1.1 Untreated/pre-treatment LDL-cholesterol (LDL-C) greater than 190 mg/dL AND 1.1.1.2 One of the following: Family history of myocardial infarction in first-degree relative less than 60 years of age • Family history of myocardial infarction in second-degree relative less than 50 years of • age Family history of LDL-C greater than 190 mg/dL in first- or second-degree relative • Family history of familial hypercholesterolemia in first- or second-degree relative ٠ Family history of tendinous xanthomata and/or arcus cornealis in first- or second-• degree relative OR **1.1.2** Both of the following: [5] 1.1.2.1 Untreated/pre-treatment LDL-cholesterol (LDL-C) greater than 190 mg/dL AND 1.1.2.2 Submission of medical records (e.g., chart notes, laboratory values) documenting one of the following: Functional mutation in the LDL receptor, ApoB, or PCSK9 gene • Tendinous xanthomata • Arcus cornealis before age 45 OR **1.2** Atherosclerotic cardiovascular disease (ASCVD) as confirmed by one of the following: [2,4] Acute coronary syndromes •

History of myocardial infarction Stable or unstable angina • Coronary or other arterial revascularization • Stroke Transient ischemic attack Peripheral arterial disease presumed to be of atherosclerotic origin AND **2** - One of the following: [4] 2.1 Patient has been receiving at least 12 consecutive weeks of HIGH-INTENSITY statin therapy [i.e., atorvastatin 40-80 mg, rosuvastatin 20-40 mg] and will continue to receive a HIGH-INTENSITY statin at maximally tolerated dose OR **2.2** Both of the following: **2.2.1** Patient is unable to tolerate high-intensity statin as evidenced by one of the following intolerable and persistent (i.e., more than 2 weeks) symptoms: Myalgia (muscle symptoms without CK elevations) • Myositis (muscle symptoms with CK elevations less than 10 times upper limit of normal [ULN]) AND 2.2.2 One of the following: Patient has been receiving at least 12 consecutive weeks of MODERATE-INTENSITY • statin therapy [i.e., atorvastatin 10-20 mg, rosuvastatin 5-10 mg, simvastatin 20-40 mg, pravastatin 40-80 mg, lovastatin 40 mg, Lescol XL (fluvastatin XL) 80 mg, fluvastatin 40 mg twice daily, or Livalo (pitavastatin) 2-4 mg] and will continue to receive a MODERATE-INTENSITY statin at maximally tolerated dose Patient has been receiving at least 12 consecutive weeks of LOW-INTENSITY statin therapy [i.e., simvastatin 10 mg, pravastatin 10-20 mg, lovastatin 20 mg, fluvastatin 20-40 mg, Livalo (pitavastatin) 1 mg] and will continue to receive a LOW-INTENSITY statin at maximally tolerated dose OR

2.3 Patient is unable to tolerate low- or moderate-, and high-intensity statins as evidenced by one of the following intolerable and persistent (i.e., more than 2 weeks) symptoms for low- or moderate-, and high-intensity statins: Myalgia (muscle symptoms without CK elevations) • Myositis (muscle symptoms with CK elevations less than 10 times ULN) • OR 2.4 Patient has a labeled contraindication to all statins OR 2.5 Patient has experienced rhabdomyolysis or muscle symptoms with statin treatment with CK elevations greater than 10 times ULN [4] AND 3 - One of the following: **3.1** Patient has been receiving at least 12 consecutive weeks of ezetimibe (Zetia) therapy as adjunct to maximally tolerated statin therapy OR 3.2 Patient has a history of contraindication or intolerance to ezetimibe AND 4 - Patient is unable to maintain adherence to proprotein convertase subtilisin/kexin type 9 (PCSK9) inhibitor therapy AND 5 - Submission of medical records (e.g., laboratory values) documenting one of the following LDL-C values while on maximally tolerated lipid lowering therapy within the last 120 days:

• LDL-C greater than or equal to 70 mg/dL for diagnosis of ASCVD [2]

• LDL-C greater than or equal to 100 mg/dL for diagnosis of HeFH [3]

AND

6 - Prescribed by or in consultation with one of the following:

- Cardiologist
- Endocrinologist
- Lipid specialist

AND

7 - Medication will not be used in combination with PCSK9 inhibitor therapy [2,3]

| Product Name: Leqvio | |
|----------------------|---|
| Diagnosis | Heterozygous Familial Hypercholesterolemia (HeFH), Atherosclerotic Cardiovascular Disease (ASCVD) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting LDL-C reduction from baseline while on therapy

AND

2 - One of the following:

2.1 Patient continues to receive other lipid-lowering therapy (e.g., statins, ezetimibe) at the maximally tolerated dose

2.2 Patient has a documented inability to take other lipid-lowering therapy (e.g., statins, ezetimibe)

AND

3 - Medication will not be used in combination with PCSK9 inhibitor therapy [2,3]

2. Revision History

| Date | Notes |
|-----------|---|
| 3/22/2022 | New program, mirrors ORx with Submission of Records requirement |

Leucovorin- Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99469 Leucovorin- A | rizona |
|------------------------|--------|
|------------------------|--------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Leucovorin tabs | | |
|---------------------------------------|---------------------|--|
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - ONE of the following: | | |
| 1.1 Methotrexate toxicity prophylaxis | | |

OR

1.2 Treatment of hematologic toxicity from folic acid antagonists (i.e., pyrimethamine toxicity treatment or trimethoprim toxicity treatment)

2. Revision History

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk copy C&S Arizona standard to Medicaid Arizona |

Lidocaine - Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Lidocaine 2% Gel | | |
|---|---------------------|--|
| Approval Length 6 month(s) | | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Used with catheters or open mucus membrane areas. | | |

| Product Name: Lidocaine 4 or 5% ointment | |
|--|---------------------|
| Guideline Type | Prior Authorization |

1 - Requests for Lidocaine 4 or 5% ointment should be denied. The plan's preferred product is Aspercreme with Lidocaine 4%.

2. Revision History

| Date | Notes |
|-----------|-------------------|
| 7/14/2021 | Updated guideline |

Livtencity (maribavir)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-113529 | Livtencity (maribavir) | |
|-----------|------------------------|--|
|-----------|------------------------|--|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 9/8/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| DiagnosisCMV infection/diseaseApproval Length8 Week(s) | Product Name: Livtencity | |
|--|--------------------------|-----------------------|
| Approval Length 8 Week(s) | Diagnosis | CMV infection/disease |
| | Approval Length | 8 Week(s) |
| Guideline Type Prior Authorization | Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of cytomegalovirus (CMV) infection/disease as confirmed by one of the following methods:

• quantitative polymerase chain reaction (qPCR)

| CMV pp65 antigenemia | | |
|--|--|--|
| AND | | |
| 2 - Patient is a recipient of one of the following: | | |
| Hematopoietic stem cell transplantSolid organ transplant | | |
| AND | | |
| 3 - Trial and failure of a minimum 2 weeks duration, contraindication, or intolerance to one of the following therapies at an appropriately indicated dose: | | |
| Intravenous (IV) ganciclovir Oral valganciclovir IV foscarnet IV cidofovir | | |
| AND | | |
| 4 - Patient is 12 years of age or older | | |
| AND | | |
| 5 - Patient weighs greater than or equal to 35kg | | |
| AND | | |
| 6 - Prescribed by or in consultation with a provider who specializes in one of the following areas: | | |
| Transplant Infectious Disease | | |

2. Revision History

| Date | Notes |
|----------|---|
| 9/8/2022 | Removed references and end note, no changes to clinical criteria. |

Long-Acting Opioid Products - AZM

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-112078 Long-Acting Opioid Products - AZM

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 8/23/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

Product Name: Generic morphine sulfate ER tablets, brand Duragesic, generic fentanyl, Xtampza, generic morphine sulfate beads cap ER, generic hydromorphone ER, brand Kadian, generic morphine sulfate ER capsules, brand MS Contin, Zohydro ER, generic oxymorphone ER (non-crush resistant), brand Methadose, brand Dolophine, generic methadone, Arymo ER, Morphabond ER, Hysingla ER, brand Oxycontin, generic oxycodone ER, Nucynta ER, brand Conzip, generic tramadol ER 24HR biphasic capsules, generic tramadol ER biphasic release tablets, generic tramadol ER tablets

| Diagnosis | PA REQUIRED for use of MAT and other Opioids (Reject 88) |
|----------------|--|
| Guideline Type | DUR |
| | |

Approval Criteria

1 - Provider attests to notify the prescriber of the MAT therapy and the prescriber of the MAT therapy approves the concurrent opioid therapy.

AND

2 - The days supply does not exceed 14 days for a surgical procedure.

AND

3 - The days supply does not exceed 5 days for all other requests.

AND

4 - There has not been a previous approval in the last 6 months.

| Notes | Approval Length: 14 Days for surgical procedure, 5 Days for all other r |
|-------|---|
| | equests |

| Product Name: Generic morphine sulfate ER tablets, generic fentanyl 12 mcg/hr, 25 mcg/hr, 50 mcg/hr, 75 mcg/hr, 100 mcg/hr, Xtampza, generic tramadol ER tablets | |
|--|---|
| Diagnosis | Cancer related pain/Hospice care/end-of-life care* |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - ONE of the following: | |
| 1.1 Patient is being treated for cancer | |
| OR | |
| 1.2 Patient is receiving hospice or end-of-life care | |
| Notes | *Note: If the member is currently taking the requested long-acting opio id for at least 30 days and does not meet the medical necessity author |

| ization criteria requirements for treatment with an opioid, a denial sho |
|--|
| uld be issued and a maximum 30 day authorization may be authorized |
| one time for the requested drug/strength combination up to the reque |
| sted quantity for transition to an alternative treatment. |

Product Name: Generic morphine sulfate beads cap ER, generic hydromorphone ER, brand Kadian, generic morphine sulfate ER capsules, brand MS Contin, generic fentanyl transdermal 37.5mcg/hr, 62.5mcg/hr, and 87.5mcg/hr, brand Zohydro ER, hydrocodone ER capsules (generic Zohydro ER), generic oxymorphone ER (non-crush resistant), brand Methadose, brand Dolophine, generic methadone, Arymo ER, Morphabond ER, Hysingla ER, hydrocodone ER tablets (generic Hysingla ER), brand Oxycontin, generic oxycodone ER, Nucynta ER, brand Duragesic

| Diagnosis | Cancer related pain/Hospice care/end-of-life care* |
|-----------------|--|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - ONE of the following:

1.1 Patient is being treated for cancer

OR

1.2 Patient is receiving hospice or end-of-life care

AND

2 - BOTH of the following:

2.1 ONE of the following:

2.1.1 The patient has a history of failure, contraindication or intolerance to a trial of at least THREE of the following (Document drugs and date of trials):*

- morphine sulfate controlled release tablets (specifically generic MS Contin)
- preferred fentanyl transdermal**
- Butrans (buprenorphine)
- Xtampza ER (oxycodone extended-release)
- tramadol extended release tablets (non-biphasic release tablets)

FENTANYL PATCH 72-HOUR 12mcg, 25mcg, 50mcg, 75mcg & 100mcg

OR

2.1.2 Patient is established on pain therapy with the requested medication for cancer, hospice care, or end-of-life care pain, and the medication is not a new regimen for treatment of cancer, hospice care, or end-of-life care pain (Document date regimen was started)

AND

2.2 Prescriber attests to the following: the information provided is true and accurate to the best of their knowledge and they understand that OptumRx may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided

| Notes | *Note: If the member is currently taking the requested long-acting opio id for at least 30 days and does not meet the medical necessity autho r ization criteria requirements for treatment with an opioid, a denial sh o uld be issued and a maximum 30-day authorization may be authoriz e d one time for the requested drug/strength combination up to the req u ested quantity for transition to an alternative treatment. *Note: If the re quest is for a non-preferred product and the member is currently ta kin g the requested long-acting opioid for at least 30 days and has me t th e medical necessity authorization criteria requirements for treatme nt w ith an opioid, but has not tried the preferred alternatives a denial shoul d be issued and a maximum 30-day authorization may be autho rized one time for the requested drug/strength combination up to the r eques ted quantity for transition to an alternative treatment. **NOTE: Fentanyl transdermal 37.5mcg/hr, 62.5mcg/hr, and 87.5m cg/hr are n on-preferred *Note: Claims bistory may be used in conjunction as doc |
|-------|---|
| | on-preferred. *Note: Claims history may be used in conjunction as doc umentation of drug, date, and duration of trial. |

| Product Name: Brand Conzip, generic tramadol ER 24HR biphasic capsules, generic tramadol ER biphasic release tablets | |
|--|--|
| Diagnosis | Cancer related pain/Hospice care/end-of-life care* |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - ONE of the following:

1.1 Patient is being treated for cancer

OR

1.2 Patient is receiving hospice or end-of-life care

AND

2 - BOTH of the following:

2.1 ONE of the following:

2.1.1 The patient has a history of failure, contraindication or intolerance to a trial of BOTH of the following (Document drugs and date of trials):*

- tramadol immediate release (IR)
- tramadol extended release tablets (non-biphasic release tablets)

OR

2.1.2 Patient is established on pain therapy with the requested medication for cancer, hospice care, or end-of-life care pain, and the medication is not a new regimen for treatment of cancer, hospice care, or end-of-life care pain (Document date regimen was started)

AND

2.2 Prescriber attests to the following: the information provided is true and accurate to the best of their knowledge and they understand that OptumRx may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided

| | *Note: If the member is currently taking the requested long-acting opio id for at least 30 days and does not meet the medical necessity autho r ization criteria requirements for treatment with an opioid, a denial sh o uld be issued and a maximum 30-day authorization may be authoriz ed one time for the requested drug/strength combination up to the req u ested quantity for transition to an alternative treatment. *Note: Claim s history may be used in conjunction as documentation of drug, date, and duration of trial. |
|--|---|
|--|---|

| 50 mcg/hr, 75 mcg/hr, 100 mcg/hr, Xtampza, generic tramadol ER tablets | |
|--|---|
| Diagnosis | Non-cancer pain/Non-hospice care/Non-end-of-life care pain* |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

1 - Prescriber attests to ALL of the following:

- The information provided is true and accurate to the best of their knowledge and they understand that OptumRx may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided
- Treatment goals are defined, including estimated duration of treatment
- Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic intervention
- Patient has been screened for substance abuse/opioid dependence
- If used in patients with medical comorbidities or if used concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression
- Pain is moderate to severe and expected to persist for an extended period of time
- Pain is chronic
- Pain is not postoperative (unless the patient is already receiving chronic opioid therapy prior to surgery, or if the postoperative pain is expected to be moderate to severe and persist for an extended period of time)
- Pain management is required around the clock with a long-acting opioid

AND

2 - ONE of the following:

2.1 Prior to the start of therapy with the long-acting opioid, the patient has failed an adequate (minimum of 2 week) trial of a short-acting opioid within the last 30 days (Document drug(s) and date of trial)*

OR

2.2 The patient is already receiving chronic opioid therapy prior to surgery for postoperative pain

OR

2.3 Postoperative pain is expected to be moderate to severe and persist for an extended period of time

AND

3 - If the request for neuropathic pain (examples of neuropathic pain include neuralgias, neuropathies, fibromyalgia), BOTH of the following:

3.1 Unless it is contraindicated, the patient has not exhibited an adequate response to 8 weeks of treatment with gabapentin titrated to a therapeutic dose (Document date of trial)*

AND

3.2 Unless it is contraindicated, the patient has not exhibited an adequate response to at least 6 weeks of treatment with a tricyclic antidepressant titrated to the maximum tolerated dose. (Document drug and date of trial)*

| Notes | *Note: If the member is currently taking the requested long-acting opio id for at least 30 days and does not meet the medical necessity autho r ization criteria requirements for treatment with an opioid, a denial sh o uld be issued and a maximum 30-day authorization may be authoriz ed one time for the requested drug/strength combination up to the req u ested quantity for transition to an alternative treatment. *Note: Claim s history may be used in conjunction as documentation of drug, date, and duration of trial **NOTE: Fentanyl transdermal 37.5mcg/hr, 62.5m cg/hr, and 87.5 mcg/hr are non-preferred. |
|-------|--|
|-------|--|

Product Name: Generic morphine sulfate beads cap ER, generic hydromorphone ER, brand Kadian, generic morphine sulfate ER capsules, brand MS Contin, generic fentanyl transdermal 37.5mcg/hr, 62.5mcg/hr, and 87.5mcg/hr, brand Zohydro ER, hydrocodone ER capsules (generic Zohydro ER), generic oxymorphone ER (non-crush resistant), brand Methadose, brand Dolophine, generic methadone, Arymo ER, Morphabond ER, Hysingla ER, hydrocodone ER tablets (generic Hysingla ER), brand Oxycontin, generic oxycodone ER, Nucynta ER, brand Duragesic

| Diagnosis | Non-cancer pain/Non-hospice care/Non-end-of-life care pain* |
|-----------------|---|
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |

| Guideline Type | Prior Authorization | |
|----------------|---------------------|--|
|----------------|---------------------|--|

1 - Prescriber attests to ALL of the following:

- The information provided is true and accurate to the best of their knowledge and they understand that OptumRx may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided
- Treatment goals are defined, including estimated duration of treatment
- Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic intervention
- Patient has been screened for substance abuse/opioid dependence
- If used in patients with medical comorbidities or if used concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression
- Pain is moderate to severe and expected to persist for an extended period of time
- Pain is not postoperative (unless the patient is already receiving chronic opioid therapy prior to surgery, or if the postoperative pain is expected to be moderate to severe and persist for an extended period of time)
- Pain management is required around the clock with a long-acting opioid

AND

2 - ONE of the following:

2.1 Prior to the start of therapy with the long-acting opioid, the patient has failed an adequate (minimum of 2 week) trial of a short-acting opioid within the last 30 days (Document drug(s) and date of trial)*

OR

2.2 The patient is already receiving chronic opioid therapy prior to surgery for postoperative pain

OR

2.3 Postoperative pain is expected to be moderate to severe and persist for an extended period of time

AND

3 - The patient has a history of failure, contraindication or intolerance to at least THREE of the following (Document drugs and date of trials):)*

- morphine sulfate controlled release tablets (specifically generic MS Contin)
- preferred fentanyl transdermal**
- Butrans (buprenorphine)
- Xtampza ER (oxycodone extended-release)
- tramadol extended release tablets (non-biphasic release tablets)
- FENTANYL PATCH 72-HOUR 12mcg, 25mcg, 50mcg, 75mcg & 100mcg

AND

4 - If the request for neuropathic pain (examples of neuropathic pain include neuralgias, neuropathies, fibromyalgia), BOTH of the following:

4.1 Unless it is contraindicated, the patient has not exhibited an adequate response to 8 weeks of treatment with gabapentin titrated to a therapeutic dose (Document date of trial))*

AND

4.2 Unless it is contraindicated, the patient has not exhibited an adequate response to at least 6 weeks of treatment with a tricyclic antidepressant titrated to the maximum tolerated dose. (Document drug and date of trial))*

| Notes | *Note: If the member is currently taking the requested long-acting opio id for at least 30 days and does not meet the medical necessity autho r ization criteria requirements for treatment with an opioid, a denial sh o uld be issued and a maximum 30-day authorization may be authoriz e d one time for the requested drug/strength combination up to the req u ested quantity for transition to an alternative treatment. *Note: If the re quest is for a non-preferred product and the member is currently ta kin g the requested long-acting opioid for at least 30 days and has me t th e medical necessity authorization criteria requirements for treatme nt w ith an opioid, but has not tried the preferred alternatives a denial shoul d be issued and a maximum 30-day authorization may be autho rized one time for the requested drug/strength combination up to the r eques ted quantity for transition to an alternative treatment. Additionall y **NOTE: Fentanyl transdermal 37.5mcg/hr, 62.5mcg/hr, and 87.5 m cg/hr are non-preferred. *Note: Claims history may be used in conjunc tion as documentation of drug, date, and duration of trial. |
|-------|--|

| Product Name: Brand Conzip, generic tramadol ER 24HR biphasic capsules, generic tramadol ER biphasic release tablets | |
|--|---|
| Diagnosis | Non-cancer pain/Non-hospice care/Non-end-of-life care pain* |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

1 - Prescriber attests to ALL of the following:

- The information provided is true and accurate to the best of their knowledge and they understand that OptumRx may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided
- Treatment goals are defined, including estimated duration of treatment
- Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic intervention
- Patient has been screened for substance abuse/opioid dependence
- If used in patients with medical comorbidities or if used concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression
- Pain is moderate to severe and expected to persist for an extended period of time
- Pain is chronic
- Pain is not postoperative (unless the patient is already receiving chronic opioid therapy prior to surgery, or if the postoperative pain is expected to be moderate to severe and persist for an extended period of time)
- Pain management is required around the clock with a long-acting opioid

AND

2 - ONE of the following:

2.1 Prior to the start of therapy with the long-acting opioid, the patient has failed an adequate (minimum of 2 week) trial of a short-acting opioid within the last 30 days (Document drug(s) and date of trial)*

OR

2.2 The patient is already receiving chronic opioid therapy prior to surgery for postoperative pain

OR

2.3 Postoperative pain is expected to be moderate to severe and persist for an extended period of time

AND

3 - If the request for neuropathic pain (examples of neuropathic pain include neuralgias, neuropathies, fibromyalgia), BOTH of the following:

3.1 Unless it is contraindicated, the patient has not exhibited an adequate response to 8 weeks of treatment with gabapentin titrated to a therapeutic dose (Document date of trial)*

AND

3.2 Unless it is contraindicated, the patient has not exhibited an adequate response to at least 6 weeks of treatment with a tricyclic antidepressant titrated to the maximum tolerated dose. (Document drug and date of trial)*

AND

3.3 The patient has a history of failure, contraindication or intolerance to BOTH of the following (Document drugs and date of trials):)*

- tramadol immediate release (IR)**
- tramadol extended release tablets (non-biphasic release tablets)**

| Notes | *Note: If the member is currently taking the requested long-acting opio id for at least 30 days and does not meet the medical necessity autho r ization criteria requirements for treatment with an opioid, a denial sh o uld be issued and a maximum 30-day authorization may be authoriz e d one time for the requested drug/strength combination up to the req u ested quantity for transition to an alternative treatment. *Note: If the re quest is for tramadol extended release capsules or tramadol extend ed release biphasic release tablets and the member is currently taking th e requested long-acting opioid for at least 30 days and has met the m edical necessity authorization criteria requirements for treatment wi th an opioid, but has not tried the preferred alternatives a denial shoul |
|-------|--|
| | th an opioid, but has not tried the preferred alternatives a denial shoul d b e issued and a maximum 30-day authorization may be authorized |

| one time for the requested drug/strength combination up to the reques ted quantity for transition to an alternative treatment. *Drug may requir e prior authorization *Note: Claims history may be used in conjunction |
|--|
| as documentation of drug, date, and duration of trial. |

| Product Name: Generic morphine sulfate ER tablets, generic fentanyl 12 mcg/hr, 25 mcg/hr, 50 mcg/hr, 75 mcg/hr, 100 mcg/hr, Xtampza, generic tramadol ER tablets | |
|--|---|
| Diagnosis | Non-cancer pain/Non-hospice care/Non-end-of-life care pain* |
| Approval Length | 6 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Patient demonstrates meaningful improvement in pain and function (Document improvement in function or pain score improvement)

AND

2 - Identify rationale for not tapering and discontinuing opioid (Document rationale)

AND

3 - Prescriber attests to ALL of the following:

- The information provided is true and accurate to the best of their knowledge and they understand that OptumRx may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided
- Treatment goals are defined, including estimated duration of treatment
- Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic intervention
- Patient has been screened for substance abuse/opioid dependence
- If used in patients with medical comorbidities or if used concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression
- Pain is moderate to severe and expected to persist for an extended period of time
- Pain is chronic

- Pain is not postoperative (unless the patient is already receiving chronic opioid therapy prior to surgery, or if the postoperative pain is expected to be moderate to severe and persist for an extended period of time)
- Pain management is required around the clock with a long-acting opioid

| Notes | *Note: If the member is currently taking the requested long-acting opio id for at least 30 days and does not meet the medical necessity autho r ization criteria requirements for treatment with an opioid, a denial sh o uld be issued and a maximum 30-day authorization may be authoriz e d one time for the requested drug/strength combination up to the req u ested quantity for transition to an alternative treatment. *Note: If the re quest is for a non-preferred product and the member is currently ta kin g the requested long-acting opioid for at least 30 days and has me t th e medical necessity authorization criteria requirements for treatme nt w ith an opioid, but has not tried the preferred alternatives a denial shoul d be issued and a maximum 30-day authorization may be autho rized one time for the requested drug/strength combination up to the r eques ted quantity for transition to an alternative treatment. **NOTE: Fentanyl transdermal 37.5mcg/hr, 62.5mcg/hr, and 87.5m cg/hr are n on-preferred. |
|-------|--|

Product Name: Generic morphine sulfate beads cap ER, generic hydromorphone ER, brand Kadian, generic morphine sulfate ER capsules, brand MS Contin, generic fentanyl transdermal 37.5mcg/hr, 62.5mcg/hr, and 87.5mcg/hr, brand Zohydro ER, hydrocodone ER capsules (generic Zohydro ER), generic oxymorphone ER (non-crush resistant), brand Methadose, brand Dolophine, generic methadone, Arymo ER, Morphabond ER, Hysingla ER, hydrocodone ER tablets (generic Hysingla ER), brand Oxycontin, generic oxycodone ER, Nucynta ER, brand Duragesic

| Diagnosis | Non-cancer pain/Non-hospice care/Non-end-of-life care pain* |
|-----------------|---|
| Approval Length | 6 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient demonstrates meaningful improvement in pain and function (Document improvement in function or pain score improvement)

AND

2 - Identify rationale for not tapering and discontinuing opioid (Document rationale)

| | AND |
|--|---|
| 3 - Prescriber atte | ests to ALL of the following: |
| The information provided is true and accurate to the best of their knowledge and they understand that OptumRx may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided Treatment goals are defined, including estimated duration of treatment Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic intervention Patient has been screened for substance abuse/opioid dependence If used in patients with medical comorbidities or if used concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression Pain is moderate to severe and expected to persist for an extended period of time Pain is not postoperative (unless the patient is already receiving chronic opioid therapy prior to surgery, or if the postoperative pain is expected to be moderate to severe and persist for an extended period of time) Pain management is required around the clock with a long-acting opioid | |
| Notes | *Note: If the member is currently taking the requested long-acting opic id for at least 30 days and does not meet the medical necessity author r ization criteria requirements for treatment with an opioid, a denial sh o uld be issued and a maximum 30-day authorization may be authoriz e d one time for the requested drug/strength combination up to the requested quantity for transition to an alternative treatment. *Note: If the re quest is for a non-preferred product and the member is currently ta kin g the requested long-acting opioid for at least 30 days and has me t th e medical necessity authorization criteria requirements for treatment nt w ith an opioid, but has not tried the preferred alternatives a denial shoul d be issued and a maximum 30-day authorization may be author rized one time for the requested drug/strength combination up to the r eques ted quantity for transition to an alternative treatment.**NOTE: F entanyl transdermal 37.5mcg/hr, 62.5mcg/hr, and 87.5m cg/hr are nor -preferred. |

| Product Name: Brand Conzip, generic tramadol ER 24HR biphasic capsules, generic tramadol ER biphasic release tablets | |
|--|---|
| Diagnosis | Non-cancer pain/Non-hospice care/Non-end-of-life care pain* |
| Approval Length | 6 month(s) |
| Therapy Stage | Reauthorization |

| Guideline Type | Prior Authorization |
|--|---|
| | |
| Approval Criteria | |
| | es meaningful improvement in pain and function (Document n or pain score improvement) |
| | AND |
| 2 - Identify rationale for | not tapering and discontinuing opioid (Document rationale) |
| | AND |
| 3 - Prescriber attests to | ALL of the following: |
| The information provided is true and accurate to the best of their knowledge and they understand that OptumRx may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided Treatment goals are defined, including estimated duration of treatment Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic intervention Patient has been screened for substance abuse/opioid dependence If used in patients with medical comorbidities or if used concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression Pain is moderate to severe and expected to persist for an extended period of time Pain is not postoperative (unless the patient is already receiving chronic opioid therapy prior to surgery, or if the postoperative pain is expected to be moderate to severe and persist for an extended period of time) Pain management is required around the clock with a long-acting opioid | |
| Notes | *Note: If the member is currently taking the requested long-acting opio id for at least 30 days and does not meet the medical necessity autho r ization criteria requirements for treatment with an opioid, a denial sh o uld be issued and a maximum 30-day authorization may be authoriz e d one time for the requested drug/strength combination up to the req u ested quantity for transition to an alternative treatment. *Note: If the re quest is for tramadol extended release capsules or tramadol extend ed release biphasic release tablets and the member is currently taking th e requested long-acting opioid for at least 30 days and has met the m edical necessity authorization criteria requirements for treatment wi |

| th an opioid, but has not tried the preferred alternatives a denial shoul d b e issued and a maximum 30-day authorization may be authorized |
|---|
| one time for the requested drug/strength combination up to the reques ted quantity for transition to an alternative treatment. |

Product Name: Generic morphine sulfate ER tablets, brand Duragesic, generic fentanyl, Xtampza, generic morphine sulfate beads cap ER, generic hydromorphone ER, brand Kadian, generic morphine sulfate ER capsules, brand MS Contin, brand Zohydro ER, hydrocodone ER capsules (generic Zohydro ER), generic oxymorphone ER (non-crush resistant), brand Methadose, brand Dolophine, generic methadone, Arymo ER, Morphabond ER, Hysingla ER, hydrocodone ER tablets (generic Hysingla ER), brand Oxycontin, generic oxycodone ER, Nucynta ER, brand Conzip, generic tramadol ER 24HR biphasic capsules, generic tramadol ER biphasic release tablets, generic tramadol ER tablets

| Diagnosis | Criteria for Quantity Limit Reviews* |
|----------------|--------------------------------------|
| Guideline Type | Quantity Limit |

Approval Criteria

1 - The requested dose cannot be achieved by moving to a higher strength of the product

AND

2 - The requested dose is within the Food and Drug Administration (FDA) maximum dose per day, where an FDA maximum dose per day exists (see Table 1 in the Background section)

| *Note: Authorization will be issued for • Cancer pain/hospice/end-of-lif e related pain: 12 months • All Tramadol ER requests: 12 months • No |
|--|
| n-cancer pain/non-hospice/non-end-of-life related pain: 6 months |

Product Name: Generic morphine sulfate ER tablets, brand Duragesic, generic fentanyl, Xtampza, generic morphine sulfate beads cap ER, generic hydromorphone ER, brand Kadian, generic morphine sulfate ER capsules, brand MS Contin, Zohydro ER, generic oxymorphone ER (non-crush resistant), brand Methadose, brand Dolophine, generic methadone, Arymo ER, Morphabond ER, Hysingla ER, brand Oxycontin, generic oxycodone ER, Nucynta ER, brand Conzip, generic tramadol ER 24HR biphasic capsules, generic tramadol ER biphasic release tablets, generic tramadol ER tablets

| Diagnosis | Opioid Naïve (Not having filled an opioid in the past 120 days)* |
|----------------|--|
| Guideline Type | Morphine Milligram Equivalents (MME)** MME 50.00 exceeded; PA Required for dosage above 50 MEDD |
| | |

1 - Opioid naïve members may receive greater than 50 morphine milligram equivalent (MME) based on the following:

1.1 If the request is for 50 MME to 90 MME, ONE of the following (NOTE: If the request exceeds 90 MME please skip this section and proceed to the Exceeding the 90 MME Cumulative Threshold Reviews section):

1.1.1 Diagnosis of ONE of the following:

- Cancer
- End of life pain (including hospice care)
- Palliative care
- Sickle cell anemia

OR

1.1.2 Patient is currently exceeding 50 MME and prescriber attests patient has been on a short-acting opioid in the past 120 days

OR

1.1.3 Document ALL of the following:

- The diagnosis associated with the need for pain management with opioid
- If used in patients with medical comorbidities or if used concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression
- The prescriber has acknowledged that they have completed an addiction risk and risk of overdose assessment
- Prescriber attests the member requires more than 50 MME per day to adequately control pain

Product Name: Generic morphine sulfate ER tablets, brand Duragesic, generic fentanyl, Xtampza, generic morphine sulfate beads cap ER, generic hydromorphone ER, brand Kadian, generic morphine sulfate ER capsules, brand MS Contin, Zohydro ER, generic oxymorphone ER (non-crush resistant), brand Methadose, brand Dolophine, generic methadone, Arymo ER, Morphabond ER, Hysingla ER, brand Oxycontin, generic oxycodone ER, Nucynta ER, brand Conzip, generic tramadol ER 24HR biphasic capsules, generic tramadol ER biphasic release tablets, generic tramadol ER tablets

| Diagnosis | Doses Exceeding the Cumulative MME of 90 mg - Cancer/Hospice/End-of-Life/Palliative Care/Skilled Nursing Facility/Traumatic Injury Related Pain* |
|-----------------|--|
| Approval Length | 12 month(s) |
| Guideline Type | Morphine Milligram Equivalent (MME)** (MME 90.00 exceeded; PA REQUIRED; Dosage Above MEDD Limit) |

1 - Doses exceeding the cumulative morphine milligram equivalent (MME) of 90 milligrams will be approved up to the requested amount for ALL opioid products if the patient has one of the following conditions:

- Active oncology diagnosis
- Hospice care
- End-of-life care (other than hospice)
- Palliative care
- Skilled nursing facility care
- Traumatic injury, including burns and excluding post-surgical procedure

AND

2 - Provider attests patient has been prescribed naloxone (may also be verified via paid pharmacy claims)

| Notes | *Note: Authorization will be issued for 12 months for one of the above |
|-------|--|
| | conditions. The authorization should be entered for an MME of 9999 s o as to prevent future disruptions in therapy if the patient's dose is inc r eased. |

Product Name: Generic morphine sulfate ER tablets, brand Duragesic, generic fentanyl, Xtampza, generic morphine sulfate beads cap ER, generic hydromorphone ER, brand Kadian, generic morphine sulfate ER capsules, brand MS Contin, Zohydro ER, generic oxymorphone ER (non-crush resistant), brand Methadose, brand Dolophine, generic methadone, Arymo ER, Morphabond ER, Hysingla ER, brand Oxycontin, generic oxycodone ER, Nucynta ER, brand Conzip, generic tramadol ER 24HR biphasic capsules, generic tramadol ER biphasic release tablets, generic tramadol ER tablets

| Diagnosis | Doses- Exceeding the Cumulative MME of 90 mg - Non-cancer/non- hospice/non-end-of-life/non-palliative care/non-skilled Nursing Facility/Traumatic Injury Related Pain* |
|----------------------------|--|
| Approval Length | 6 month(s) |
| Therapy Stage Initial Auth | Initial Authorization |

| Guideline Type | Morphine Milligram Equivalent (MME)** MME 90.00 exceeded; PA REQUIRED; Dosage Above MEDD Limit | |
|--|---|--|
| Approval Criteria | | |
| 1 - Prescriber attests to | ALL of the following: | |
| understand that information nece Treatment goals Treatment plan intervention | • Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic | |
| if used in patien benzodiazepine prescriber has a | Patient has been screened for substance abuse/opioid dependence if used in patients with medical comorbidities or if used concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression | |
| | AND | |
| 2 - BOTH of the followir | ng: | |
| 2.1 Patient has tried and failed non-opioid pain medication (document drug name and date of trial) | | |
| AND | | |
| 2.2 Opioid medication doses of less than 90 morphine milligram equivalent (MME) have been tried and did not adequately control pain (document drug regimen or MME and dates of therapy) | | |
| | AND | |
| 3 - Provider attests patient has been prescribed naloxone (may also be verified via paid pharmacy claims) | | |
| Notes | *Note: If the member has been established on the requested MME do se for at least 30 days and does not meet the medical necessity autho rization criteria requirements, a denial should be issued and a maximu m 30 -day authorization may be authorized one time for the requested MME dose. **Note: Authorization will be issued for 6 months for non- | |

| | cancer/non-hospice/non-end-of-life/non-palliative care/non-skilled nur si ng facility/non-traumatic injury related pain up to the current request |
|--|--|
| | ed MME plus 90 MME. |

Product Name: Generic morphine sulfate ER tablets, brand Duragesic, generic fentanyl, Xtampza, generic morphine sulfate beads cap ER, generic hydromorphone ER, brand Kadian, generic morphine sulfate ER capsules, brand MS Contin, Zohydro ER, generic oxymorphone ER (non-crush resistant), brand Methadose, brand Dolophine, generic methadone, Arymo ER, Morphabond ER, Hysingla ER, brand Oxycontin, generic oxycodone ER, Nucynta ER, brand Conzip, generic tramadol ER 24HR biphasic capsules, generic tramadol ER biphasic release tablets, generic tramadol ER tablets

| Desse Eveneding the Cumulative MME of 00 mg. Non concer/non |
|---|
| Doses Exceeding the Cumulative MME of 90 mg - Non-cancer/non- hospice/non-end-of-life/non-palliative Nursing Facility/Traumatic Injury Related Pain* care/non-skilled |
| 6 month(s) |
| Reauthorization |
| Morphine Milligram Equivalent (MME)** MME 90.00 exceeded; PA REQUIRED; Dosage Above MEDD Limit |
| |

Approval Criteria

- 1 Prescriber attests to ALL of the following:
 - The information provided is true and accurate to the best of their knowledge and they understand that OptumRx may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided
 - Treatment goals are defined, including estimated duration of treatment
 - Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic intervention
 - Patient has been screened for substance abuse/opioid dependence
 - if used in patients with medical comorbidities or if used concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression

AND

2 - Identify rationale for not tapering and discontinuing opioid (Document rationale)

AND

3 - Patient demonstrates meaningful improvement in pain and function (Document improvement in function or pain score improvement)

AND

4 - Provider attests patient has been prescribed naloxone (may also be verified via paid pharmacy claims)

| Notes | *Note: If the member has been established on the requested MME do se for at least 30 days and does not meet the medical necessity autho rization criteria requirements, a denial should be issued and a maximu m 30 -day authorization may be authorized one time for the requested MME dose. **Note: Authorization will be issued for 6 months for non- cancer/non-hospice/non-end-of-life/non-palliative care/non-skilled nur si ng facility/non-traumatic injury related pain up to the current request ed MME plus 90 MME. |
|-------|---|
|-------|---|

2. Background

| FDA Label Max Daily Doses None |
|--------------------------------|
| None |
| |
| None |
| None |
| |
| None |
| |
| None |
| |

| Oxymorphone | None |
|---|--|
| Oxycodone | Xtampza Only =288mg |
| *Doses are not considered equianalgesic an chart. | d table does not represent a dose conversion |
| | ed on morphine milligram equivalents allowed specialist. Max MME is based upon the CDC e product strengths. Fentanyl is dosed in |

3. Revision History

| Date | Notes |
|-----------|--|
| 8/22/2022 | Added attestation criteria for naloxone rx requirement to 90 MME Exc eeded sections. |

Lonhala and Yupelri- Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Lonha | la Magnair, Yupleri |
|---------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of moderate to severe chronic obstructive pulmonary disease (COPD

AND

2 - ONE of the following:

2.1 History of failure, contraindication or intolerance to Spiriva Handihaler (tiotropium)

OR

2.2 BOTH of the following:

2.2.1 Patient is unable to use a metered-dose, dry powder or slow mist inhaler (e.g. Spiriva Handihaler) to control his/her COPD due to ONE of the following

- Cognitive or physical impairment limiting coordination of handheld devices (e.g., cognitive decline, arthritis in the hands) (Document impairment)
- Patient is unable to generate adequate inspiratory force (e.g., peak inspiratory flow rate (PIFR) resistance is less than 60 Liters per minute)

AND

2.2.2 History of failure, contraindication or intolerance to ipratropium nebulized solution (generic Atrovent)

| Product Name: Lonhala Magnair, Yupleri | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | · |

Approval Criteria

1 - Documentation of positive clinical response to therapy

2. Revision History

| Date | Notes |
|------|-------|
| | |

| 3/11/2021 | Bulk copy C&S Arizona standard to Medicaid Arizona |
|-----------|--|
| | |

Lucemyra

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-104515 Lucemyra

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 3/9/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Lucemyra | | |
|--|---------------------|--|
| Approval Length | 14 Day(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - For symptoms of abrupt opioid withdrawal | | |
| | | |
| AND | | |
| | | |

2 - Opioids have been discontinued

AND

3 - BOTH of the following:

3.1 History of failure, contraindication, or intolerance to clonidine as verified by recent clonidine claims history in the past 180 days

AND

3.2 Lucemyra was initiated in the inpatient setting*

AND

4 - Prescriber must verify patient has been screened for hepatic and renal impairment and that dosing is appropriate for the patient's degree of hepatic and renal function

AND

5 - Prescriber must verify patient's vital signs have been monitored and that the patient is capable of and has been instructed on self-monitoring for hypotension, orthostasis, bradycardia, and associated symptoms

AND

6 - Patient does not have severe coronary insufficiency, a recent myocardial infarction, cerebrovascular disease, chronic renal failure, or marked bradycardia

AND

7 - Patient does not have congenital long QT syndrome

| Notes | *NOTE: Authorization will be issued for 14 days of therapy. If Lucemyr |
|-------|--|
| | a was initiated in the inpatient setting, the total course of therapy shou |
| | ld not exceed 14 days. |

| Date | Notes |
|----------|--------------------------------------|
| 3/9/2022 | Updated with patient safety criteria |

Lumizyme -Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99471 Lumizyme - Arizona

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Lumizyme | |
|--|---------------------|
| Diagnosis | Pompe disease |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of Pompe disease (acid alpha-glucosidase [GAA] deficiency) | |

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk copy C&S Arizona standard to Medicaid Arizona |

Lyrica

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-105529 Lyrica

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 4/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Brand Lyrica | |
|-----------------------------------|---------------------|
| Diagnosis | Seizure Disorder |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of seizure disorder | |

2 - History of failure, contraindication, or intolerance to generic pregabalin immediate-release capsules or generic pregabalin solution

| Product Name: Brand Lyrica | |
|----------------------------|---|
| Diagnosis | Neuropathic Pain Associated with Spinal Cord Injury |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of neuropathic pain associated with spinal cord injury

AND

2 - One of the following:

- History of failure to generic pregabalin immediate-release capsules or solution at a minimum dose of 300mg daily for 4 weeks
- Contraindication or intolerance to generic pregabalin immediate-release capsules or solution

| Product Name: Brand Lyrica | |
|----------------------------|---------------------|
| Diagnosis | Fibromyalgia |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of fibromyalgia

2 - One of the following:

- History of failure to generic pregabalin immediate-release capsules or solution at a minimum dose of 300mg daily for 4 weeks
- Contraindication or intolerance to generic pregabalin immediate-release capsules or solution

| Product Name: Brand Lyrica | |
|--------------------------------------|--|
| Diabetic peripheral neuropathy (DPN) | |
| 12 month(s) | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Diagnosis of diabetic peripheral neuropathy (DPN)

AND

2 - One of the following:

- History of failure to generic pregabalin immediate-release capsules or solution at a minimum dose of 300mg daily for 4 weeks
- Contraindication or intolerance to generic pregabalin immediate-release capsules or solution

| Product Name: Brand Lyrica | |
|----------------------------|-------------------------------|
| Diagnosis | Post herpetic neuralgia (PHN) |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of post herpetic neuralgia (PHN)

AND

- **2** One of the following:
 - History of failure to generic pregabalin immediate-release capsules or solution at a minimum dose of 300mg daily for 4 weeks
 - Contraindication or intolerance to generic pregabalin immediate-release capsules or solution

| Product Name: Lyrica CR | |
|-------------------------|--------------------------------------|
| Diagnosis | Diabetic peripheral neuropathy (DPN) |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of diabetic peripheral neuropathy (DPN)

AND

2 - History of failure, contraindication, or intolerance to gabapentin (generic Neurontin) at a minimum dose of 1800 milligrams daily for 4 weeks

AND

3 - History of failure, contraindication, or intolerance to treatment with ONE of the following:

- Tricyclic antidepressant at the maximum tolerated dose for 6 to 8 weeks, or intolerance to a tricyclic antidepressant
- Serotonin and norepinephrine reuptake inhibitor (SNRI) antidepressant (i.e. duloxetine, venlafaxine)

4 - History of failure, contraindication, or intolerance to generic pregabalin immediate-release capsules or generic pregabalin solution

| Product Name: Lyrica CR | |
|-------------------------|-------------------------------|
| Diagnosis | Post herpetic neuralgia (PHN) |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of post herpetic neuralgia (PHN)

AND

2 - History of failure, contraindication, or intolerance to gabapentin (generic Neurontin) at a minimum dose of 1800 milligrams daily for 4 weeks

AND

3 - History of failure, contraindication, or intolerance to a tricyclic antidepressant at the maximum tolerated dose for 6 to 8 weeks

AND

4 - History of failure, contraindication, or intolerance to generic pregabalin immediate-release capsules or generic pregabalin solution

| Date | Notes |
|------|-------|
| | |

| 3/31/2022 | Added step through generic for seizure indication. Updated all indicat |
|-----------|--|
| | ions to allow for any manufacturer of generic immediate-release caps |
| | ules or solution. |

Lysteda

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99473 Lysteda

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Lysteda, generic tranexamic acid | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of cyclic heavy menstrual bleeding | |

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk copy C&S Arizona standard to Medicaid Arizona |

Lyvispah (baclofen granules)

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-114468 | Lyvispah (baclofen granules) |
|-----------|------------------------------|
|-----------|------------------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Lyvispah | | |
|---|---------------------|--|
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Trial and failure, or intolerance to baclofen tablets | | |

| Date | Notes |
|-----------|-------------|
| 9/26/2022 | New program |

Makena- AZM

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-114911 Makena- AZM

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/5/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Makena*, generic hydroxyprogesterone caproate* | | | |
|---|---------------------|--|--|
| Guideline Type | Prior Authorization | | |
| | | | |
| Approval Criteria | Approval Criteria | | |
| 1 - Current singleton pregnancy | | | |
| | | | |
| AND | | | |
| | | | |
| 2 - History of a prior spontaneous preterm birth of a singleton pregnancy | | | |

3 - Treatment is initiated between 16 weeks, 0 days of gestation and 20 weeks, 6 days of gestation

AND

4 - Administration is to continue weekly until week 37 (through 36 weeks, 6 days) of gestation or delivery, whichever occurs first

AND

5 - Applies to generic hydroxyprogesterone caproate ONLY: patient has a history of failure, contraindication or intolerance to Brand Makena

| Notes | *NOTE: Approval duration is up to 21 weeks; approval duration should |
|-------|--|
| | take into account gestation week when Makena will be started and on |
| | ly authorized up to week 37. |

| Date | Notes |
|-----------|--|
| 10/4/2022 | Updated gestational days for drug initiation to align w PI |

Marinol, Syndros

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-108625 Marinol, Syndros

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 6/23/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Marinol, Syndros | | |
|--|--|--|
| Diagnosis | Chemotherapy-induced nausea and vomiting | |
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Patient is receiving cancer chemotherapy | | |

2 - ONE of the following:

2.1 History of failure, contraindication, or intolerance to formulary generic dronabinol

OR

2.2 Patient is unable to swallow capsules

AND

3 - History of failure, contraindication, or intolerance to a 5HT-3 (5-hydroxytriptamine) receptor antagonist [eg, Anzemet (dolasetron), Kytril (granisetron), or Zofran (ondansetron)]

AND

4 - History of failure, contraindication, or intolerance to ONE of the following:

- Ativan (lorazepam)
- Compazine (prochlorperazine)
- Decadron (dexamethasone)
- Haldol (haloperidol)
- Phenergan (promethazine)
- Reglan (metoclopramide)
- Zyprexa (olanzapine)

| Product Name: Generic Dronabinol | |
|----------------------------------|--|
| Diagnosis | Chemotherapy-induced nausea and vomiting |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Patient is receiving cancer chemotherapy

AND

2 - History of failure, contraindication, or intolerance to a 5HT-3 (5-hydoxytriptamine) receptor antagonist [eg, Anzemet (dolasetron), Kytril (granisetron), or Zofran (ondansetron)]

AND

3 - History of failure, contraindication, or intolerance to ONE of the following:

- Ativan (lorazepam)
- Compazine (prochlorperazine)
- Decadron (dexamethasone)
- Haldol (haloperidol)
- Phenergan (promethazine)
- Reglan (metoclopramide)
- Zyprexa (olanzapine)

| Product Name: Brand Marinol, Syndros | |
|--------------------------------------|--------------------------------|
| Diagnosis | Anorexia in Patients with AIDS |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of anorexia with weight loss in patients with AIDS (acquired immunodeficiency syndrome)

AND

2 - Patient is on antiretroviral therapy

AND

3 - ONE of the following:

3.1 Patient is 65 years of age or greater

OR

3.2 BOTH of the following:

- Patient is less than 65 years of age
- History of failure, contraindication, or intolerance to Megace (megestrol)

AND

4 - ONE of the following:

4.1 History of failure, contraindication, or intolerance to formulary generic dronabinol

OR

4.2 Patient is unable to swallow capsules

| Product Name: Generic dronabinol | |
|----------------------------------|--------------------------------|
| Diagnosis | Anorexia in Patients with AIDS |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of anorexia with weight loss in patients with AIDS (acquired immunodeficiency syndrome)

AND

2 - Patient is on antiretroviral therapy

| AND |
|--|
| 3 - ONE of the following: |
| 3.1 Patient is 65 years of age or greater |
| OR |
| 3.2 BOTH of the following: |
| Patient is less than 65 years of age History of failure, contraindication, or intolerance to Megace (megestrol) |

| Date | Notes |
|-----------|---|
| 6/23/2022 | Removed cesamet from guideline name. Added Brand Marinol as NP target |

Mavenclad - AZ

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99624 Wavenciad - AZ | GL-99624 | Mavenclad - AZ |
|-------------------------|----------|----------------|
|-------------------------|----------|----------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Mavenclad | |
|-------------------------|--|
| 2 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Diagnosis of relapsing form of multiple sclerosis (MS) (e.g., relapsing-remitting MS, secondary progressive MS with relapses)

2 - Prescribed by, or in consultation with, a specialist in the treatment of MS (e.g., neurologist)

AND

3 - ONE of the following:

3.1 Trial and failure (after trial of at least 4 weeks), contraindication, or intolerance to TWO of the following disease-modifying therapies for MS (document medication used, dose, and duration):

- Interferon beta-1a (Avonex, Rebif)
- Interferon beta-1b (Betaseron, Extavia)*
- Peginterferon beta-1a (Plegridy)
- Glatiramer acetate products (e.g., Copaxone, Glatopa)*
- A preferred dimethyl fumarate product (e.g., Tecfidera)
- Aubagio (teriflunomide)
- Gilenya (fingolimod)
- Mayzent (siponimod)
- Tysabri (natalizumab)**
- Ocrevus (ocrelizumab)**
- Lemtrada (alemtuzumab)**
- Zeposia (ozanimod)*
- Kesimpta (ofatumumab)*
- Bafiertam (monomethyl fumarate)*

OR

3.2 Patient is currently on Mavenclad

AND

4 - Patient is NOT receiving Mavenclad in combination with another disease modifying therapy [e.g., interferon beta preparations, glatiramer acetate products, Tecfidera (dimethyl fumarate), Tysabri (natalizumab), Gilenya (fingolimod), Mayzent (siponimod), Ocrevus (ocrelizumab), Lemtrada (alemtuzumab), or Aubagio (teriflunomide)]

| Notes | *Copaxone 40mg, Glatopa 20mg, glatiramer acetate, Bafiertam, Kesi |
|-------|--|
| | mpta, Zeposia, and Extavia are non-preferred and should not be inclu |

| ded in denial to provider. **Tysabri, Ocrevus, and Lemtrada are medic |
|---|
| al benefit and should not be included in denial to provider. |

| Product Name: Mavenclad | |
|-------------------------|---------------------|
| Approval Length | 2 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Mavenclad treatment

AND

2 - Patient is NOT receiving Mavenclad in combination with another disease modifying therapy [e.g., interferon beta preparations, glatiramer acetate products, Tecfidera (dimethyl fumarate), Tysabri (natalizumab), Gilenya (fingolimod), Mayzent (siponimod), Ocrevus (ocrelizumab), Lemtrada (alemtuzumab), or Aubagio (teriflunomide)]

AND

3 - Patient has not exceeded the FDA (Food and Drug Administration)-recommended limit of 2 treatment courses (4 treatment cycles) of Mavenclad

| Notes | Duration of coverage will be limited to 1 reauthorization to allow 2 cum |
|-------|--|
| | ulative treatment courses (4 treatment cycles) of Mavenclad therapy |

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Mepron

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99474 Mepron

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Mepron, generic atovaquone | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - ONE of the following: | |
| 1.1 BOTH of the following: | |

1.1.1 The patient has a diagnosis (e.g. human immunodeficiency virus [HIV]) warranting Pneumocystis jirovecii pneumonia (PCP) infection prophylaxis

1.1.2 The patient has a documented intolerance or contraindication to trimethoprimsulfamethoxazole (TMP-SMX) and dapsone

OR

1.2 BOTH of the following:

1.2.1 The patient has a diagnosis of mild to moderate pneumonia caused by P. jirovecii

AND

1.2.2 The patient has a documented intolerance, contraindication, or history of treatment failure to TMP-SMX

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk copy C&S Arizona standard to Medicaid Arizona |

Metformin products - AZM

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-115355 | Metformin | products - AZM |
|-----------|-----------|----------------|
|-----------|-----------|----------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/13/2022 |
|-----------------|------------|
|-----------------|------------|

1. Criteria

| Product Name: generic metformin 625 mg immediate-release tablets | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - History of greater than or equal to 12 week trial of preferred metformin immediate-release products

Product Name: generic metformin extended-release (generic for Fortamet and generic for Glumetza)

| Approval Length | 12 month(s) |
|-----------------|---------------------|
| Guideline Type | Prior Authorization |

Approval Criteria

1 - ALL of the following:

1.1 History of greater than or equal to 12 week trial of metformin extended-release (generic Glucophage XR)

AND

1.2 ONE of the following:

1.2.1 Submission of medical records (e.g. chart notes, laboratory values) documenting an inadequate response to metformin extended-release (generic Glucophage XR), in diabetic patients, as evidenced by the hemoglobin A1c level being above the patient's goal

OR

1.2.2 Submission of medical records (e.g. chart notes, laboratory values) documenting an intolerance to metformin extended-release (generic Glucophage XR) which is unable to be resolved with attempts to minimize the adverse effects where appropriate (e.g. dose reduction)

AND

1.3 History of greater than or equal to 12 week trial of metformin immediate-release

AND

1.4 One of the following:

1.4.1 Submission of medical records (e.g. chart notes, laboratory values) documenting an inadequate response to metformin immediate-release, in diabetic patients, as evidenced by the hemoglobin A1c level being above the patient's goal

OR

1.4.2 Submission of medical records (e.g. chart notes, laboratory values) documenting an intolerance to metformin immediate-release which is unable to be resolved with attempts to minimize the adverse effects where appropriate (e.g. dose reduction)

| Product Name: Brand Glumetza, Brand Fortamet | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - ALL of the following:

1.1 History of greater than or equal to 12 week trial of metformin extended-release (generic Glucophage XR)

AND

1.2 ONE of the following:

1.2.1 Submission of medical records (e.g. chart notes, laboratory values) documenting an inadequate response to metformin extended-release (generic Glucophage XR), in diabetic patients, as evidenced by the hemoglobin A1c level being above the patient's goal

OR

1.2.2 Submission of medical records (e.g. chart notes, laboratory values) documenting an intolerance to metformin extended-release (generic Glucophage XR) which is unable to be resolved with attempts to minimize the adverse effects where appropriate (e.g. dose reduction)

AND

1.3 History of greater than or equal to 12 week trial of metformin extended-release (generic Fortamet)

1.4 One of the following:

1.4.1 Submission of medical records (e.g. chart notes, laboratory values) documenting an inadequate response to metformin extended-release (generic Fortamet), in diabetic patients, as evidenced by the hemoglobin A1c level being above the patient's goal

OR

1.4.2 Submission of medical records (e.g. chart notes, laboratory values) documenting an intolerance to metformin extended-release (generic Fortamet) which is unable to be resolved with attempts to minimize the adverse effects where appropriate (e.g. dose reduction)

AND

1.5 History of greater than or equal to 12 week trial of metformin immediate-release

AND

1.6 One of the following:

1.6.1 Submission of medical records (e.g. chart notes, laboratory values) documenting an inadequate response to metformin immediate-release, in diabetic patients, as evidenced by the hemoglobin A1c level being above the patient's goal

OR

1.6.2 Submission of medical records (e.g. chart notes, laboratory values) documenting an intolerance to metformin immediate-release which is unable to be resolved with attempts to minimize the adverse effects where appropriate (e.g. dose reduction)

AND

1.7 Submission of article(s) published in the peer-reviewed medical literature showing that

the requested drug is likely to be more efficacious to this patient than metformin extendedrelease (generic Glucophage XR)

| Date | Notes |
|------------|---------------------------------------|
| 10/13/2022 | Removed Brand Glucophage XR as target |

Migranal

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99475 Migranal

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Migranal, Generic dihydroergotamine mesylate | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of migraine headaches with or without aura | |
| | |
| AND | |
| | |

2 - History of failure, contraindication, or intolerance to TWO preferred 5-HT1 (5hydroxytryptamine-1) receptor agonist (triptan) alternatives [eg, Imitrex (sumatriptan), Maxalt or Maxalt-MLT (rizatriptan)]

| Product Name: Brand I | Migranal*, Generic dihydroergotamine mesylate* |
|--|---|
| Approval Length | 12 month(s) |
| Guideline Type | Quantity Limit |
| | |
| Approval Criteria | |
| 1 - Diagnosis of migrain | ne headaches with or without aura |
| | AND |
| 2 - Prescribed by, or in | consultation with, ONE of the following: |
| NeurologistPain management specialist | |
| | AND |
| 3 - ONE of the following | g: |
| - | g prophylactic therapy with at least ONE of the following agents in wo or more migraines monthly: |
| Amitriptyline (E ONE of the follo timolol** | lavil) owing beta-blockers: atenolol, metoprolol, nadolol, propranolol, or |
| Divalproex sodium (Depakote/Depakote ER) Topiramate (Topamax) | |
| | fexor/Effexor XR) |
| | OR |
| 3.2 Patient has a contexperiencing two or mo | traindication or intolerance to ALL of the following, in patients ore migraines monthly: |

| timolol** | g beta-blockers: atenolol, metoprolol, nadolol, propranolol, or (Depakote/Depakote ER) ax) |
|---|---|
| | AND |
| 4 - BOTH of the following: | |
| 4.1 ONE of the following: | |
| 4.1.1 Higher dose or qua | ntity is supported by the manufacturer's prescribing information |
| | OR |
| 4.1.2 Higher dose or qua | ntity is supported by ONE of following compendia: |
| American Hospital I Micromedex DRUG Clinical pharmacolo | • |
| | OR |
| maximum doses as docum | evidence to support safety and additional efficacy at higher than ented in published biomedical literature demonstrating safety and greater than those approved by the Food and Drug Administration icated |
| | AND |
| 4.2 Physician acknowledg higher dose or quantity | es that the potential benefit outweighs the risk associated with the |
| ntly add | uantity requests exceeding the limited amount per month for freque occurring migraines will be approved by a clinical pharmacist. **N olol and timolol are non-preferred and should not be included in den to provider |
| | |

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk copy C&S Arizona standard to Medicaid Arizona |

Monurol

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-108624 Monurol

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 6/23/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Monurol | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - The provider has submitted labs showing the culture and sensitivity is positive for Monural and negative to Ciprofloxacin or Nitrofurantoin | |
| OR | |

2 - Trial and failure, contraindication, or intolerance to ONE of the following:

- •
- Ciprofloxacin Nitrofurantoin •

| Date | Notes |
|-----------|---|
| 6/23/2022 | Added product name to criteria section, no change to criteria |

Mozobil

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99625 Mozobil

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

| Product Name: Mozobil | | |
|--|---------------------|--|
| Approval Length | 4 Days* | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - ONE of the following: | | |
| Patients with non-Hodgkin's lymphoma (NHL) who will be undergoing autologous hematopoietic stem cell (HSC) transplantation | | |

| Patients with matrix transplantation | ultiple myeloma (MM) who will be undergoing autologous HSC |
|--|---|
| | AND |
| 2 - Used in combination (filgrastim)] | n with granulocyte-colony stimulating factor (G-CSF) [e.g., Zarxio |
| | AND |
| 3 - Prescribed by, or in | consultation with, a hematologist/oncologist |
| Notes | *Authorization will be issued for 1 course of therapy (up to four days of therapy). |

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

MS Agents - AZM

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-110769 MS Agents - AZM

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 8/15/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Gilenya, Brand Copaxone 20 mg, Brand Glatopa 40 mg, Avonex, Rebif, Betaseron, Extavia | | |
|---|------------------------------------|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | herapy Stage Initial Authorization | |
| Guideline Type | Prior Authorization | |
| Approval Criteria | | |

1 - Diagnosis of multiple sclerosis (MS)

| Product Name: GLATIRAMER 20mg, Brand GLATOPA 20mg, GLATIRAMER 40mg, Brand COPAXONE 40mg | | |
|---|---------------------------------------|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| Approval Criteria 1 - Diagnosis of multiple sclerosis (MS) | | |
| AND | | |
| 2 - Patient has a history of failure, contraindication, or intolerance to a trial of one of the preferred alternatives * NOTE: Drug May Require PA | | |
| Interferon Beta-1B (Extavia) Fingolimod (Gilenya) Interferon Beta-1A (Refib, Avonex) | | |
| AND | | |
| 3 - If the request is for GLATIRAMER 20mg or brand GLATOPA 20mg, patient must have tried and failed brand COPAXONE 20mg If the request is for GLATIRAMER 40mg or brand COPAXONE 40mg, patient must have tried and failed brand GLATOPA 40mg. | | |
| Notes | * Note: Preferred Drug may require PA | |

| Product Name: Tascenso ODT | |
|-----------------------------|-----------------------|
| Approval Length 12 month(s) | |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of multiple sclerosis (MS)

AND

2 - Patient has a history of failure, contraindication, or intolerance to Gilenya*

| Notes | * Note: Preferred Drug may require PA |
|-------|---------------------------------------|
|-------|---------------------------------------|

| Product Name: Vumerity, Bafiertam, Kesimpta, Tecfidera, Plegridy, Aubagio, Mayzent | |
|--|-----------------------|
| Approval Length 12 month(s) | |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of multiple sclerosis (MS)

AND

2 - Patient has a history of failure, contraindication, or intolerance to a trial of at least TWO of the preferred alternatives * NOTE: Drug May Require PA

- Interferon Beta-1B (Extavia)
- Fingolimod (Gilenya)
- Brand Copaxone 20mg
- Brand Glatopa 40mg
- Interferon Beta-1A (Refib, Avonex)

|--|

| Product Name: Gilenya, Brand Copaxone 20 mg, Brand Glatopa 40 mg, Avonex, Rebif, Betaseron, Extavia, GLATIRAMER 20mg, Brand GLATOPA 20mg, GLATIRAMER 40mg, Brand COPAXONE 40mg, Vumerity, Bafiertam, Kesimpta, Tecfidera, Plegridy, Aubagio, Mayzent, Tascenso ODT | |
|---|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to therapy

| Date | Notes |
|----------|---------------------------------|
| 8/4/2022 | Added Tascenso ODT as NP target |

Mulpleta

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99521 Mulpleta

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

| Product Name: Mulpleta | | |
|-----------------------------------|---------------------|--|
| Diagnosis | Thrombocytopenia | |
| Approval Length | 1 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of thrombocytopenia | | |

| | AND | | |
|---|---------------------------------------|--|--|
| 2 - Patient has chronic | 2 - Patient has chronic liver disease | | |
| | AND | | |
| 3 - Patient is scheduled to undergo a procedure | | | |
| | AND | | |
| 4 - History of failure, contraindication, or intolerance to the preferred alternatives Nyplate (romiplostim)* Promacta (eltrombopag olamine)* | | | |
| Notes | *Drugs may require PA | | |

| Date | Notes |
|-----------|-------------------------------------|
| 5/13/2021 | Arizona Medicaid 7.1 Implementation |

Multaq

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99476 Multaq

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

| Product Name: Multaq | | |
|--|---------------------|--|
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - ONE of the following: | | |
| 1.1 All of the following: | | |
| 1.1.1 Diagnosis of ONE of the following: | | |

| Paroxysmal Atrial Fibrillation (AF) Persistent AF defined as AF less than 6 months duration |
|--|
| AND |
| 1.1.2 ONE of the following: |
| Patient is in sinus rhythmPatient is planned to undergo cardioversion to sinus rhythm |
| AND |
| 1.1.3 Patient does not have New York Heart Association (NYHA) Class IV heart failure |
| AND |
| 1.1.4 Patient does not have symptomatic heart failure with recent decompensation requiring hospitalization |
| OR |
| 1.2 For continuation of current therapy |

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk copy C&S Arizona standard to Medicaid Arizona |

Myalept

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99626 Myalept

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Myalept | |
|-----------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of ONE of the following:

• Congenital generalized lipodystrophy associated with leptin deficiency

Acquired generalized lipodystrophy associated with leptin deficiency AND 2 - Used as an adjunct to diet modification AND 3 - Prescribed by an endocrinologist AND 4 - Documentation demonstrates that patient has at least ONE of the following: 4.1 Diabetes mellitus or insulin resistance with persistent hyperglycemia (hemoglobin A1C greater than 7.0%) despite BOTH of the following: Dietary intervention • Optimized insulin therapy at maximum tolerated doses • OR 4.2 Persistent hypertriglyceridemia (triglycerides greater than 250 milligrams per deciliter) despite BOTH of the following: Dietary intervention • Optimized therapy with at least two triglyceride-lowering agents from different classes • (e.g., fibrates, statins) at maximum tolerated doses

| Product Name: Myalept | |
|-----------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

| 1 - Documentation of positive clinical respo | nse to Myalept therapy |
|--|------------------------|
| | AND |
| 2 - Used as an adjunct to diet modification | |
| | AND |
| 3 - Prescribed by an endocrinologist | |

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Myfembree (relugolix, estradiol, and norethindrone acetate)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-114480 | Myfembree (relugolix, estradiol, and norethindrone acetate) |
|-----------|---|
|-----------|---|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Myfembree | |
|-------------------------|-----------------------|
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of heavy menstrual bleeding associated with uterine leiomyomas (fibroids)

AND AND Combination (estrogen/progestin) contraceptive • Progestins • Tranexamic acid • OR 3.2 Patient has had a previous interventional therapy to reduce bleeding AND 4 - Treatment duration of therapy has not exceeded a total of 24 months

AND

5 - Trial and failure, contraindication, or intolerance to Oriahnn

| Product Name: Myfembree | |
|-------------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

2 - Patient is premenopausal

3 - One of the following:

3.1 History of inadequate control of bleeding following a trial of at least 3 months, or history of intolerance or contraindication to one of the following:

1 - Patient has improvement in bleeding associated with uterine leiomyomas (fibroids) (e.g., significant/sustained reduction in menstrual blood loss per cycle, improved quality of life, etc.)

AND

2 - Treatment duration of therapy has not exceeded a total of 24 months

| Date | Notes |
|-----------|-------------|
| 9/26/2022 | New program |

Mytesi

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99477 Mytesi

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Mytesi | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of human immunodeficiency virus (HIV)/acquired immunodeficiency syndrome (AIDS) associated diarrhea | |

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk copy C&S Arizona standard to Medicaid Arizona |

Nadolol

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-109903 Nadolol

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 8/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Nadolol | |
|-----------------------|---|
| Diagnosis | PA required for patients 18 years of age or older |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - History of failure, contraindication, or intolerance to 3 of the following:

- atenolol
- atenolol/chlorthalidone
- bisoprolol fumarate

- bisoprolol/hydrochlorothiazide •
- carvedilol •
- labetalol HCI •
- metoprolol succinate •
- •
- metoprolol tartrate metoprolol/hydrochlorothiazide •
- propranolol HCI •
- propranolol/hydrochlorothiazide •
- sotalol HCI •

| Date | Notes |
|-----------|--|
| 7/28/2022 | Updated indication verbiage, no change to clinical criteria. |

Namzaric

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99478 Namzaric

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

| Product Name: Namzaric | |
|---------------------------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - BOTH of the following: | |
| 1.1 History of BOTH of the following: | |
| 1.1.1 Memantine (generic Namenda) | |

| ANI |) |
|--|-----------|
| 1.1.2 Donepezil (generic Aricept) | |
| | |
| ANI | |
| | |
| 1.2 Patient is stabilized on 10mg of donepezil o | nce daily |

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Standard to Medicaid Arizona Standard for 7 /1 go live |

Natpara

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99627 Natpara

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

| Product Name: Natpara | |
|---|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - ALL of the following: | |
| 1.1 Diagnosis of hypocalcemia resulting from chronic hypoparathyroidism | |

AND

1.2 25-hydroxy vitamin D level is above the lower limit of the normal laboratory reference range

AND

1.3 Patient is currently on active vitamin D (calcitriol) therapy

AND

1.4 Total serum calcium level (albumin corrected) is above 7.5 milligrams per deciliter

AND

2 - ONE of the following:

2.1 Patient is currently on calcium supplementation of 1-2 grams per day of elemental calcium in divided doses

OR

2.2 Patient has a contraindication to calcium supplementation

AND

- **3** Prescribed by ONE of the following:
 - Endocrinologist
 - Nephrologist

| Product Name: Natpara | |
|-----------------------|-------------|
| Approval Length | 12 month(s) |

| Therapy Stage | Reauthorization |
|--|---|
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Total serum calcium (approximately 8 to 9 m | n level (albumin corrected) within the lower half of the normal range nilligrams per deciliter) |
| | AND |
| 2 - Patient continues to take concomitant calcium supplementation that is sufficient to meet daily requirements | |
| | AND |
| 3 - Prescribed by ONE | of the following: |
| EndocrinologistNephrologist | |

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Nayzilam and Valtoco

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

| Product Name: Nayzilam | | |
|---------------------------|-----------------------|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of epilepsy | | |

AND

2 - Nayzilam is being prescribed for the acute treatment of intermittent, stereotypic episodes of frequent seizure activity that are distinct from a patient's usual seizure pattern

AND

3 - The prescriber provides a reason or special circumstance that precludes the use of diazepam rectal gel

| Product Name: Nayzilam | |
|------------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| | |

Approval Criteria

1 - Documentation of positive clinical response to therapy

| Product Name: Valtoco | |
|-----------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of epilepsy

AND

2 - Valtoco is being prescribed for the acute treatment of intermittent, stereotypic episodes of frequent seizure activity that are distinct from a patient's usual seizure pattern

AND

3 - The prescriber provides a reason or special circumstance that precludes the use of diazepam rectal gel

AND

4 - One of the following:

4.1 Patient is less than 12 years of age

OR

4.2 History of failure, contraindication, or intolerance to Nayzilam

| Product Name: Valtoco | |
|-----------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Documentation of positive clinical response to therapy

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona Standard to Medicaid Arizona Standard for 7 /1 go live |

Nexavar

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99758 Nexavar

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

| Product Name: Nexavar | |
|---|----------------------------|
| Diagnosis | Renal Cell Carcinoma (RCC) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of renal cell carcinoma (RCC) | |

AND

2 - ONE of the following:

2.1 Disease has relapsed

OR

2.2 BOTH of the following:

Medically or surgically unresectable tumor Diagnosis of Stage IV disease •

•

| Product Name: Nexavar | | |
|--|--------------------------|--|
| Diagnosis | Hepatocellular Carcinoma | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of hepato | ocellular carcinoma | |
| AND | | |
| 2 - ONE of the following: | | |
| 2.1 Patient has metastatic disease | | |
| | OR | |
| 2.2 Patient has extensive liver tumor burden | | |

OR

2.3 Patient is inoperable by performance status or comorbidity (local disease or local disease with minimal extrahepatic disease only)

OR

2.4 BOTH of the following:

- Patient is not a transplant candidate
- Disease is unresectable

| Product Name: Nexavar | |
|-----------------------|-----------------------|
| Diagnosis | Thyroid Cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - ONE of the following:

1.1 ALL of the following:

1.1.1 Diagnosis of ONE of the following:

- Follicular carcinoma
- Hürthle cell carcinoma
- Papillary carcinoma

AND

1.1.2 ONE of the following:

- Unresectable recurrent disease
- Persistent locoregional disease

| Metastatic disease |
|--|
| |
| AND |
| 1.1.3 ONE of the following: |
| Patient has symptomatic diseasePatient has progressive disease |
| AND |
| 1.1.4 Disease is refractory to radioactive iodine treatment |
| OR |
| 1.2 ALL of the following: |
| 1.2.1 Diagnosis of medullary thyroid carcinoma |
| AND |
| 1.2.2 ONE of the following: |
| Disease is progressive Disease is symptomatic with distant metastases |
| AND |
| 1.2.3 History of failure, contraindication, or intolerance to ONE of the following: |
| Caprelsa (vandetanib)Cometriq (cabozantinib) |

| Product Name: Nexavar | |
|-----------------------|---------------------|
| Diagnosis | Soft Tissue Sarcoma |

| Approval Length | 12 month(s) | | |
|--|---|--|--|
| Therapy Stage | Initial Authorization | | |
| Guideline Type | Prior Authorization | | |
| Approval Criteria | | | |
| | | | |
| 1 - Diagnosis of angios | arcoma | | |
| | OR | | |
| 2 - Diagnosis of desmo | 2 - Diagnosis of desmoid tumors / aggressive fibromatosis | | |
| | OR | | |
| 3 - BOTH of the following | ng: | | |
| 3.1 Diagnosis of progr | 3.1 Diagnosis of progressive gastrointestinal stromal tumors (GIST) | | |
| AND | | | |
| 3.2 History of failure, contraindication, or intolerance to ONE of the following: | | | |
| Gleevec (imatinib) Sutent (sunitinib) Stivarga (regorafenib) | | | |
| OR | | | |
| 4 - Diagnosis of solitary | / fibrous tumor/hemangiopericytoma | | |

| Product Name: Nexavar | |
|-----------------------|-----------------------|
| Diagnosis | Bone Cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

| Approval Criteria |
|---|
| 1 - BOTH of the following: |
| 1.1 Diagnosis of chordoma |
| AND |
| 1.2 Disease is recurrent |
| OR |
| 2 - BOTH of the following: |
| 2.1 ONE of the following: |
| Diagnosis of osteosarcoma Diagnosis of dedifferentiated chondrosarcoma Diagnosis of high-grade undifferentiated pleomorphic sarcoma (UPS) |
| AND |

2.2 Not used as first-line therapy

| Product Name: Nexavar | |
|-----------------------|------------------------|
| Diagnosis | Acute Myeloid Leukemia |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of acute myeloid leukemia (AML)

AND

2 - Patient has FLT3-ITD mutation-positive disease

AND

- **3** ONE of the following:
 - Patient has relapsed disease
 - Patient has refractory disease

AND

4 - Used in combination with ONE of the following:

- Vidaza (azacitidine)
- Dacogen (decitabine)

AND

5 - Patient is unable to tolerate more aggressive treatment regimens

| Product Name: Nexavar | | |
|-----------------------|-----------------------|--|
| Diagnosis | Ovarian Cancer | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

- **1** Diagnosis of ONE of the following:
 - Ovarian cancer
 - Fallopian tube cancer

| Primary peritoneal cancer | |
|--|--|
| AND | |
| 2 - ONE of the following: | |
| Patient has persistent diseasePatient has recurrent disease | |
| AND | |
| 3 - Disease is platinum-resistant | |
| AND | |
| 4 - Used in combination with topotecan | |

| Product Name: Nexavar | | |
|-----------------------|--|--|
| Diagnosis | Renal Cell Carcinoma (RCC), Hepatocellular Carcinoma, Thyroid Cancer, Soft Tissue Sarcoma, Bone Cancer, Acute Myeloid Leukemia, Ovarian Cancer | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |
| | | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Nexavar therapy

| Product Name: Nexavar | |
|-----------------------|--------------------------|
| Diagnosis | NCCN Recommended Regimen |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Nexavar will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Nexavar | |
|-----------------------|--------------------------|
| Diagnosis | NCCN Recommended Regimen |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Nexavar therapy

| Date | Notes |
|----------|-------------------------------------|
| 6/2/2021 | Arizona Medicaid 7.1 Implementation |

Nexiclon XR (clonidine ER)

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 6/1/2022 |
|-----------------|----------|
|-----------------|----------|

| Product Name: Nexiclon XR | |
|---|--|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| Approval Criteria 1 - Requested drug is to indication | being used for a Food and Drug Administration (FDA)-approved |
| AND | |

2 - Trial and failure, contraindication, or intolerance to one of the following (verified via paid pharmacy claims or submitted chart notes):

- generic clonidine oral tablet
- generic clonidine topical patch

| Date | Notes |
|-----------|-------------|
| 5/24/2022 | New Program |

Nexletol, Nexlizet

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99480 | Nexletol, Nexlizet |
|----------|--------------------|
|----------|--------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

| Product Name: Nexletol, Nexlizet | |
|---|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - ONE of the following diagnoses: | |
| Heterozygous familial hypercholesterolemia (HeFH) | |

• Atherosclerotic cardiovascular disease (ASCVD)

AND

2 - ONE of the following:

2.1 Patient has been receiving at least 12 consecutive weeks of high intensity statin therapy [i.e. atorvastatin 40-80 mg (milligrams), rosuvastatin 20-40 mg] and will continue to receive a high intensity statin at maximally tolerated dose

OR

2.2 BOTH of the following:

2.2.1 Patient is unable to tolerate high-intensity statin as evidenced by ONE of the following intolerable and persistent (i.e. more than 2 weeks) symptoms:

- Myalgia (muscle symptoms without CK [creatine kinase] elevations)
- Myositis (muscle symptoms with CK elevations less than 10 times upper limit of normal [ULN])

AND

2.2.2 ONE of the following:

2.2.2.1 Patient has been receiving at least 12 consecutive weeks of moderate- intensity statin therapy [i.e. atorvastatin 10-20 mg, rosuvastatin 5-10 mg, simvastatin greater than or equal to 20 mg, pravastatin greater than or equal to 40 mg, lovastatin 40 mg, Lescol XL (fluvastatin XL) 80 mg, fluvastatin 40 mg twice daily or Livalo (pitavastatin) greater than or equal to 2 mg] and will continue to receive a moderate-intensity statin at maximally tolerated dose

OR

2.2.2. Patient has been receiving at least 12 consecutive weeks of low-intensity statin therapy [i.e. simvastatin 10 mg, pravastatin 10-20 mg, lovastatin 20 mg, fluvastatin 20-40 mg, or Livalo (pitavastatin) 1 mg] statin therapy and will continue to receive a low-intensity statin at maximally tolerated dose

OR

2.3 Patient is unable to tolerate low or moderate-, and high-intensity statins as evidenced by ONE of the following:

2.3.1 ONE of the following intolerable and persistent (i.e. more than 2 weeks) symptoms for low or moderate-, and high-intensity statins:

- Myalgia (muscle symptoms without CK [creatine kinase] elevations)
- Myositis (muscle symptoms with CK elevations less than 10 times upper limit of normal [ULN])

OR

2.3.2 Patient has a labeled contraindication to all statins as documented in medical records

OR

2.3.3 Patient has experienced rhabdomyolysis or muscle symptoms with statin treatment with CK elevations greater than 10 times ULN

AND

3 - ONE of the following:

3.1 Documentation of ONE of the following LDL-C (low-density lipoprotein cholesterol) values while on maximally tolerated lipid lowering therapy within the last 120 days:

- LDL-C greater than or equal to 100 mg/dL (milligrams per deciliter) with ASCVD
- LDL-C greater than or equal to 130 mg/dL without ASCVD

OR

3.2 BOTH of the following:

3.2.1 Documentation of ONE of the following LDL-C values while on maximally tolerated lipid lowering therapy within the last 120 days:

- LDL-C between 70 mg/dL and 99 mg/dL with ASCVD
- LDL-C between 100 mg/dL and 129 mg/dL without ASCVD

3.2.2 Documentation of ONE of the following:

3.2.2.1 Patient has been receiving at least 12 consecutive weeks of ezetimibe (Zetia) therapy as adjunct to maximally tolerated statin therapy

OR

3.2.2.2 Patient has a history of contraindication, or intolerance to ezetimibe

| Product Name: Nexletol, Nexlizet | |
|----------------------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of a positive clinical response to therapy

AND

2 - Patient continues to receive statin at maximally tolerated dose (unless patient has documented inability to take statins)

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Standard to Medicaid Arizona Standard for 7 /1 go live |

Nityr- Arizona

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99628 Nityr- Arizona

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

| Product Name: Nityr | |
|--|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of hereditary tyrosinemia type 1 | |

2 - Prescriber provides a reason or special circumstance the patient cannot use Orfadin (nitisinone) capsules or suspension

| Product Name: Nityr | |
|---------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient shows evidence of positive clinical response (e.g. decrease in urinary/plasma succinylacetone and alpha-1-microglobulin levels) while on Nityr therapy

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Nocdurna

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99505 Nocdurna

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Nocdurna | |
|------------------------|--|
| 3 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Diagnosis of nocturia due to nocturnal polyuria (as defined by nighttime urine production that exceeds one-third of the 24-hour urine production)

2 - Patient wakes at least twice per night on a reoccurring basis to void

AND

3 - Documented serum sodium level is currently within normal limits of the normal laboratory reference range and has been within normal limits over the previous six months

AND

4 - The patient has been evaluated for other medical causes and has either not responded to, tolerated, or has a contraindication to treatments for identifiable medical causes [e.g., overactive bladder, benign prostatic hyperplasia/lower urinary tract symptoms (BPH/LUTS), elevated post-void residual urine, and heart failure]

AND

5 - Prescriber attests that the risks have been assessed and benefits outweigh the risks

| Product Name: Nocdurna | |
|------------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to therapy

AND

2 - Patient has routine monitoring for serum sodium levels

3 - Prescriber attests that the risks of hyponatremia have been assessed and benefits outweigh the risks

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk copy from C&S Medicaid to Arizona Medicaid for 7/1 eff |

Non-Preferred Drugs - Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-104403 Non-Preferred Drugs - Arizona

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 3/4/2022 |
|-----------------|----------|
|-----------------|----------|

| Product Name: Non-Preferred Drugs | | |
|--|----------------|--|
| Approval Length | 12 months* | |
| Guideline Type | Administrative | |
| | | |
| Approval Criteria | | |
| 1 - ALL of the following: | | |
| 1.1 ONE of the following: | | |
| If there are at least three preferred alternatives, history of trial per member's pharmacy claims resulting in a therapeutic failure, contraindication, or intolerance to at least | | |

THREE preferred alternatives (Prior trials of formulary/preferred drug list (PDL) alternatives must sufficiently demonstrate that the formulary/PDL alternatives are either ineffective or inappropriate at the time of the request)*

- If there are fewer than three preferred alternatives, the patient must have a history of trial per member's pharmacy claims resulting in a therapeutic failure, contraindication, or intolerance to ALL of the preferred products (Prior trials of formulary/preferred drug list (PDL) alternatives must sufficiently demonstrate that the formulary/PDL alternatives are either ineffective or inappropriate at the time of the request)*
- There are no preferred formulary alternatives for the requested drug*

AND

1.2 If the request is for a multi-source brand medication (i.e., MSC O) ONE of the following:

1.2.1 BOTH of the following:

- The brand is being requested because of an adverse reaction, allergy or sensitivity to the generic and the prescriber must attest to submitting the FDA MedWatch Form for allergic reactions to the medications.
- If there are generic product(s), the member has tried at least three (if available)

OR

1.2.2 ONE of the following:

- The brand is being requested due to a therapeutic failure with the generic (please provide reason for therapeutic failure).
- The brand is being requested because transition to the generic could result in destabilization of the patient (rationale must be provided)
- Special clinical circumstances exist that preclude the use of the generic equivalent of the multi-source brand medication for the patient (rationale must be provided)

AND

1.3 ONE of the following:

1.3.1 The requested drug must be used for a Food and Drug Administration (FDA)-approved indication.

1.3.2 The use of this drug is supported by information from ONE of the following appropriate compendia of current literature:

- The requested drug must be used for a Food and Drug Administration (FDA)-approved indication.
- Food and Drug Administration (FDA) approved indications and limits
- Published practice guidelines and treatment protocols
- Comparative data evaluating the efficacy, type and frequency of side effects and potential drug interactions among alternative products as well as the risks, benefits and potential member outcomes
- Drug Facts and Comparisons
- American Hospital Formulary Service Drug Information
- United States Pharmacopeia Drug Information
- DRUGDEX Information System
- UpToDate
- MicroMedex
- Peer-reviewed medical literature, including randomized clinical trials, outcomes, research data and pharmacoeconomic studies
- Other drug reference resources

AND

1.4 The drug is being prescribed for a medically accepted indication that is recognized as a covered benefit by the applicable health plan's program**

OR

2 - If the requested medication is a behavioral health medication, ONE of the following:

- The patient has been receiving treatment with the requested non-preferred behavioral health medication and is new to the plan (enrollment effective date within the past 90 days).
- The patient is currently receiving treatment with the requested non-preferred behavioral health medication in the hospital and must continue upon discharge.

| Notes | *Anti-infectives: Approve for the requested time frame, or if duration is not specified approve the request for 30 days. *Controlled Substance s shall be approved for the requested time. If there is not a requested time period and it is not clear in the directions, approve for one time o nly. *Other medications: Approved for the requested time frame, or if d uration is not specified, approve for 12 months. * For Non-Preferred G enerics (i.e. MSC=Y) approvals: Please approve at MSC=Y only. For preferred alternatives, use the non-preferred alternatives grid to identif y appropriate alternatives: https://uhgazure.sharepoint.com/sites/CST/ |
|-------|---|
|-------|---|

| | CSDM/Shared%20Documents/Forms/AllItems.aspx?FolderCTID=0x0 1200027C80175A8369D44AC45A99A99328B80&View=%7B4B6D25 AD%2D6A95%2D496D%2D9937%2D65CECD43AFE7%7D&viewid= c2ad0afa%2D814c%2D499e%2Dbf25%2D3411fac9171f&id=%2Fsite s%2FCST%2FCSDM%2FShared%20Documents%2FAZM%2FNF%2 0Alt%20Tables **Medications used solely for anti-obesity/weight loss, cosmetic (e.g., alopecia, actinic keratosis, vitiligo), erectile dysfunction , or sexual dysfunction purposes are NOT medically accepted indicati ons and are NOT recognized as a covered benefit. Erectile dysfunction n drugs (Cialis/Tadalafil) are covered for clinical diagnoses other than ED. |
|--|---|
|--|---|

| Date | Notes |
|----------|--|
| 3/4/2022 | Updated MSC criterion verbiage. Attached to Specialty formulary. |

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-104403 Non-Preferred Drugs - Arizona

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 3/4/2022 |
|-----------------|----------|
|-----------------|----------|

| Product Name: Non-Preferred Drugs | | | |
|--|----------------|--|--|
| Approval Length | 12 months* | | |
| Guideline Type | Administrative | | |
| | | | |
| Approval Criteria | | | |
| 1 - ALL of the following: | | | |
| 1.1 ONE of the following: | | | |
| If there are at least three preferred alternatives, history of trial per member's pharmacy claims resulting in a therapeutic failure, contraindication, or intolerance to at least THREE preferred alternatives (Prior trials of formulary/preferred drug list (PDL) | | | |

alternatives must sufficiently demonstrate that the formulary/PDL alternatives are either ineffective or inappropriate at the time of the request)*

- If there are fewer than three preferred alternatives, the patient must have a history of trial per member's pharmacy claims resulting in a therapeutic failure, contraindication, or intolerance to ALL of the preferred products (Prior trials of formulary/preferred drug list (PDL) alternatives must sufficiently demonstrate that the formulary/PDL alternatives are either ineffective or inappropriate at the time of the request)*
- There are no preferred formulary alternatives for the requested drug*

AND

1.2 If the request is for a multi-source brand medication (i.e., MSC O) ONE of the following:

1.2.1 BOTH of the following:

- The brand is being requested because of an adverse reaction, allergy or sensitivity to the generic and the prescriber must attest to submitting the FDA MedWatch Form for allergic reactions to the medications.
- If there are generic product(s), the member has tried at least three (if available)

OR

1.2.2 ONE of the following:

- The brand is being requested due to a therapeutic failure with the generic (please provide reason for therapeutic failure).
- The brand is being requested because transition to the generic could result in destabilization of the patient (rationale must be provided)
- Special clinical circumstances exist that preclude the use of the generic equivalent of the multi-source brand medication for the patient (rationale must be provided)

AND

1.3 ONE of the following:

1.3.1 The requested drug must be used for a Food and Drug Administration (FDA)-approved indication.

1.3.2 The use of this drug is supported by information from ONE of the following appropriate compendia of current literature:

- The requested drug must be used for a Food and Drug Administration (FDA)-approved indication.
- Food and Drug Administration (FDA) approved indications and limits
- Published practice guidelines and treatment protocols
- Comparative data evaluating the efficacy, type and frequency of side effects and potential drug interactions among alternative products as well as the risks, benefits and potential member outcomes
- Drug Facts and Comparisons
- American Hospital Formulary Service Drug Information
- United States Pharmacopeia Drug Information
- DRUGDEX Information System
- UpToDate
- MicroMedex
- Peer-reviewed medical literature, including randomized clinical trials, outcomes, research data and pharmacoeconomic studies
- Other drug reference resources

AND

1.4 The drug is being prescribed for a medically accepted indication that is recognized as a covered benefit by the applicable health plan's program**

OR

2 - If the requested medication is a behavioral health medication, ONE of the following:

- The patient has been receiving treatment with the requested non-preferred behavioral health medication and is new to the plan (enrollment effective date within the past 90 days).
- The patient is currently receiving treatment with the requested non-preferred behavioral health medication in the hospital and must continue upon discharge.

| not sp s shal time p nly. *C uration eneric prefer | nfectives: Approve for the requested time frame, or if duration is becified approve the request for 30 days. *Controlled Substance I be approved for the requested time. If there is not a requested eriod and it is not clear in the directions, approve for one time o Other medications: Approved for the requested time frame, or if d in is not specified, approve for 12 months. * For Non-Preferred G is (i.e. MSC=Y) approvals: Please approve at MSC=Y only. For red alternatives, use the non-preferred alternatives grid to identif ropriate alternatives: https://uhgazure.sharepoint.com/sites/CST/ |
|--|--|
|--|--|

| | CSDM/Shared%20Documents/Forms/AllItems.aspx?FolderCTID=0x0 1200027C80175A8369D44AC45A99A99328B80&View=%7B4B6D25 AD%2D6A95%2D496D%2D9937%2D65CECD43AFE7%7D&viewid= c2ad0afa%2D814c%2D499e%2Dbf25%2D3411fac9171f&id=%2Fsite s%2FCST%2FCSDM%2FShared%20Documents%2FAZM%2FNF%2 0Alt%20Tables **Medications used solely for anti-obesity/weight loss, cosmetic (e.g., alopecia, actinic keratosis, vitiligo), erectile dysfunction , or sexual dysfunction purposes are NOT medically accepted indicati ons and are NOT recognized as a covered benefit. Erectile dysfunction n drugs (Cialis/Tadalafil) are covered for clinical diagnoses other than ED. |
|--|---|
|--|---|

| Date | Notes |
|----------|--|
| 3/4/2022 | Updated MSC criterion verbiage. Attached to Specialty formulary. |

Non-Preferred Prenatal Vitamins

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99528 Non-Preferred Prenatal Vitamins

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

| Product Name: Non-Preferred Prenatal Vitamins | | | | | |
|--|--|--|--|--|--|
| Approval Length | 12 month(s) | | | | |
| Guideline Type | Prior Authorization | | | | |
| | | | | | |
| Approval Criteria | | | | | |
| 1 - History of failure, contraindication, or intolerance to ALL of the following preferred products:* | | | | | |
| Notes | *Please refer to the background table for the alternatives | | | | |

2. Background

۱L

| Benefit/Coverage/Program Information | | | |
|--------------------------------------|-----------------|-----------------------|--|
| Preferred Products: | | | |
| GPI-14 | Product ID | Product Label | GPI-14 Description |
| 785120000003 15 | 7331710500 9 | PRENATVITE TA B RX | *PRENATAL MULTIVITAMINS & MINERALS W/IRON & FA TAB 0.8 MG*** |
| 785120100003 | 6954302679 | PNV TABS TAB | *PRENATAL VIT W/ IRON |
| 30 | 0 | 29-1MG | CARBONYL-FA TAB 29-1 MG*** |
| 785120100003 | 6025801930 | PRENATABS RX | *PRENATAL VIT W/ IRON |
| 30 | 9 | TAB | CARBONYL-FA TAB 29-1 MG*** |
| 785120100003 | 4293707051 | PRENATAL+FE T | *PRENATAL VIT W/ IRON |
| 30 | 0 | AB 29-1MG | CARBONYL-FA TAB 29-1 MG*** |
| 785120100003 | 4293707051 | PRENATAL+FE T | *PRENATAL VIT W/ IRON |
| 30 | 6 | AB 29-1MG | CARBONYL-FA TAB 29-1 MG*** |
| 785120100003 | 4293707051 | PRENATAL+FE T | *PRENATAL VIT W/ IRON |
| 30 | 8 | AB 29-1MG | CARBONYL-FA TAB 29-1 MG*** |
| 785120100003 | 5865701339 | THRIVITE RX TAB | *PRENATAL VIT W/ IRON |
| 30 | 0 | 29-1MG | CARBONYL-FA TAB 29-1 MG*** |
| 785120100003 | 7118600192 | VIL-RX TAB 29- | *PRENATAL VIT W/ IRON |
| 30 | 4 | 1MG | CARBONYL-FA TAB 29-1 MG*** |
| 785120100003 | 1381105169 | VOL-TAB RX TAB | *PRENATAL VIT W/ IRON |
| 30 | 0 | | CARBONYL-FA TAB 29-1 MG*** |
| 785120100003 | 1381100271 | ELITE-OB TAB | *PRENATAL VIT W/ IRON |
| 52 | 0 | | CARBONYL-FA TAB 50-1.25 MG*** |
| 785120100003 | 6802500101 | OB | *PRENATAL VIT W/ IRON |
| 52 | 0 | COMPLETE TAB | CARBONYL-FA TAB 50-1.25 MG*** |
| 785120150003 | 5865701700 | M-NATAL PLUS | *PRENATAL VIT W/ FE |
| 24 | 1 | TAB | FUMARATE-FA TAB 27-1 MG*** |
| 785120150003 | 1283008000 | M-VIT TAB 27- | *PRENATAL VIT W/ FE |
| 24 | 1 | 1MG | FUMARATE-FA TAB 27-1 MG*** |
| 785120150003 | 7089802200 | NEONATAL TAB | *PRENATAL VIT W/ FE |
| 24 | 1 | COMPLTE | FUMARATE-FA TAB 27-1 MG*** |

| 785120150003 | 7089801150 | NEONATAL PLS | *PRENATAL VIT W/ FE |
|--------------------|-----------------|-------------------------------|---|
| 24 | 1 | TAB 27-1MG | FUMARATE-FA TAB 27-1 MG*** |
| 785120150003 | 7583400500 | NIVA-PLUS TAB | *PRENATAL VIT W/ FE |
| 24 | 1 | | FUMARATE-FA TAB 27-1 MG*** |
| 785120150003 | 0081393160 | O-CAL FA TAB | *PRENATAL VIT W/ FE |
| 24 | 1 | | FUMARATE-FA TAB 27-1 MG*** |
| 785120150003 | 7139962460 | ONE VITE TAB | *PRENATAL VIT W/ FE |
| 24 | 9 | 1MG PLUS | FUMARATE-FA TAB 27-1 MG*** |
| 785120150003 | 3932801061 | PRENATAL TAB | *PRENATAL VIT W/ FE |
| 24 | 0 | 27-1MG | FUMARATE-FA TAB 27-1 MG*** |
| 785120150003 | 3932801065 | PRENATAL TAB | *PRENATAL VIT W/ FE |
| 24 | 0 | 27-1MG | FUMARATE-FA TAB 27-1 MG*** |
| 785120150003 | 6304401500 | PRENATAL VIT | *PRENATAL VIT W/ FE |
| 24 | 1 | TAB LOW IRON | FUMARATE-FA TAB 27-1 MG*** |
| 785120150003 | 6304401500 | PRENATAL VIT | *PRENATAL VIT W/ FE |
| 24 | 5 | TAB LOW IRON | FUMARATE-FA TAB 27-1 MG*** |
| 785120150003 | 6954302581 | PREPLUS TAB | *PRENATAL VIT W/ FE |
| 24 | 0 | 27-1MG | FUMARATE-FA TAB 27-1 MG*** |
| 785120150003 | 6954302585 | PREPLUS TAB | *PRENATAL VIT W/ FE |
| 24 | 0 | 27-1MG | FUMARATE-FA TAB 27-1 MG*** |
| 785120150003 | 6711201010 | TRICARE TAB | *PRENATAL VIT W/ FE |
| 24 | 0 | PRENATAL | FUMARATE-FA TAB 27-1 MG*** |
| 785120150003 | 1713908003 | VITATHELY TAB | *PRENATAL VIT W/ FE |
| 24 | 0 | | FUMARATE-FA TAB 27-1 MG*** |
| 785120150003 | 1381105191 | VOL-PLUS TAB | *PRENATAL VIT W/ FE |
| 24 | 0 | | FUMARATE-FA TAB 27-1 MG*** |
| 785120150003 | 1381105195 | VOL-PLUS TAB | *PRENATAL VIT W/ FE |
| 24 | 0 | | FUMARATE-FA TAB 27-1 MG*** |
| 785120150003 24 | 6936702670 1 | WESTAB PLUS TAB 27- 1MG | *PRENATAL VIT W/ FE FUMARATE-FA TAB 27-1 MG*** |
| 785120150003 | 6025801920 | TRINATE TAB | *PRENATAL VIT W/ FE |
| 29 | 1 | | FUMARATE-FA TAB 28-1 MG*** |
| 785120150003 | 1381105141 | VOL-NATE TAB | *PRENATAL VIT W/ FE |
| 29 | 0 | | FUMARATE-FA TAB 28-1 MG*** |

| 705400450000 | 4000700700 | | |
|--------------|------------|----------------|----------------------------|
| 785120150003 | 1026722700 | CO-NATAL | *PRENATAL VIT W/ FE |
| 32 | 1 | FA TAB 29-1MG | FUMARATE-FA TAB 29-1 MG*** |
| 785120150003 | 7331782860 | NEONATAL TAB | *PRENATAL VIT W/ FE |
| 32 | 1 | COMPLETE | FUMARATE-FA TAB 29-1 MG*** |
| 02 | • | | |
| 785120150003 | 6954302591 | PRETAB TAB | *PRENATAL VIT W/ FE |
| 32 | 0 | 29-1MG | FUMARATE-FA TAB 29-1 MG*** |
| 705400450000 | 4004400074 | | |
| 785120150003 | 1381100071 | | *PRENATAL VIT W/ FE |
| 60 | 0 | RX TAB 1 | FUMARATE-FA TAB 60-1 MG*** |
| 785120150003 | 5199105660 | VINATE ONE TAB | *PRENATAL VIT W/ FE |
| 60 | 1 | | FUMARATE-FA TAB 60-1 MG*** |
| 00 | • | | |
| 785120150003 | 5860708112 | MYNATAL PLUS | *PRENATAL VIT W/ FE |
| 66 | 0 | TAB | FUMARATE-FA TAB 65-1 MG*** |
| 705400450000 | 5000704050 | | |
| 785120150003 | 5860701056 | MYNATAL-Z TAB | *PRENATAL VIT W/ FE |
| 66 | 5 | | FUMARATE-FA TAB 65-1 MG*** |
| 785120150003 | 0064200791 | VITAFOL-OB TAB | *PRENATAL VIT W/ FE |
| | 2 | 65-1MG | FUMARATE-FA TAB 65-1 MG*** |
| 00 | - | | |
| 785120150005 | 1381100149 | COMPLETENATE | *PRENATAL VIT W/ FE |
| 30 | 0 | CHW | FUMARATE-FA CHEW TAB 29-1 |
| | | | MG*** |
| 705400450005 | 4000707074 | | |
| | 4293707071 | PRENATAL | *PRENATAL VIT W/ FE |
| 30 | 0 | 19 CHW 29-1MG | FUMARATE-FA CHEW TAB 29-1 |
| | | | MG*** |
| 785120150005 | 4293707071 | PRENATAL | *PRENATAL VIT W/ FE |
| 30 | 6 | 19 CHW 29-1MG | FUMARATE-FA CHEW TAB 29-1 |
| | - | | MG*** |
| | | | |
| 785120150005 | 4293707071 | PRENATAL | *PRENATAL VIT W/ FE |
| 30 | 8 | 19 CHW 29-1MG | FUMARATE-FA CHEW TAB 29-1 |
| | | | MG*** |
| 705400450005 | 0005004070 | DDENATAL | |
| 785120150005 | 6025801970 | | *PRENATAL VIT W/ FE |
| 30 | 1 | 19 CHW TAB | FUMARATE-FA CHEW TAB 29-1 |
| | | | MG*** |
| 785120150005 | 1392501170 | SE-NATAL | *PRENATAL VIT W/ FE |
| 30 | 1 | 19 CHW | FUMARATE-FA CHEW TAB 29-1 |
| | | | MG*** |
| | | | |

| 785120160001 30 | 1381100493 0 | ULTIMATECARE CAP ONE | *PRENATAL VIT W/ FE CBN-FE ASP GLYC-FA-OMEGA 3 CAP 27- 1MG*** |
|--------------------|-----------------|--------------------------------|--|
| 785120180001 16 | 2335901053 0 | C-NATE DHA CAP 28-1- 200 | *PRENATAL VIT W/ FE FUM-FA- OMEGA 3 CAP 28-1-200 MG*** |
| 785120180001 | 2335902003 | RELNATE | *PRENATAL VIT W/ FE FUM-FA- |
| 16 | 0 | DHA CAP | OMEGA 3 CAP 28-1-200 MG*** |
| 785120180001 | 6954303703 | VIRT-NATE CAP | *PRENATAL VIT W/ FE FUM-FA- |
| 16 | 0 | DHA | OMEGA 3 CAP 28-1-200 MG*** |
| 785120180001 | 6466100803 | VIVA DHA CAP | *PRENATAL VIT W/ FE FUM-FA- |
| 16 | 0 | | OMEGA 3 CAP 28-1-200 MG*** |
| 785120220003 20 | 6954302419 0 | VIRT-PN TAB | *PRENATAL VIT W/ FE FUM- METHYLFOLATE-FA TAB 27-0.6- 0.4 MG*** |
| 785120460003 30 | 5549501250 1 | ATABEX OB TAB 29-1MG | *PRENATAL VIT W/ FE BISGLYCINATE CHELATE-FA TAB 29-1 MG*** |
| 785120460003 30 | 5199101780 1 | VINATE II TAB | *PRENATAL VIT W/ FE BISGLYCINATE CHELATE-FA TAB 29-1 MG*** |
| 785120510003 27 | 0017808589 0 | CITRANATAL TA B RX | *PRENATAL W/O A W/ FE CARBONYL-FE GLUC-DSS-FA TAB 27-1MG*** |
| 785120580001 | 5274706203 | CONCEPT | *PRENATAL W/O A W/FE FUM-FE |
| 50 | 0 | OB CAP | POLY-FA CAP 130-92.4-1 MG*** |
| 785120580001 | 1381105353 | FOLIVANE- | *PRENATAL W/O A W/FE FUM-FE |
| 50 | 0 | OB CAP | POLY-FA CAP 130-92.4-1 MG*** |
| 785120600003 | 5199101550 | VINATE M TAB | *PRENATAL VIT W/ SEL-FE |
| 25 | 1 | | FUMARATE-FA TAB 27-1 MG*** |
| 785120700003 | 4293707061 | PRENATAL | *PRENATAL VIT W/ DSS-FE |
| 30 | 0 | 19 TAB 29-1MG | FUMARATE-FA TAB 29-1 MG*** |
| 785120700003 | 4293707061 | PRENATAL | *PRENATAL VIT W/ DSS-FE |
| 30 | 6 | 19 TAB 29-1MG | FUMARATE-FA TAB 29-1 MG*** |
| 785120700003 | 4293707061 | PRENATAL | *PRENATAL VIT W/ DSS-FE |
| 30 | 8 | 19 TAB 29-1MG | FUMARATE-FA TAB 29-1 MG*** |

| 785120700003 | 1392501160 | SE-NATAL 19 TAB | *PRENATAL VIT W/ DSS-FE |
|--------------------|-----------------|------------------------|--|
| 30 | 1 | | FUMARATE-FA TAB 29-1 MG*** |
| 785120910001 | 5274706213 | CONCEPT | *PRENATAL W/FE FUM-FE POLY - |
| 35 | 0 | DHA CAP | FA-OMEGA 3 CAP 53.5-38-1 MG*** |
| 785120910001 | 5865701213 | DOTHELLE DHA | *PRENATAL W/FE FUM-FE POLY - |
| 35 | 0 | CAP | FA-OMEGA 3 CAP 53.5-38-1 MG*** |
| 785120910001 | 1381105363 | TARON-C | *PRENATAL W/FE FUM-FE POLY - |
| 35 | 0 | DHA CAP | FA-OMEGA 3 CAP 53.5-38-1 MG*** |
| 785120910001 | 7643903313 | VIRT-C DHA CAP | *PRENATAL W/FE FUM-FE POLY - |
| 35 | 0 | | FA-OMEGA 3 CAP 53.5-38-1 MG*** |
| 785160200063 30 | 0064200763 0 | VITAFOL-OB PAK +DHA | *PRENATAL MV W/FE FUM-FA TAB 65-1 MG & DHA CAP 250 MG PACK * |
| 785160320001 30 | 0064200703 0 | VITAFOL- ONE CAP | *PRENATAL MV W/ FE POLYSAC CMPLX-FA-DHA CAP 29-1-200 MG*** |
| 785160320063 25 | 0064200753 0 | SELECT- OB+ PAK DHA | *PRENATAL MV W/FE POLY-FA CHW 29-1 MG & DHA CAP 250 MG PAK * |

| Date | Notes |
|-----------|-------------------------------------|
| 5/18/2021 | Arizona Medicaid 7.1 Implementation |

Norliqva (amlodipine oral solution)

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-107420 Norliqva (amlodipine oral solution)

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 5/25/2022 |
|-----------------|-----------|
|-----------------|-----------|

| Product Name: Norliqva | | | |
|---|---------------------|--|--|
| Approval Length | th 12 month(s) | | |
| Guideline Type | Prior Authorization | | |
| Approval Criteria 1 - Requested drug is being used for a Food and Drug Administration (FDA)-approved indication | | | |
| AND | | | |

2 - One of the following:

2.1 Trial and failure, contraindication, or intolerance to generic amlodipine tablets (verified via paid pharmacy claims or submitted chart notes)

OR

2.2 One of the following:

- ٠
- Patient is 8 years of age or younger Patient is unable to swallow oral tablets/capsules •

| Date | Notes |
|-----------|-------------|
| 5/23/2022 | New Program |

Northera

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99629 | Northera |
|----------|----------|
| | |

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: |
|-----------------|
|-----------------|

1. Criteria

| Product Name: Northera | |
|------------------------|-----------------------|
| Approval Length | 3 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of symptomatic neurogenic orthostatic hypotension (nOH) as defined by ONE of the following when an upright position is assumed or when using a head-up tilt-table testing at an angle of at least 60 degrees:

• At least a 20 millimeters of mercury (mm Hg) fall in systolic pressure

| | • | At least a | 10 mm | Hg fall in | diastolic | pressure |
|--|---|------------|-------|------------|-----------|----------|
|--|---|------------|-------|------------|-----------|----------|

2 - nOH caused by ONE of the following:

- Primary autonomic failure (e.g., Parkinson's disease, multiple system atrophy, and pure autonomic failure)
- Dopamine beta-hydroxylase deficiency
- Non-diabetic autonomic neuropathy

AND

3 - Diagnostic evaluation has excluded other causes associated with orthostatic hypotension (e.g., congestive heart failure, fluid restriction, malignancy)

AND

- 4 The patient has tried at least TWO of the following non-pharmacologic interventions:
 - Discontinuation of drugs which can cause orthostatic hypotension [e.g., diuretics, antihypertensive medications (primarily sympathetic blockers), anti-anginal drugs (nitrates), alpha-adrenergic antagonists, and antidepressants]
 - Raising the head of the bed 10 to 20 degrees
 - Compression garments to the lower extremities or abdomen
 - Physical maneuvers to improve venous return (e.g., regular modest-intensity exercise)
 - Increased salt and water intake, if appropriate
 - Avoiding precipitating factors (e.g., overexertion in hot weather, arising too quickly from supine to sitting or standing)

AND

5 - No previous diagnosis of supine hypertension

AND

- 6 Prescribed by, or in consultation with, ONE of the following specialists:
 - Cardiologist

- Neurologist
- Nephrologist

7 - History of failure (after a trial of at least 30 days), contraindication or intolerance to BOTH of the following medications:

- Florinef (fludrocortisone)
- ProAmatine (midodrine)

| Product Name: Northera | |
|------------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Northera therapy

AND

2 - Physiological countermeasures for neurogenic orthostatic hypotension (nOH) continue to be employed

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Nourianz

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99481 Nourianz

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

| Product Name: Nourianz | | |
|--------------------------------------|-----------------------|--|
| Approval Length | 6 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of Parkinson's disease | | |

2 - Used as adjunctive treatment to levodopa/carbidopa in patients experiencing "off" episodes

AND

3 - History of failure, contraindication, or intolerance to TWO anti-Parkinson's disease therapies from the following adjunctive pharmacotherapy classes (trial must be from two different classes):

- Dopamine agonists (e.g., pramipexole, ropinirole)
- Catechol-O-methyl transferase (COMT) inhibitors (e.g., entacapone)
- Monoamine oxidase (MAO) B inhibitors (e.g., rasagiline, selegiline)

| Product Name: Nourianz | | |
|------------------------|---------------------|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Documentation of positive clinical response to Nourianz therapy

AND

2 - Patient will continue to receive treatment with a carbidopa/levodopa-containing medication

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona Standard to Medicaid Arizona Standard for 7 /1 go live |

Nucala (mepolizumab)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 7/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Nucala | | |
|----------------------|-----------------------|--|
| Diagnosis | Severe Asthma | |
| Approval Length | 6 Months [G] | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of severe asthma

2 - Asthma is an eosinophilic phenotype as defined by one of the following:

- Baseline (pre-treatment) peripheral blood eosinophil level is greater than or equal to 150 cells/microliter
- Peripheral blood eosinophil levels were greater than or equal to 300 cells/microliter within the past 12 months

AND

3 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting one of the following:

3.1 Patient has had at least two or more asthma exacerbations requiring systemic corticosteroids (e.g., prednisone) within the past 12 months

OR

3.2 Prior asthma-related hospitalization within the past 12 months

AND

4 - Patient is currently being treated with one of the following unless there is a contraindication or intolerance to these medications (verified via paid pharmacy claims):

4.1 Both of the following:

- High-dose inhaled corticosteroid (ICS) (e.g., greater than 500 mcg fluticasone propionate equivalent/day)
- Additional asthma controller medication (e.g., leukotriene receptor antagonist [e.g., montelukast], long-acting beta-2 agonist [LABA] [e.g., salmeterol], tiotropium)

OR

4.2 One maximally-dosed combination ICS/LABA product (e.g., Advair [fluticasone propionate/salmeterol], Symbicort [budesonide/formoterol], Breo Ellipta [fluticasone/vilanterol])

5 - Age greater than or equal to 6 years

AND

6 - Prescribed by or in consultation with one of the following:

- Pulmonologist
- Allergist/Immunologist

| Product Name: Nucala | |
|----------------------|---------------------|
| Diagnosis | Severe Asthma |
| Approval Length | 12 Months |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting positive clinical response to therapy (e.g., reduction in exacerbations, improvement in forced expiratory volume in 1 second [FEV1], decreased use of rescue medications) [C]

AND

2 - Patient continues to be treated with an inhaled corticosteroid (ICS) (e.g., fluticasone, budesonide) with or without additional asthma controller medication (e.g., leukotriene receptor antagonist [e.g., montelukast], long-acting beta-2 agonist [LABA] [e.g., salmeterol], tiotropium) unless there is a contraindication or intolerance to these medications (verified via paid pharmacy claims)

AND

3 - Prescribed by or in consultation with one of the following:

- Pulmonologist
- Allergist/Immunologist

| Product Name: Nucala | | |
|----------------------|---|--|
| Diagnosis | Chronic rhinosinusitis with nasal polyps (CRSwNP) | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of chronic rhinosinusitis with nasal polyps (CRSwNP)

AND

2 - Unless contraindicated, the patient has had an inadequate response to 2 months of treatment with an intranasal corticosteroid (e.g., fluticasone, mometasone)

AND

3 - Used in combination with another agent for CRSwNP

AND

4 - Prescribed by or in consultation with one of the following:

- Allergist/Immunologist
- Otolaryngologist
- Pulmonologist

| Product Name: Nucala | |
|----------------------|---|
| Diagnosis | Chronic rhinosinusitis with nasal polyps (CRSwNP) |

| Approval Length | 12 month(s) |
|-----------------|---------------------|
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting positive clinical response to therapy (e.g., reduction in nasal polyps score [NPS; 0-8 scale], improvement in nasal obstruction symptoms via visual analog scale [VAS; 0-10 scale])

AND

2 - Used in combination with another agent for CRSwNP

AND

3 - Prescribed by or in consultation with one of the following:

- Allergist/Immunologist
- Otolaryngologist
- Pulmonologist

| Product Name: Nucala | | |
|----------------------|--|--|
| Diagnosis | Eosinophilic Granulomatosis with Polyangiitis (EGPA) | |
| Approval Length | 12 Months | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of eosinophilic granulomatosis with polyangiitis (EGPA)

AND

2 - Patient's disease has relapsed or is refractory to standard of care therapy (i.e., corticosteroid treatment with or without immunosuppressive therapy)

AND

3 - Patient is currently receiving corticosteroid therapy (e.g., prednisolone, prednisone)

AND

4 - Prescribed by or in consultation with one of the following:

- Pulmonologist
- Rheumatologist
- Allergist/Immunologist

| Product Name: Nucala | | |
|----------------------|--|--|
| Diagnosis | Eosinophilic Granulomatosis with Polyangiitis (EGPA) | |
| Approval Length | 12 Months | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting positive clinical response to therapy (e.g., increase in remission time)

| Product Name: Nucala | | |
|----------------------|----------------------------------|--|
| Diagnosis | Hypereosinophilic Syndrome (HES) | |
| Approval Length | 12 Months | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| | | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of hypereosinophilic syndrome (HES)

AND

2 - Patient has been diagnosed for at least 6 months

AND

3 - Verification that other non-hematologic secondary causes have been ruled out (e.g., drug hypersensitivity, parasitic helminth infection, HIV infection, non-hematologic malignancy)

AND

4 - Patient is Fip1-like1-platelet-derived growth factor receptor alpha (FIP1L1-PDGFRA)negative

AND

5 - Patient has uncontrolled HES defined as both of the following:

- History of 2 or more flares within the past 12 months [I]
- Pre-treatment blood eosinophil count greater than or equal to 1000 cells/microliter

AND

6 - Trial and failure, contraindication, or intolerance to one of the following:

- Corticosteroid therapy (e.g., prednisone)
- Cytotoxic/immunosuppressive therapy (e.g., hydroxyurea, cyclosporine, imatinib)

AND

7 - Prescribed by or in consultation with one of the following:

- Allergist/Immunologist
- Hematologist

| Product Name: Nucala | | |
|------------------------------------|----------------------------------|--|
| Diagnosis | Hypereosinophilic Syndrome (HES) | |
| Approval Length | 12 Months | |
| Therapy Stage | Reauthorization | |
| Guideline Type Prior Authorization | | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting positive clinical response to therapy (e.g., reduction in flares, decreased blood eosinophil count, reduction in corticosteroid dose)

2. Background

The Global Initiative for Asthma Global Strategy for Asthma Management and Prevention: Table 1. Low, medium and high daily doses of inhaled corticosteroids in adolescents and adults 12 years and older [6]

| Inhaled corticosteroid | Total Daily ICS Dose (mcg) | | |
|---|----------------------------|------------|--------|
| | Low | Medium | High |
| Beclometasone dipropionate (pMDI, standard particle, HFA) | 200-500 | > 500-1000 | > 1000 |
| Beclometasone dipropionate (pMDI, extrafine particle*, HFA) | 100-200 | > 200-400 | > 400 |
| Budesonide (DPI) | 200-400 | > 400-800 | > 800 |
| Ciclesonide (pMDI, extrafine particle*, HFA) | 80-160 | > 160-320 | > 320 |

| Fluticasone furoate (DPI) | | 100 | 200 |
|---|--|---|------------------------|
| Fluticasone propionate (DPI) | 100-250 | > 250-500 | > 500 |
| Fluticasone propionate (pMDI, standard particle, HFA) | 100-250 | > 250-500 | > 500 |
| Mometasone furoate (DPI) | | 200 | 400 |
| Mometasone furoate (pMDI, standard particle, HFA) | 20 | 0-400 | > 400 |
| with a spacer *See product information |) . | | |
| This is not a table of equivalence, bu | | • • | |
| This is not a table of equivalence, but for the 'low', 'medium' and 'high' dose l based on available studies and product potency are not readily available and th potency equivalence. Doses may be co availability, regulatory labelling and clir | ICS options fo t information. herefore this ta puntry -specific | r adults/adolesce Data on compara able does NOT in c depending on lo | ents, ative nply |

| Date | Notes |
|-----------|---|
| 6/22/2022 | Updated criteria for all approved indications |

Nuedexta

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99482 Nuedexta

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Nuedexta | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of pseudobulbar affect (PBA) | |

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Standard to Medicaid Arizona Standard for 7 /1 go live |

Nuplazid

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99483 Nuplazid

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Nuplazid | |
|--------------------------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of Parkinson's disease | |

2 - Patient is currently experiencing hallucinations and delusions associated with Parkinson's disease psychosis (i.e., hallucination and delusion symptoms started after Parkinson's disease diagnosis)

| Product Name: Nuplazid | |
|------------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Nuplazid therapy

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona Standard to Medicaid Arizona Standard for 7 /1 go live |

Nurtec, Ubrelvy - AZM

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-112910 Nurtec, Ubrelvy - AZM

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 9/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Ubrelvy | |
|-----------------------|-----------------------------|
| Diagnosis | Acute Treatment of Migraine |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| | |

1 - Used for acute treatment of migraine

2 - Patient has a history of a one-month trial resulting in therapeutic failure, contraindication, or intolerance to TWO of the following as evidenced by submission of medical records or claims history:

- naratriptan tablets
- rizatriptan tablets/ODT (Oral Disintegrating Tablets)
- sumatriptan auto injection/cartridge
- zolmitriptan tablets/ODT
- Zomig nasal spray (Brand only)
- Imitrex nasal spray (Brand only)

AND

3 - Prescribed by or in consultation with one of the following specialists with expertise in the acute treatment of migraine:

- Neurologist
- Pain specialist
- Headache specialist*

AND

4 - One of the following:

4.1 Patient is currently treated with TWO of the following prophylactic therapies as evidenced by submission of medical records or claims history:

- amitriptyline (Elavil)
- A beta-blocker (i.e., atenolol, metoprolol, or propranolol)
- divalproex sodium [(Depakote/Depakote ER (extended release)]
- topiramate (Topamax)
- venlafaxine [Effexor/Effexor XR (extended release)]

OR

4.2 The patient has less than 4 migraine days per month

OR

4.3 Both of the following:

4.3.1 The patient has greater than or equal to 4 migraine days per month

AND

4.3.2 Patient has a history of failure, contraindication, or intolerance to TWO of the following prophylactic therapies as evidenced by submission of medical records or claims history:

- amitriptyline (Elavil)
- A beta-blocker (i.e., atenolol, metoprolol, or propranolol)
- divalproex sodium (Depakote/Depakote ER)
- topiramate (Topamax)
- venlafaxine (Effexor/Effexor XR)

AND

5 - Patient has tried ALL of the following calcitonin gene-related peptide receptor (CGRP) antagonist for preventive treatment of migraine:

- Ajovy (fremanezumab)
- Emgality (galacanezumab)
- Aimovig (erenumab-aooe)

AND

6 - This medication will not be used in combination with another acute calcitonin gene-related peptide receptor (CGRP) antagonist (i.e., Nurtec)

| Notes | *Headache specialists are physicians certified by the United Council f |
|-------|--|
| | or Neurologic Subspecialties (UCNS). |

| Product Name: Nurtec ODT | |
|--------------------------|-------------|
| Diagnosis | Migraine |
| Approval Length | 12 month(s) |

| Therapy Stage | Initial Authorization | |
|--|---|--|
| Guideline Type | Prior Authorization | |
| Approval Criteria 1 - Submission of medical records (e.g., chart notes) documenting one of the following | | |
| diagnoses: | | |
| 1.1 Used for acute tre | atment of migraine | |
| | OR | |
| 1.2 Used for preventiv | e treatment of episodic migraine | |
| | AND | |
| 2 - BOTH of the following | ng: | |
| 2.1 Patient has a history of a one-month trial resulting in therapeutic failure, contraindication, or intolerance to FOUR of the following as evidenced by submission of medical records or claims history: | | |
| sumatriptan aut | ts/ODT (Oral Disintegrating Tablets) o injection/cartridge ray (Brand only) lets/ODT | |
| AND | | |
| 2.2 Patient has a history of a one-month trial resulting in therapeutic failure, contraindication, or intolerance to Ubrelvy as evidenced by submission of medical records or claims history** | | |
| | AND | |
| 3 - Prescribed by or in a acute treatment of migr | consultation with one of the following specialists with expertise in the raine: | |

| Neurologist Pain specialist Headache specialist* |
|---|
| AND |
| 4 - One of the following: |
| 4.1 Patient is currently treated with TWO of the following prophylactic therapies as evidenced by submission of medical records or claims history: |
| amitriptyline (Elavil) A beta-blocker (i.e., atenolol, metoprolol, or propranolol divalproex sodium [(Depakote/Depakote ER (extended release)] topiramate (Topamax) venlafaxine [Effexor/Effexor XR (extended release)] |
| OR |
| 4.2 The patient has less than 4 migraine days per month |
| OR |
| 4.3 Both of the following: |
| 4.3.1 The patient has greater than or equal to 4 migraine days per month |
| AND |
| 4.3.2 Patient has a history of failure, contraindication, or intolerance to TWO of the following prophylactic therapies as evidenced by submission of medical records or claims history: |
| amitriptyline (Elavil) A beta-blocker (i.e., atenolol, metoprolol, or propranolol) divalproex sodium (Depakote/Depakote ER) topiramate (Topamax) venlafaxine (Effexor/Effexor XR) |

5 - Patient has tried ALL of the following calcitonin gene-related peptide receptor (CGRP) antagonist*** for preventive treatment of migraine:

- Ajovy (fremanezumab)
- Emgality (galacanezumab)
- Aimovig (erenumab-aooe)

AND

6 - This medication will not be used in combination with another acute calcitonin gene-related peptide receptor (CGRP) antagonist (i.e., Ubrelvy)

| or N hori actu m o equ erap | eadache specialists are physicians certified by the United Council f Neurologic Subspecialties (UCNS). **Patients requesting initial aut ization who were established on therapy via the receipt of a manuf urer supplied sample at no cost in the prescriber's office or any for of assistance from the manufacturer sponsored programs shall be r uired to meet initial authorization criteria as if patient were new to th py. ***CGRP antagonists for preventive treatment of migraines req a prior authorization. |
|--|---|
|--|---|

| Product Name: Ubrelvy, Nurtec ODT | |
|-----------------------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to therapy

AND

2 - This medication will not be used in combination with another calcitonin gene-related peptide receptor (CGRP) antagonist (i.e., Nurtec ODT for Ubrelvy requests, Ubrelvy for Nurtec ODT requests, Aimovig, Ajovy, & Emgality)

| Date | Notes |
|-----------|--|
| 8/29/2022 | Changed number of prerequisite triptan therapies for both targets. |

Nuzyra

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99522 Nuzyra

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Nuzyra | | |
|---|--|--|
| Diagnosis | Community-Acquired Bacterial Pneumonia | |
| Approval Length | 14 Day(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - ONE of the following: | | |
| 1.1 For continuation of therapy upon hospital discharge | | |

OR

1.2 As continuation of therapy when transitioning from intravenous antibiotics that are shown to be sensitive to the cultured organism for the requested indication

OR

1.3 ALL of the following:

1.3.1 Diagnosis of community-acquired bacterial pneumonia (CABP)

AND

1.3.2 Infection caused by an organism that is confirmed to be or likely to be susceptible to treatment with Nuzyra

AND

1.3.3 History of failure, contraindication, or intolerance to THREE of the following antibiotics or antibiotic regimens:

- Amoxicillin
- A macrolide
- Doxycycline
- A fluoroquinolone
- Combination therapy with amoxicillin/clavulanate or cephalosporin AND a macrolide or doxycycline

| Product Name: Nuzyra | |
|----------------------|--|
| Diagnosis | Acute Bacterial Skin and Skin Structure Infections |
| Approval Length | 14 Day(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - ONE of the following:

1.1 For continuation of therapy upon hospital discharge

OR

1.2 As continuation of therapy when transitioning from intravenous antibiotics that are shown to be sensitive to the cultured organism for the requested indication

OR

1.3 ALL of the following:

1.3.1 ONE of the following diagnoses:

1.3.1.1 BOTH of the following:

- Acute bacterial skin and skin structure infections
- Infection caused by methicillin-resistant Staphylococcus aureus (MRSA) documented by culture and sensitivity report

OR

1.3.1.2 BOTH of the following:

- Empirical treatment of patients with acute bacterial skin and skin structure infections
- Presence of MRSA infection is likely

AND

1.3.2 History of failure, contraindication, or intolerance to linezolid (generic Zyvox)

AND

1.3.3 History of failure, contraindication, or intolerance to ONE of the following antibiotics:

- Sulfamethoxazole-trimethoprim (SMZ-TMP)
- A tetracycline

• Clindamycin

OR

1.4 ALL of the following:

1.4.1 Diagnosis of acute bacterial skin and skin structure infections

AND

1.4.2 Infection caused by an organism that is confirmed to be or likely to be susceptible to treatment with Nuzyra

AND

1.4.3 History of failure, contraindication, or intolerance to THREE of the following antibiotics:

- A penicillin
- A cephalosporin
- A tetracycline
- Sulfamethoxazole-trimethoprim (SMZ-TMP)
- Clindamycin

| Product Name: Nuzyra | |
|----------------------|--|
| Off-Label Uses* | |
| Prior Authorization | |
| | |

Approval Criteria

- **1** ONE of the following:
- **1.1** For continuation of therapy upon hospital discharge

1.2 As continuation of therapy when transitioning from intravenous antibiotics that are shown to be sensitive to the cultured organism for the requested indication

OR

1.3 The medication is being prescribed by or in consultation with an infectious disease specialist.

| Notes | *Note: Authorization duration based on provider treatment durations, n |
|-------|--|
| | ot to exceed 6 months. |

| Date | Notes |
|-----------|-------------------------------------|
| 5/13/2021 | Arizona Medicaid 7.1 Implementation |

OAB Agents- Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-100643 | OAB Agents- Arizona |
|-----------|---------------------|
|-----------|---------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|--------------------|-----------|
| P&T Approval Date: | |
| P&T Revision Date: | |

1. Criteria

| Product Name: Product Name: Oxytrol (Rx) patch, trospium ER, brand Enablex, generic darifenacin ER, Brand Vesicare, generic solifenacin, Myrbetriq, Gelnique, brand Ditropan XL, Flavoxate | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| Approval Criteria | |

1 - The patient has a history of failure, contraindication, or intolerance to a trial of THREE preferred products

- oxybutynin (generic Ditropan)
 oxybutynin ER (generic Ditropan XL)
 Brand Detrol
- Brand Detrol LA •
- Toviaz (fesoterodine) •

| Date | Notes |
|------------|---|
| 12/16/2021 | update guideline for new formulary no changes to criteria |

Ocaliva

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99630 Ocaliva

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Ocaliva | |
|-----------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of primary biliary cholangitis (aka primary biliary cirrhosis)

AND 2.1.2 Used in combination with ursodeoxycholic acid (e.g., Urso, ursodiol) OR 2.2 History of contraindication or intolerance to ursodeoxycholic acid (e.g., Urso, ursodiol)

AND

3 - Prescribed by ONE of the following:

Hepatologist

Gastroenterologist

| Product Name: Ocaliva | |
|-----------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | * |

Approval Criteria

1 - Submission of medical records (e.g., laboratory values) documenting a reduction in

AND

2 - ONE of the following:

2.1 BOTH of the following:

2.1.1 Patient has failed to achieve an alkaline phosphatase (ALP) level of less than 1.67 times the upper limit of normal after at least 12 consecutive months of treatment with ursodeoxycholic acid(e.g., Urso, ursodiol)

alkaline phosphatase (ALP) level from pre-treatment baseline (i.e., prior to Ocaliva therapy) while on Ocaliva therapy

AND

- **2** Prescribed by ONE of the following:
 - •
 - Hepatologist Gastroenterologist •

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Olumiant (baricitinib)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-114464 | Olumiant (baricitinib) |
|-----------|------------------------|
|-----------|------------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Olumiant | |
|------------------------|-----------------------|
| Diagnosis | Rheumatoid Arthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes) confirming diagnosis of moderately to severely active rheumatoid arthritis

2 - Prescribed by or in consultation with a rheumatologist

AND

3 - Paid claims or submission of medical records (e.g., chart notes) confirming trial and failure, contraindication, or intolerance to ONE nonbiologic disease-modifying antirheumatic drug (DMARD) (e.g., methotrexate, leflunomide, sulfasalazine)

AND

- **4** One of the following:
- **4.1** All of the following:

4.1.1 Paid claims or submission of medical records (e.g., chart notes) confirming trial and failure, contraindication, or intolerance to TWO of the following, or attestation demonstrating a trial may be inappropriate*

- Humira (adalimumab)
- Enbrel (etanercept)
- Xeljanz (tofacitinib)

AND

4.1.2 Paid claims or submission of medical records (e.g., chart notes) confirming trial and failure, contraindication, or intolerance to Orencia (abatacept)

OR

4.2 Paid claims or submission of medical records (e.g., chart notes) confirming continuation of prior Olumiant therapy

AND

| 5 - Not used in combination with other Janus kinase (JAK) inhibitors, biologic DMARDs, or potent immunosuppressants (e.g., azathioprine or cyclosporine)** | |
|---|--|
| Notes | *Includes attestation that a total of two TNF inhibitors have already be en tried in the past, and the patient should not be made to try a third T NF inhibitor. **Olumiant may be used with concomitant methotrexate, topical or inhaled corticosteroids, and/or low stable dosages of oral co rticosteroids (equivalent to 10 mg or less of prednisone daily). |

| Product Name: Olumiant | |
|------------------------|----------------------|
| Diagnosis | Rheumatoid Arthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to therapy

AND

2 - Not used in combination with other JAK inhibitors, biologic DMARDs, or potent immunosuppressants (e.g., azathioprine or cyclosporine)**

| Notes | **Olumiant may be used with concomitant methotrexate, topical or inh |
|-------|---|
| | aled corticosteroids, and/or low stable dosages of oral corticosteroids |
| | (equivalent to 10 mg or less of prednisone daily). |

| Product Name: Olumiant | |
|------------------------|-------------------------------------|
| Diagnosis | Coronavirus disease 2019 (COVID-19) |
| Approval Length | 14 Day(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of COVID-19

2 - Patient is hospitalized

AND

- **3** Patient requires one of the following:
 - •
 - Supplemental oxygen Non-invasive mechanical ventilation •
 - Invasive mechanical ventilation •
 - Extracorporeal membrane oxygenation (ECMO) •

| Product Name: Olumiant | |
|---|-------------------------------------|
| Diagnosis | Coronavirus disease 2019 (COVID-19) |
| Approval Length | 14 Day(s) |
| Guideline Type | Non Formulary |
| | |
| Approval Criteria | |
| 1 - Diagnosis of COVID | D-19 |
| | |
| | AND |
| | |
| 2 - Patient is hospitalized | |
| | |
| | AND |
| 3 - Patient requires one of the following: | |
| Supplemental oxygen Non-invasive mechanical ventilation Invasive mechanical ventilation | |

Extracorporeal membrane oxygenation (ECMO)

| Product Name: Olumiant | |
|--|--|
| Diagnosis | Alopecia Areata |
| Approval Length | N/A - Requests for non-approvable diagnoses should not be approved |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Requests for coverage for diagnosis of Alopecia Areata are not authorized and will not be approved | |
| Notos | Approval Longth: N/A - Poquests for Alonocia Areata should not be ap |

| Notes | Approval Length: N/A - Requests for Alopecia Areata should not be ap |
|-------|--|
| | proved. Deny as a benefit exclusion. |

| Date | Notes |
|-----------|--|
| 9/26/2022 | Updated criteria for RA. Added criteria for Covid-19 and alopecia are ata indications. |

Opzelura (ruxolitinib)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-114493 | Opzelura (ruxolitinib) |
|-----------|------------------------|
|-----------|------------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Opzelura | |
|------------------------|-----------------------|
| Diagnosis | Atopic Dermatitis |
| Approval Length | 12 weeks [A] |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of mild to moderate atopic dermatitis

2 - One of the following:

- Greater than or equal to 3% body surface area (BSA) involvement
- Involvement of sensitive body areas (e.g., face, hands, feet, scalp, groin)

AND

3 - Patient is 12 years of age or older

AND

4 - Prescribed by or in consultation with one of the following:

- Dermatologist
- Allergist/Immunologist

AND

5 - Trial and failure of a minimum 30-day supply of non-pharmacologic topical therapies (e.g., moisturizers) [2]

AND

6 - Trial and failure of a minimum 30-day supply (14-day supply for topical corticosteroids), contraindication, or intolerance to at least TWO of the following:

- Medium or higher potency topical corticosteroid
- Elidel (pimecrolimus) cream*
- Tacrolimus ointment
- Eucrisa (crisaborole) ointment*

7 - Patient is not receiving Opzelura in combination with a potent immunosuppressant (e.g., azathioprine or cyclosporine)

AND

8 - Opzelura will only be used for short-term and/or non-continuous chronic treatmentNotes*Product may require step therapy

| Product Name: Opzelura | |
|------------------------|---------------------|
| Diagnosis | Atopic Dermatitis |
| Approval Length | 6 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of a positive clinical response to therapy as evidenced by at least ONE of the following:

- Reduction in body surface area involvement from baseline
- Reduction in pruritus severity from baseline
- Improvement in quality of life from baseline

AND

2 - Patient is not receiving Opzelura in combination with a potent immunosuppressant (e.g., azathioprine or cyclosporine)

AND

3 - Opzelura will only be used for short-term and/or non-continuous chronic treatment

| Product Name: Opzelura | |
|------------------------|-----------------------|
| Diagnosis | Nonsegmental Vitiligo |

| Approval Length | N/A - Requests for non-approvable diagnoses should not be approved |
|--------------------|--|
| Guideline Type | Prior Authorization |
| | |
| Arene vel Oritoria | |

Approval Criteria

1 - Requests for coverage for diagnosis of nonsegmental vitiligo are not authorized and will not be approved

| Notes | Approval Length: N/A - Requests for nonsegmental vitiligo should not |
|-------|--|
| | be approved. Deny as a benefit exclusion. |

2. Background

| Class | Drug | Dosage Form | Strength (%) |
|----------------------|--------------------------------------|---------------------------------|-----------------|
| Very high potency | Augmented betamethasone dipropionate | Ointment, gel | 0.05 |
| | Clobetasol propionate | Cream, foam, ointment | 0.05 |
| | Diflorasone diacetate | Ointment | 0.05 |
| | Halobetasol propionate | Cream, ointment | 0.05 |
| High | Amcinonide | Cream, lotion, ointment | 0.1 |
| Potency | Augmented betamethasone dipropionate | Cream, lotion | 0.05 |
| | Betamethasone dipropionate | Cream, foam, ointment, solution | 0.05 |
| | Desoximetasone | Cream, ointment | 0.25 |
| | Desoximetasone | Gel | 0.05 |
| | Diflorasone diacetate | Cream | 0.05 |
| | Fluocinonide | Cream, gel, ointment, solution | 0.05 |

| | Halcinonide | Cream, ointment | 0.1 |
|------------------|----------------------------|-----------------------------------|--------------|
| | Mometasone furoate | Ointment | 0.1 |
| | Triamcinolone acetonide | Cream, ointment | 0.5 |
| Medium | Betamethasone valerate | Cream, foam, lotion, ointment | 0.1 |
| potency | Clocortolone pivalate | Cream | 0.1 |
| | Desoximetasone | Cream | 0.05 |
| | Fluocinolone acetonide | Cream, ointment | 0.025 |
| | Flurandrenolide | Cream, ointment, lotion | 0.05 |
| | Fluticasone propionate | Cream | 0.05 |
| | Fluticasone propionate | Ointment | 0.005 |
| | Mometasone furoate | Cream, lotion | 0.1 |
| | Triamcinolone acetonide | Cream, ointment, lotion | 0.1 |
| Lower- medium | Hydrocortisone butyrate | Cream, ointment, solution | 0.1 |
| potency | Hydrocortisone probutate | Cream | 0.1 |
| | Hydrocortisone valerate | Cream, ointment | 0.2 |
| | Prednicarbate | Cream | 0.1 |
| Low | Alclometasone dipropionate | Cream, ointment | 0.05 |
| potency | Desonide | Cream, gel, foam, ointment | 0.05 |
| | Fluocinolone acetonide | Cream, solution | 0.01 |
| Lowest | Dexamethasone | Cream | 0.1 |
| potency | Hydrocortisone | Cream, lotion, ointment, solution | 0.25, 0.5, 1 |
| | Hydrocortisone acetate | Cream, ointment | 0.5-1 |

| Date Notes | |
|------------|--|
|------------|--|

| 9/26/2022 | Added denial criteria for nonsegmental vitiligo |
|-----------|---|
| | |

Oral Oncology Agents

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-113505 | Oral Oncology Agents |
|-----------|----------------------|
|-----------|----------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 9/8/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

Product Name: Non-Preferred Oral Oncology Drugs: Alunbrig, Avyakit, Balversa, Bosulif,
Braftovi, Brukinsa, Cabometyx, Calquence, Cometriq, Copiktra, Cotellic, Daurismo, Erleada,
Farydak, Gavreto, Gilotrif, Hycamtin capsules, Ibrance, Iclusig, Idhifa, Inrebic, Kisqali, Kisqali-
Femara Co-pack, Koselugo, Lenvima, Lonsurf, Lorbrena, Lynparaza, Mekinist, Mektovi,
Nerlynx, Ninlaro, Nubeqa, Odomzo, Pemazyre, Piqray, Pomalyst, Qinlock, Retevmo,
Rozlytrek, Rubraca, Rydapt, Stivarga, Tabrecta, Tafinlar, Tagrisso, Talzenna, Tazverik,
Tepmetko, Tibsovo, Tukysa, Turalio, Venclexta, Verzenio, Vitrakvi, Vizimpro, Xospata,
Xpovio, Xtandi, Zejula, Zydelig, ZykadiaDiagnosisCancer IndicationsApproval Length12 month(s)Guideline TypePrior Authorization

Approval Criteria

1 - The drug is being used as indicated by National Comprehensive Cancer Network (NCCN) guidelines with a Category of Evidence and Consensus of 1, 2A, or 2B

| Date | Notes |
|----------|---|
| 9/8/2022 | Updated with additional NP targets. Caprelsa and Xalkori (Preferred) moved to drug-specific guidelines. |

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

Effective Date: 9/8/2022

1. Criteria

Product Name: Non-Preferred Oral Oncology Drugs: Alunbrig, Avyakit, Balversa, Bosulif,
Braftovi, Brukinsa, Cabometyx, Calquence, Cometriq, Copiktra, Cotellic, Daurismo, Erleada,
Farydak, Gavreto, Gilotrif, Hycamtin capsules, Ibrance, Iclusig, Idhifa, Inrebic, Kisqali, Kisqali-
Femara Co-pack, Koselugo, Lenvima, Lonsurf, Lorbrena, Lynparaza, Mekinist, Mektovi,
Nerlynx, Ninlaro, Nubeqa, Odomzo, Pemazyre, Piqray, Pomalyst, Qinlock, Retevmo,
Rozlytrek, Rubraca, Rydapt, Stivarga, Tabrecta, Tafinlar, Tagrisso, Talzenna, Tazverik,
Tepmetko, Tibsovo, Tukysa, Turalio, Venclexta, Verzenio, Vitrakvi, Vizimpro, Xospata,
Xpovio, Xtandi, Zejula, Zydelig, ZykadiaDiagnosisCancer IndicationsApproval Length12 month(s)Guideline TypePrior Authorization

1 - The drug is being used as indicated by National Comprehensive Cancer Network (NCCN) guidelines with a Category of Evidence and Consensus of 1, 2A, or 2B

| Date | Notes |
|----------|---|
| 9/8/2022 | Updated with additional NP targets. Caprelsa and Xalkori (Preferred) moved to drug-specific guidelines. |

Orencia (abatacept)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-114537 Orencia (abatacept)

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Orencia IV or Orencia SC | |
|--|---------------------------|
| Diagnosis | Rheumatoid Arthritis (RA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of moderately to severely active rheumatoid arthritis

2 - Prescribed by or in consultation with a rheumatologist

AND

3 - Trial and failure, contraindication, or intolerance to ONE nonbiologic disease-modifying antirheumatic drug (DMARD) (e.g., methotrexate [Rheumatrex/Trexall], Arava [leflunomide], Azulfidine [sulfasalazine])

| Product Name: Orencia IV or Orencia SC | |
|--|---------------------------|
| Diagnosis | Rheumatoid Arthritis (RA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to therapy

| Product Name: Orencia IV or Orencia SC | |
|--|--|
| Diagnosis | Polyarticular Juvenile Idiopathic Arthritis (PJIA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of moderately to severely active polyarticular juvenile idiopathic arthritis

AND

 ${\bf 2}$ - Prescribed by or in consultation with a rheumatologist

AND

3 - Trial and failure, contraindication, or intolerance to ONE of the following nonbiologic disease modifying anti-rheumatic drugs (DMARDs):

- •
- leflunomide (Arava) methotrexate (Rheumatrex/Trexall) •

| Product Name: Orencia IV or Orencia SC | |
|--|--|
| Diagnosis | Polyarticular Juvenile Idiopathic Arthritis (PJIA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | • |

Approval Criteria

1 - Documentation of positive clinical response to therapy

| Product Name: Orencia IV or Orencia SC | |
|--|---------------------------|
| Diagnosis | Psoriatic Arthritis (PsA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of active psoriatic arthritis (PsA)

AND

2 - Prescribed by or in consultation with one of the following:

- •
- Dermatologist Rheumatologist •

| Product Name: Orencia IV or Orencia SC | |
|--|---------------------------|
| Diagnosis | Psoriatic Arthritis (PsA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to therapy

| Product Name: Orencia IV | | |
|--|---|--|
| Diagnosis | Prophylaxis for Acute Graft versus Host Disease (aGVHD) | |
| Approval Length | 2 month(s) | |
| Guideline Type | Prior Authorization | |
| Approval Criteria 1 - Used for prophylaxis of acute graft versus host disease (aGVHD) | | |
| | AND | |
| 2 - Patient is 2 years of age or older | | |
| | AND | |
| 3 - Patient will receive hematopoietic stem cell transplantation (HSCT) from a matched or 1 allele-mismatched unrelated donor | | |

4 - Recommended antiviral prophylactic treatment for Epstein-Barr Virus (EBV) reactivation (e.g., acyclovir) will be administered prior to Orencia and continued for six months after HSCT

AND

5 - Used in combination with both of the following:

- calcineurin inhibitor (e.g., cyclosporine, tacrolimus)
- methotrexate

| Date | Notes |
|-----------|---|
| 9/26/2022 | Added criteria for GVHD. Removed embedded step through preferre d agents. |

Orfadin- Arizona

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99631 O | rfadin- Arizona |
|------------|-----------------|
|------------|-----------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Orfadin, generic nitisinone | |
|---|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of hereditary tyrosinemia type 1 | |

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Oriahnn

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99484 Oriahnn

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Oriahnn | |
|--|-----------------------|
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of uterine fibroids (leiomyomas) | |

2 - Used for the management of heavy menstrual bleeding

AND

3 - Patient is premenopausal

AND

4 - History of trial and failure, contraindication, or intolerance after a three-month trial to ONE of the following:

- Estrogen/progestin contraceptive (e.g., Loestrin FE)
- Progestin-releasing intrauterine devices (IUDs) (e.g., Mirena)*
- Progestin-only contraceptive [e.g., norethindrone (generic Aygestin)]

AND

5 - History of trial and failure, contraindication or intolerance after a three-month trial of tranexamic acid (e.g., Lysteda)

AND

6 - Prescribed by or in consultation with ONE of the following:

- Obstetrics/Gynecologist (OB/GYN)
- Reproductive endocrinologist

Notes *This is a medical benefit, should not be included in denial to provider

| Product Name: Oriahnn | |
|-----------------------|-----------------|
| Approval Length | 6 months* |
| Therapy Stage | Reauthorization |

| Guideline Type | Prior Authorization | |
|--|--|--|
| | | |
| Approval Criteria | | |
| 1 - Documentation of p | ositive clinical response to therapy | |
| | | |
| AND | | |
| 2 - Impact to bone mineral density has been considered | | |
| AND | | |
| 3 - Treatment duration has not exceeded a total of 24 months** | | |
| Notes | *Authorization will be issued for 6 months up to a maximum of 24 mon ths **Oriahnn is indicated for a maximum of 24 months | |

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Standard to Medicaid Arizona Standard for 7 /1 go live |

Orilissa

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99485 Orilissa

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Orilissa 150 mg | |
|-------------------------------|-----------------------|
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of moderate to severe pain associated with endometriosis

2 - Patient is premenopausal

AND

3 - History of trial and failure (e.g., inadequate pain relief), contraindication or intolerance after a three month trial of TWO analgesics (e.g., ibuprofen, meloxicam, naproxen)

AND

4 - History of trial and failure, contraindication, or intolerance after a three month trial to ONE of the following:

- Hormonal contraceptives
- Progestins [e.g., norethindrone (generic Aygestin)]

AND

- 5 Prescribed by or in consultation with ONE of the following:
 - Obstetrics/Gynecologist (OB/GYN)
 - Reproductive endocrinologist

| Product Name: Orilissa 150 mg | |
|-------------------------------|--|
| 6 months* | |
| Reauthorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Documentation of positive clinical response to therapy

| | AND | |
|--|---|--|
| 2 - Impact to bone mineral density has been considered | | |
| | AND | |
| 3 - Treatment duration has not exceeded a total of 24 months** | | |
| Notes | *NOTE: Authorization for Orilissa 150 mg will be issued for 6 months up to a maximum of 24 months. **NOTE: Orilissa 150 mg once daily i s indicated for a maximum of 24 months. | |

| Product Name: Orilissa | Product Name: Orilissa 200 mg | |
|--|-------------------------------|--|
| Approval Length | 6 months* | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of moderate to severe pain associated with endometriosis | | |
| | AND | |
| 2 - Patient is premenopausal | | |
| | AND | |
| 3 - History of trial and failure (e.g., inadequate pain relief), contraindication or intolerance after a three month trial of TWO analgesics (e.g., ibuprofen, meloxicam, naproxen) | | |
| | AND | |
| 4 - History of trial and failure, contraindication, or intolerance after a three month trial to ONE of the following: | | |

- •
- Hormonal contraceptives Progestins [e.g., norethindrone (generic Aygestin)] •

- **5** Prescribed by or in consultation with ONE of the following:
 - Obstetrics/Gynecologist (OB/GYN) Reproductive endocrinologist ٠
 - •

| *NOTE: Orilissa 200 mg twice daily is indicated for a maximum of 6 m onths. |
|---|
| |

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona Standard to Medicaid Arizona Standard for 7 /1 go live |

Orkambi

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99632 Orkambi

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Orkambi | |
|---------------------------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of cystic fibrosis (CF) | |

2 - Submission of laboratory results confirming that patient is homozygous for the F508del mutation in the cystic fibrosis transmembrane conductance regulator (CFTR) gene

AND

3 - The patient is greater than or equal to 2 years of age

AND

4 - Prescribed by, or in consultation with, a specialist affiliated with a CF care center

| Product Name: Orkambi | |
|-----------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Provider attests that the patient has achieved a clinically meaningful response while on Orkambi therapy to ONE of the following:

- Lung function as demonstrated by percent predicted expiratory volume in 1 second (ppFEV1)
- Body mass index (BMI)
- Pulmonary exacerbations
- Quality of life as demonstrated by Cystic Fibrosis Questionnaire-Revised (CFQ-R) respiratory domain score

AND

2 - Prescribed by, or in consultation with, a specialist affiliated with a cystic fibrosis (CF) care center

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Osphena - Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Osphena | |
|-----------------------|--|
| 12 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Treatment of moderate to severe vaginal dryness, a symptom of vulvar and vaginal atrophy (VVA), due to menopause*

AND 2 - History of failure, contraindication, or intolerance to BOTH of the following: Estradiol vaginal cream Estradiol vaginal tablet Notes *Treatment of dyspareunia is a benefit exclusion.

| Product Name: Osphena | | |
|--|---------------------|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Documentation of positive clinical response to therapy | | |

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Standard to Medicaid Arizona Standard for 7 /1 go live |

Otezla

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99724 Otezla

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Otezla | |
|---|-----------------------|
| Diagnosis | Psoriatic Arthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of active psoriatic arthritis | |

2 - History of failure to a 3 month trial of methotrexate at the maximally indicated dose within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*

AND

3 - Patient is not receiving Otezla in combination with one of the following:

- Biologic disease-modifying antirheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]

AND

4 - Prescribed by or in consultation with ONE of the following:

Rheumatologist

Dermatologist

| Notes | *Note: Claims history may be used in conjunction as documentation of drug, date, and duration of trial |
|-------|--|
| | drug, date, and duration of that |

| Product Name: Otezla | |
|----------------------|-----------------------|
| Diagnosis | Behcet's Disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of Behcet's Disease

2 - Patient has active oral ulcers

AND

3 - History of failure, contraindication, or intolerance to one non-biologic (e.g., corticosteroids, colchicine) within the last 3 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*

AND

4 - Patient is not receiving Otezla in combination with one of the following:

- Biologic disease-modifying antirheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]

AND

5 - Prescribed by or in consultation with ONE of the following:

- Rheumatologist
- Dermatologist

| Notes | *Note: Claims history may be used in conjunction as documentation of |
|-------|--|
| | drug, date, and duration of trial |

| Product Name: Otezla | |
|----------------------|---------------------------------------|
| Diagnosis | Psoriatic Arthritis, Behcet's Disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Otezla therapy

AND

- **2** Patient is not receiving Otezla in combination with one of the following:
 - Biologic disease-modifying antirheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
 - Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]

AND

3 - Prescribed by or in consultation with ONE of the following:

- Rheumatologist
- Dermatologist

| Product Name: Otezla | |
|----------------------|-----------------------|
| Diagnosis | Plaque Psoriasis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of moderate to severe chronic plaque psoriasis

AND

2 - Greater than or equal to 3% body surface area involvement, palmoplantar, facial, or genital involvement, or severe scalp psoriasis

AND

3 - Both of the following:

3.1 History of failure to one of the following topical therapies, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial):*

- Corticosteroids (e.g., betamethasone, clobetasol, desonide)
- Vitamin D analogs (e.g., calcitriol, calcipotriene)
- Tazarotene
- Calcineurin inhibitors (e.g., tacrolimus, pimecrolimus)
- Anthralin
- Coal tar

AND

3.2 History of failure to a 3 month trial of methotrexate at the maximally indicated dose within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*

AND

4 - Patient is not receiving Otezla in combination with one of the following:

- Biologic disease-modifying antirheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]

AND

5 - Prescribed by or in consultation with a dermatologist

| Notes | *Note: Claims history may be used in conjunction as documentation of |
|-------|--|
| | drug, date, and duration of trial |

| Product Name: Otezla | |
|----------------------|---------------------|
| Diagnosis | Plaque Psoriasis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

| Approval Criteria |
|--|
| 1 - Documentation of positive clinical response to Otezla therapy |
| AND |
| 2 - Patient is not receiving Otezla in combination with one of the following: |
| Biologic disease-modifying antirheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)] Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)] |
| AND |
| 3 - Prescribed by or in consultation with a dermatologist |

2. Revision History

| Date | Notes |
|-----------|-------------------------------------|
| 5/13/2021 | Arizona Medicaid 7.1 Implementation |

Oxbryta (voxelotor)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-102750 | Oxbryta (voxelotor) |
|-----------|---------------------|
|-----------|---------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 2/3/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Oxbryta | | |
|--------------------------------------|-----------------------|--|
| Diagnosis | Sickle Cell Disease | |
| Approval Length | 6 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of sickle cell disease | | |

2 - Patient is 4 years of age or older

AND

3 - One of the following:

3.1 Patient is currently receiving hydroxyurea therapy

OR

3.2 Patient has a history of treatment failure, intolerance, or contraindication to hydroxyurea therapy

AND

4 - Patient has previously experienced 1 or more sickle cell-related vaso occlusive crises within the previous 12 months

AND

5 - Baseline hemoglobin (Hb) less than or equal to 10.5 grams per deciliter

AND

6 - Patient is not receiving concomitant chronic, prophylactic blood transfusion therapy

AND

7 - Patient is not to receive Oxbryta in combination with Adakveo (crizanlizumab-tmca)

8 - Prescribed by, or in consultation with, a hematologist or other specialist with expertise in the diagnosis and management of sickle cell disease

| Product Name: Oxbryta | |
|-----------------------|---------------------|
| Diagnosis | Sickle Cell Disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Oxbryta therapy as demonstrated by at least one of the following:

1.1 Increase in hemoglobin (Hb) by greater than or equal to 1 gram per deciliter from baseline

OR

1.2 Decrease in indirect bilirubin from baseline

OR

1.3 Decrease in percent reticulocyte count from baseline

OR

1.4 Patient has experienced a reduction in sickle cell-related vaso occlusive crises

AND

2 - Patient is not receiving Oxbryta in combination with Adakveo (crizanlizumab-tmca)

AND

3 - Patient is not receiving concomitant chronic, prophylactic blood transfusion therapy

AND

4 - Prescribed by, or in consultation with, a hematologist, or other specialist with expertise in the diagnosis and management of sickle cell disease

| Date | Notes |
|----------|--|
| 2/3/2022 | Updated age criterion due to expanded indication |

Oxervate

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99634 Oxervate

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

| Product Name: Oxervate | |
|--|------------------------|
| Diagnosis | Neurotrophic keratitis |
| Approval Length | 8 Week(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of Stage 2 or 3 neurotrophic keratitis | |

2 - History of failure to at least one OTC ocular artificial tear product (e.g., Systane® Ultra, Akwa® Tears, Refresh Optive®, Soothe® XP)

AND

- **3** Prescribed by or in consultation with ONE of the following:
 - Ophthalmologist
 - Optometrist

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Palforzia

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99635 Palforzia

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Palforzia | |
|-------------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis and clinical history of peanut allergy as documented by BOTH of the following:

1.1 A serum peanut-specific IgE level of greater than or equal to 0.35 kUA/L (kilo units of allergen per liter)

1.2 A meal wheal diameter that is at least 3mm (millimeters) larger than the negative control on skin-prick testing for peanut

AND

2 - ONE of the following:

2.1 BOTH of the following:

- Patient is 4 to 17 years of age
- Patient is in the initial dose escalation phase of therapy

OR

2.2 BOTH of the following:

- Patient is 4 years of age and older
- Patient is in the up-dosing or maintenance phase of therapy

AND

3 - Used in conjunction with a peanut-avoidant diet

AND

- 4 Patient does not have one of the following:
 - History of eosinophilic esophagitis (EoE) or eosinophilic gastrointestinal disease
 - History of severe or life-threatening episode(s) of anaphylaxis or anaphylactic shock within the past 2 months
 - Severe or poorly controlled asthma

AND

5 - Prescribed by or in consultation with an allergist or immunologist

AND

6 - Prescriber is certified/enrolled in the Palforzia REMS (Risk Evaluation and Mitigation Strategy) Program

| Product Name: Palforzia | | |
|--|--|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Documentation of p | ositive clinical response to Palforzia therapy | |
| | AND | |
| 2 - Used in conjunction with a peanut-avoidant diet | | |
| AND | | |
| 3 - Prescribed by or in consultation with an allergist or immunologist | | |
| | AND | |
| 4 - Prescriber is certified/enrolled in the Palforzia REMS (Risk Evaluation and Mitigation Strategy) Program | | |

| Date | Notes | | |
|------|-------|--|--|
|------|-------|--|--|

| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |
|-----------|--|
| | |

Palynziq

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99636 Palynziq

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

| Product Name: Palynziq | |
|--|-----------------------|
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of phenylketonuria (PKU) | |

2 - Patient is actively on a phenylalanine-restricted diet

AND

3 - Physician attestation that the patient will not be receiving Palynziq in combination with Kuvan (sapropterin dihydrochloride)

AND

4 - Submission of medical records (e.g. chart notes, laboratory values) documenting that the patient has a blood phenylalanine concentration greater than 600 micromoles per liter

| Product Name: Palynziq | |
|------------------------|---------------------|
| Approval Length | 6 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient is actively on a phenylalanine-restricted diet

AND

2 - ONE of the following:

2.1 Submission of medical records (e.g. chart notes, laboratory values) documenting that the patient has a blood phenylalanine concentration less than 600 micromoles per liter

OR

2.2 Submission of medical records (e.g. chart notes, laboratory values) documenting that the

patient has achieved a 20% reduction in blood phenylalanine concentration from pretreatment baseline

OR

2.3 BOTH of the following:

2.3.1 Patient is in initial titration/maintenance phase of dosing regimen (week 1-33)

AND

2.3.2 Patient will receive maximum labeled dosage of 40 milligrams (mg) once daily if response has not been obtained after 24 weeks of 20 mg once daily maintenance dosing

AND

3 - Submission of medical records (e.g. chart notes, laboratory values) documenting that the patient is not receiving Palynziq in combination with Kuvan (sapropterin dihydrochloride) [Prescription claim history that does not show any concomitant Kuvan claim within 60 days of reauthorization request may be used as documentation]

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Panretin

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99511 Panretin

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|--------------------|-----------|
| P&T Approval Date: | |
| P&T Revision Date: | |

1. Criteria

| Product Name: Panretin | |
|------------------------|------------------------------------|
| Diagnosis | AIDS-related Kaposi's Sarcoma (KS) |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of acquired immunodeficiency syndrome (AIDS)-related Kaposi's Sarcoma (KS)

2 - Patient is not receiving systemic anti-KS treatment

| Product Name: Panretin | |
|------------------------|--------------------------|
| Diagnosis | NCCN Recommended Regimen |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Panretin will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Panretin | |
|------------------------|--------------------------|
| Diagnosis | NCCN Recommended Regimen |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Panretin therapy

| Date | Notes |
|----------|--------------------|
| 4/8/2021 | 7/1 Implementation |

Pediculicides - AZM

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-105258 Pediculicides - AZM

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 4/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Sklice, Brand Natroba, generic spinosad susp | |
|--|---------------------|
| Diagnosis | Head lice |
| Approval Length | 30 Day(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of topical treatment of head lice infestations

2 - For Brand Natroba requests ONLY: Trial and failure to generic spinosad suspension (verified via paid pharmacy claims or submission of medical records/chart notes)

| Date | Notes |
|-----------|---|
| 3/28/2022 | Added step through generic for Brand Natroba. |

Praluent

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-102901 Praluent

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 2/3/2022 |
|-----------------|----------|
|-----------------|----------|

| Product Name: Praluent | |
|------------------------|---|
| Diagnosis | Primary Hyperlipidemia [Including Heterozygous Familial Hypercholesterolemia (HeFH), Atherosclerotic Cardiovascular Disease (ASCVD), and Secondary Prevention of Cardiovascular Events in Patients with ASCVD] |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of ONE of the following:

1.1 Heterozygous familial hypercholesterolemia (HeFH) as confirmed by ONE of the following*:

1.1.1 BOTH of the following:

1.1.1.1 Pre-treatment low density lipoprotein cholesterol (LDL-C) of ONE of the following:

- Greater than 190 milligrams per deciliter (mg/dL)
- Greater than 155 mg/dL if less than 16 years of age

AND

1.1.1.2 ONE of the following:

- Family history of myocardial infarction in first-degree relative less than 60 years of age
- Family history of myocardial infarction in second-degree relative less than 50 years of age
- Family history of LDL-C greater than 190 mg/dL in first- or second-degree relative
- Family history of heterozygous or homozygous familial hypercholesterolemia in first- or second-degree relative
- Family history of tendinous xanthomata and/or arcus cornealis in first- or second degree relative

OR

1.1.2 BOTH of the following:

1.1.2.1 Pre-treatment LDL-C of ONE of the following:

- Greater than 190 mg/dL
- Greater than 155 mg/dL if less than 16 years of age

AND

1.1.2.2 Submission of medical records (e.g., chart notes, laboratory values) documenting ONE of the following:

- Functional mutation in LDL (low density lipoprotein), apoB (apolipoprotein B), or PCSK9 (proprotein convertase subtilisin/kexin type 9) gene*
- Tendinous xanthomata

• Arcus cornealis before age 45

OR

1.2 Atherosclerotic cardiovascular disease (ASCVD) as confirmed by ONE of the following:

- Acute coronary syndromes
- History of myocardial infarction
- Stable or unstable angina
- Coronary or other arterial revascularization
- Stroke
- Transient ischemic attack
- Peripheral arterial disease presumed to be of atherosclerotic origin

AND

2 - Submission of medical records (e.g., chart notes, laboratory values) documenting ONE of the following [prescription claims history may be used in conjunction as documentation of medication use, dose, and duration]:

2.1 Patient has been receiving at least 12 consecutive weeks of high-intensity statin therapy [i.e. atorvastatin 40-80 milligrams (mg), rosuvastatin 20-40mg] and will continue to receive high intensity statin at maximally tolerated dose

OR

2.2 BOTH of the following:

2.2.1 Patient is unable to tolerate high-intensity statin as evidenced by one of the following intolerable and persistent (i.e. more than 2 weeks) symptoms:

- Myalgia (muscle symptoms without creatine kinase [CK] elevations)
- Myositis (muscle symptoms with CK elevations less than 10 times upper limit of normal [ULN])

AND

2.2.2 ONE of the following:

2.2.2.1 Patient has been receiving at least 12 consecutive weeks of moderate-intensity statin [i.e. atorvastatin 10-20 mg, rosuvastatin 5-10 mg, simvastatin greater than or equal to

20 mg, pravastatin greater than or equal to 40 mg, lovastatin 40 mg, Lescol XL (fluvastatin XL) 80 mg, fluvastatin 40 mg twice daily or Livalo (pitavastatin) greater than or equal to 2 mg] and will continue to receive a moderate-intensity statin at maximally tolerated dose

OR

2.2.2.2 Patient has been receiving at least 12 consecutive weeks of low-intensity statin [i.e. simvastatin 10 mg, pravastatin 10-20 mg, lovastatin 20 mg, fluvastatin 20-40 mg, or Livalo (pitavastatin) 1 mg] therapy and will continue to receive a low-intensity statin at maximally tolerated dose

OR

2.3 Patient is unable to tolerate low or moderate-, and high-intensity statins as evidenced by ONE of the following:

2.3.1 ONE of the following intolerable and persistent (i.e. more than 2 weeks) symptoms for low or moderate-, and high-intensity statins:

- Myalgia (muscle symptoms without CK elevations)
- Myositis (muscle symptoms with CK elevations less than 10 times upper limit of normal [ULN])

OR

2.3.2 Patient has a labeled contraindication to all statins as documented in medical records

OR

2.3.3 Patient has experienced rhabdomyolysis or muscle symptoms with statin treatment with CK elevations greater than 10 times ULN

AND

3 - ONE of the following:

3.1 Submission of medical records (e.g., laboratory values) documenting ONE of the following LDL-C values while on maximally tolerated lipid lowering therapy for a minimum of at least 12 weeks within the last 120 days:

• LDL-C greater than or equal to 100 mg/dL with ASCVD

• LDL-C greater than or equal to 130 mg/dL without ASCVD

OR

3.2 BOTH of the following:

3.2.1 Submission of medical records (e.g., laboratory values) documenting ONE of the following LDL-C values while on maximally tolerated lipid lowering therapy for a minimum of at least 12 weeks within the last 120 days:

• LDL-C between 70 mg/dL and 99 mg/dL with ASCVD

• LDL-C between 100 mg/dL and 129 mg/dL without ASCVD

AND

3.2.2 Submission of medical records (e.g., laboratory values) documenting ONE of the following [prescription claims history may be used in conjunction as documentation of medication use, dose, and duration]:

3.2.2.1 Patient has been receiving at least 12 consecutive weeks of ezetimibe (Zetia) therapy as adjunct to maximally tolerated statin therapy

OR

3.2.2.2 Patient has a history of contraindication or intolerance to ezetimibe

AND

4 - Used as an adjunct to a low-fat diet and exercise

AND

5 - Prescribed by ONE of the following:

Cardiologist

Endocrinologist

• Lipid specialist

AND

6 - Not used in combination with another proprotein convertase subtilisin/kexin type 9 (PCSK9) inhibitor (e.g., Repatha (evolocumab))

| Notes | *Note: Results of prior genetic testing can be submitted as confirmatio |
|-------|---|
| | n of diagnosis of HeFH. |

| Product Name: Praluent | |
|------------------------|---|
| Diagnosis | Primary Hyperlipidemia [Including Heterozygous Familial Hypercholesterolemia (HeFH), Atherosclerotic Cardiovascular Disease (ASCVD), and Secondary Prevention of Cardiovascular Events in Patients with ASCVD] |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient continues to receive statin at maximally tolerated dose (unless patient has documented inability to take statins)

AND

2 - Patient is continuing a low-fat diet and exercise regimen

AND

- **3** Prescribed by ONE of the following:
 - Cardiologist
 - Endocrinologist
 - Lipid specialist

4 - Submission of medical records (e.g. chart notes, laboratory values) documenting low density lipoprotein cholesterol (LDL-C) reduction while on Praluent therapy

AND

5 - Not used in combination with another proprotein convertase subtilisin/kexin type 9 (PCSK9) inhibitor (e.g., Repatha (evolocumab))

| Product Name: Praluent | |
|------------------------|--|
| Diagnosis | Homozygous Familial Hypercholesterolemia |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of homozygous familial hypercholesterolemia (HoFH) as confirmed by submission of medical records (e.g., chart notes, laboratory values) documenting BOTH of the following:*

1.1 ONE of the following:

- Pre-treatment LDL-C (low-density lipoprotein cholesterol) greater than 500 mg/dL (milligrams per deciliter)
- Treated LDL-C greater than 300 mg/dL

AND

1.2 ONE of the following:

- Xanthoma before 10 years of age
- Evidence of heterozygous familial hypercholesterolemia (HeFH) in both parents

2 - Used as an adjunct to a low-fat diet and exercise

AND

3 - Patient is receiving other lipid-lowering therapy (e.g., statin, ezetimibe, LDL [low-density lipoprotein] apheresis)

AND

- **4** Prescribed by ONE of the following:
 - Cardiologist
 - Endocrinologist
 - Lipid specialist

AND

5 - Not used in combination with another proprotein convertase subtilisin/kexin type 9 (PCSK9) inhibitor (e.g., Repatha (evolocumab))

| *Results of prior genetic testing can be submitted as confirmation of di agnosis of HoFH. |
|---|
| |

| Product Name: Praluent | |
|------------------------|--|
| Diagnosis | Homozygous Familial Hypercholesterolemia |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient continues to receive other lipid-lowering therapy (e.g., statin, LDL apheresis)

| 2 - Patient is continuing a low-fat diet and exercise regimen |
|--|
| AND |
| 3 - Prescribed by ONE of the following: |
| Cardiologist Endocrinologist Lipid specialist |
| AND |
| 4 - Submission of medical records (e.g. chart notes, laboratory values) documenting low density lipoprotein cholesterol (LDL-C) reduction while on Praluent therapy |
| AND |
| 5 - Not used in combination with another proprotein convertase subtilisin/kexin type 9 (PCSK9) inhibitor (e.g., Repatha (evolocumab)) |

| Date | Notes |
|----------|---|
| 2/3/2022 | Added criteria for HoFH indication, removed 'prescriber attestation' cr iterion from all sections |

Preferred Drugs- Arizona

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99782 | Preferred Drugs- Arizona |
|----------|--------------------------|
|----------|--------------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Diagnosis | Prior Authorization Administrative Guideline for Preferred Drugs Without Drug-Specific Criteria |
|--------------------------|--|
| Approval Length | 12 month(s) |
| Guideline Type | Administrative |
| | |
| Approval Criteria | |
| 1 - ALL of the following | ng: |
| | |

1.1 ONE of the following:

1.1.1 The requested drug must be used for a Food and Drug Administration (FDA)-approved indication

OR

1.1.2 The use of this drug is supported by information from ONE of the following appropriate compendia of current literature:

- American Hospital Formulary Service Drug Information
- National Comprehensive Cancer Network Drugs and Biologics Compendium
- Thomson Micromedex DrugDex
- Clinical pharmacology

AND

1.2 The drug is being prescribed for a medically accepted indication that is recognized as a covered benefit by the applicable health plans' program

AND

1.3 If the patient is less than FDA minimum age, the prescriber attests they are aware of FDA labeling and feels the treatment with the requested product is medically necessary. (Document rationale for use)

| Notes | Medications used solely for anti-obesity/weight loss, cosmetic (e.g., al opecia, actinic keratosis, vitiligo), erectile dysfunction, and sexual dysf |
|-------|--|
| | unction purposes are NOT medically accepted indications and are NO |
| | T recognized as a covered benefit. Erectile dysfunction drugs (Cialis/T |
| | adalafil) are covered for clinical diagnoses other than ED. |

| Date | Notes |
|----------|-------------------------------------|
| 6/2/2021 | Arizona Medicaid 7.1 Implementation |

Preferred Drugs- Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

| Diagnosis | Prior Authorization Administrative Guideline for Preferred Drugs Without Drug-Specific Criteria |
|---------------------------|--|
| Approval Length | 12 month(s) |
| Guideline Type | Administrative |
| | |
| Approval Criteria | |
| 1 - ALL of the following: | |
| 1.1 ONE of the following: | |

1.1.1 The requested drug must be used for a Food and Drug Administration (FDA)-approved indication

OR

1.1.2 The use of this drug is supported by information from ONE of the following appropriate compendia of current literature:

- American Hospital Formulary Service Drug Information
- National Comprehensive Cancer Network Drugs and Biologics Compendium
- Thomson Micromedex DrugDex
- Clinical pharmacology

AND

1.2 The drug is being prescribed for a medically accepted indication that is recognized as a covered benefit by the applicable health plans' program

AND

1.3 If the patient is less than FDA minimum age, the prescriber attests they are aware of FDA labeling and feels the treatment with the requested product is medically necessary. (Document rationale for use)

| Notes | Medications used solely for anti-obesity/weight loss, cosmetic (e.g., al opecia, actinic keratosis, vitiligo), erectile dysfunction, and sexual dysf |
|-------|--|
| | unction purposes are NOT medically accepted indications and are NO |
| | T recognized as a covered benefit. Erectile dysfunction drugs (Cialis/T |
| | adalafil) are covered for clinical diagnoses other than ED. |

| Date | Notes |
|----------|-------------------------------------|
| 6/2/2021 | Arizona Medicaid 7.1 Implementation |

Pretomanid

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99488 | Pretomanid |
|----------|------------|
| 02-33400 | rictomania |

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

| Product Name: Pretomanid | | |
|---|---------------------|--|
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - One of the following: | | |
| 1.1 Diagnosis of pulmonary extensively drug resistant (XDR) tuberculosis (TB) | | |

OR

1.2 Treatment-intolerant or nonresponsive multidrug-resistant (MDR) tuberculosis (TB)

AND

2 - Pretomanid will be used in combination with bedaquiline and linezolid

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona Standard to Medicaid Arizona Standard for 7 /1 go live |

Prevymis

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99507 Prevymis

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

| Product Name: Prevymis | | |
|--|---------------------|--|
| Approval Length | 6 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Patient is a recipient of an allogeneic hematopoietic stem cell transplant | | |
| | | |
| AND | | |
| | | |

2 - Patient is cytomegalovirus (CMV)-seropositive

AND

3 - Provider attests that Prevymis will be initiated between Day 0 and Day 28 posttransplantation (before or after engraftment) and is being prescribed as prophylaxis and not treatment of CMV infection

| Date | Notes |
|-----------|-------------------------------------|
| 5/21/2021 | Arizona Medicaid 7.1 Implementation |

Procysbi

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99725 Procysbi

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

| Product Name: Procysbi | |
|--|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of nephropathic cystinosis | |

| | AND | |
|---------------------------------------|---|--|
| 2 - Patient is 1 year of age or older | | |
| | AND | |
| 3 - History of failure or | intolerance to Cystagon (immediate-release cysteamine bitartrate)* | |
| Notes | *Note: AZM generally does not consider frequency of dosing and/or la ck of compliance to dosing regimens, an indication of medical necessi ty | |

| Product Name: Procysbi | |
|------------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Procysbi therapy

| Date | Notes |
|-----------|-------------------------------------|
| 5/14/2021 | Arizona Medicaid 7.1 Implementation |

Progesterone - Non-Oral

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99489 | Progesterone - Non-Oral | | | |
|----------|-------------------------|--|-------------|--|
| | | | (. | |

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Crinone, Endometrin | | |
|-----------------------------------|---------------------|--|
| Approval Length | 6 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |

1 - Treatment is for non-infertility use (e.g., secondary amenorrhea, reduce the risk of recurrent spontaneous preterm birth)

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona Standard to Medicaid Arizona Standard for 7 /1 go live |

Promacta

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99637 Promacta

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

| Product Name: Promacta | |
|--|---------------------------------------|
| Diagnosis | Chronic Immune Thrombocytopenia (ITP) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of chronic idiopathic thrombocytopenic purpura (ITP) | |

2 - History of failure, contraindication, or intolerance to at least ONE of the following:

- Corticosteroids
- Immunoglobulins
- Splenectomy

| Product Name: Promacta | | |
|------------------------|---------------------------------------|--|
| Diagnosis | Chronic Immune Thrombocytopenia (ITP) | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |
| | · | |

Approval Criteria

1 - Documentation of positive clinical response to Promacta therapy

| Product Name: Promacta | | |
|------------------------|---|--|
| Diagnosis | Chronic Hepatitis C-Associated Thrombocytopenia | |
| Approval Length | 6 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Diagnosis of chronic hepatitis C-associated thrombocytopenia

AND

- **2** ONE of the following:
 - Planning to initiate and maintain interferon-based treatment

• Currently receiving interferon-based treatment

| Product Name: Promacta | |
|------------------------|---|
| Diagnosis | Chronic Hepatitis C-Associated Thrombocytopenia |
| Approval Length | 6 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Promacta therapy

AND

2 - Patient is currently on antiviral interferon therapy for treatment of chronic hepatitis C

| Product Name: Promacta | |
|------------------------|-----------------------|
| Diagnosis | Aplastic Anemia |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of severe aplastic anemia

AND

2 - One of the following:

2.1 Used in combination with standard immunosuppressive therapy [e.g., Atgam (antithymocyte globulin equine), Thymoglobulin (antithymocyte globulin rabbit), cyclosporine]

OR

2.2 History of failure, contraindication, or intolerance to at least one course of immunosuppressive therapy [e.g., Atgam (antithymocyte globulin equine), Thymoglobulin (antithymocyte globulin rabbit), cyclosporine]

| Product Name: Promacta | |
|------------------------|---------------------|
| Diagnosis | Aplastic Anemia |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Promacta therapy

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Provigil, Nuvigil

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99490 Provigil, Nuvigil

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Provigil, generic modafinil, Brand Nuvigil, generic armodafinil | |
|---|--|
| Diagnosis | Narcolepsy, Obstructive Sleep Apnea, Shift Work Disorder, Idiopathic Hypersomnia (off label) |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - ONE of the following diagnoses:

- Narcolepsy
- Excessive sleepiness due to obstructive sleep apnea

- Excessive sleepiness due to shift work disorder (circadian rhythm sleep disorder, shift work type)
- Idiopathic hypersomnia

AND

2 - If the request is for modafinil, the patient has a history of failure, contraindication, or intolerance to armodafinil

| Product Name: Brand Provigil, generic modafinil, Brand Nuvigil, generic armodafinil | |
|---|---------------------|
| Diagnosis Fatigue due to Multiple Sclerosis (off-label) | |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of multiple sclerosis (MS)

AND

2 - Patient is experiencing fatigue

AND

3 - If the request is for modafinil, the patient has a history of failure, contraindication, or intolerance to armodafinil

| Product Name: Brand Provigil, generic modafinil, Brand Nuvigil, generic armodafinil | |
|---|---|
| Diagnosis | Adjunctive Therapy for the Treatment of Major Depressive Disorder or Bipolar Depression (off-label) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| | |

Approval Criteria

1 - Treatment-resistant depression, defined as BOTH of the following:

1.1 Diagnosis of ONE of the following:

- Major depressive disorder (MDD)
- Bipolar depression

AND

1.2 History of failure, contraindication, or intolerance to at least TWO antidepressants from different classes (e.g., SSRIs [selective serotonin reuptake inhibitors], SNRIs [serotonin-norepinephine reuptake inhibitors], bupropion)

AND

2 - Used as adjunctive therapy

AND

3 - If the request is for modafinil, the patient has a history of failure, contraindication, or intolerance to armodafinil

| Product Name: Brand Provigil, generic modafinil, Brand Nuvigil, generic armodafinil | |
|---|---|
| Diagnosis | Adjunctive Therapy for the Treatment of Major Depressive Disorder or Bipolar Depression (off-label) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to therapy

2 - Used as adjunctive therapy

AND

3 - If the request is for modafinil, the patient has a history of failure, contraindication, or intolerance to armodafinil

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona Standard to Medicaid Arizona Standard for 7 /1 go live |

Pulmonary Arterial Hypertension (PAH) Agents

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-114471 Pulmonary Arterial Hypertension (PAH) Agents

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

Product Name: Preferred Products: Brand Adcirca brand Letairis brand Revatio Suspension brand Tracleer, Sildenafil Citrate Tablets (Generic Revatio). Non Preferred: Generic Alyq tablet, Adempas tablet, Brand Flolan injection, Generic epoprostenol injection, Opsumit tablet, Orenitram tablet, Brand Remodulin injection, Generic treprostinil injection, Brand Revatio tablet, Tracleer tablet for suspension, Tyvaso inhalation solution, Tyvaso Refill inhalation solution, Tyvaso Starter inhalation, Tyvaso DPI, solution, Veletri injection, Ventavis inhalation solution, Brand Revatio Injection, Generic Sildenafil injection, Uptravi

| Diagnosis | Pulmonary Arterial Hypertension |
|-----------------|---------------------------------|
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria 1 - Diagnosis of pulmonary arterial hypertension AND 2 - Pulmonary arterial hypertension is symptomatic AND 3 - One of the following: **3.1** Diagnosis of pulmonary arterial hypertension was confirmed by right heart catheterization OR **3.2** Patient is currently on any therapy for the diagnosis of pulmonary arterial hypertension AND 4 - Prescribed by or in consultation with one of the following: Pulmonologist Cardiologist AND 5 - If the patient is requesting a non preferred product, patient has a history of failure, contraindication or intolerance to at least THREE of the following preferred alternatives* (NOTE: In instances where there are fewer than three preferred alternatives, the patient must have a history of failure, contraindication, or intolerance to ALL of the preferred products.) brand Adcirca • brand Letairis • brand Revatio Suspension ٠

• brand Tracleer

• Sildenafil Citrate Tablets (Generic Revatio)

AND

6 - If the request is for generic Adcirca, patient must have tried and failed brand Adcirca. If the request is for generic LETAIRIS, patient must have tried and failed brand LETAIRIS. If the request is for generic TRACLEER, patient must have tried and failed brand TRACLEER

| Product Name: Adempa | Product Name: Adempas tablet | | |
|---|--|--|--|
| Diagnosis | Chronic Thromboembolic Pulmonary Hypertension (CTEPH) | | |
| Approval Length | 6 month(s) | | |
| Therapy Stage | Initial Authorization | | |
| Guideline Type | Prior Authorization | | |
| | | | |
| Approval Criteria | | | |
| 1 - One of the following | : | | |
| 1.1 Both of the following | ng: | | |
| 1.1.1 Diagnosis of inc hypertension (CTEPH) | 1.1.1 Diagnosis of inoperable or persistent/recurrent chronic thromboembolic pulmonary hypertension (CTEPH) | | |
| AND | | | |
| 1.1.2 CTEPH is symptomatic | | | |
| OR | | | |
| 1.2 Patient is currently on any therapy for the diagnosis of CTEPH | | | |
| AND | | | |
| 2 - Prescribed by or in consultation with one of the following: | | | |
| Pulmonologist | | | |
| | | | |

Cardiologist

Product Name: Preferred Products: Brand Adcirca brand Letairis brand Revatio Suspension brand Tracleer, Sildenafil Citrate Tablets (Generic Revatio). Non Preferred: Generic Alyq tablet, Adempas tablet, Brand Flolan injection, Generic epoprostenol injection, Opsumit tablet, Orenitram tablet, Brand Remodulin injection, Generic treprostinil injection, Brand Revatio tablet, Tracleer tablet for suspension, Tyvaso inhalation solution, Tyvaso Refill inhalation solution, Tyvaso Starter inhalation, Tyvaso DPI, solution, Veletri injection, Ventavis inhalation solution, Brand Revatio Injection, Generic Sildenafil injection, Uptravi

| Diagnosis | All indications listed above |
|-----------------|------------------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to therapy

| Date | Notes |
|-----------|-----------------------------|
| 9/26/2022 | Updated Tyvaso product list |

Pulmozyme

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99638 Pulmozyme

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Pulmozyme | |
|----------------------------------|---------------------|
| Diagnosis | Cystic Fibrosis |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of Cystic Fibrosis | |

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Pyrukynd (mitapivat)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-107467 | Pyrukynd (mitapivat) |
|-----------|----------------------|
|-----------|----------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 6/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Pyrukynd | |
|------------------------|-----------------------|
| Diagnosis | Hemolytic Anemia |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting ALL of the following:

| 1.1 Diagnosis of hemolytic anemia confirmed by the presence of chronic hemolysis (e.g., increased indirect bilirubin, elevated lactated dehydrogenase [LDH], decreased haptoglobin, increased reticulocyte count) | | |
|--|--|--|
| AND | | |
| 1.2 Diagnosis of pyruvate kinase deficiency confirmed by molecular testing of ALL the following mutations on the PKLR gene: | | |
| Presence of at least 2 variant alleles in the pyruvate kinase liver and red blood cell (PKLR) gene, of which at least 1 was a missense variant Patients is not homozygous for the c.1436G>A (p.R479H) variant Patient does not have 2 non-missense variants (without the presence of another missense variant) in the PKLR gene | | |
| AND | | |
| 1.3 Hemoglobin is less than or equal to 10g/dL | | |
| AND | | |
| 1.4 Patient has symptomatic anemia or is transfusion dependent | | |
| AND | | |
| 1.5 Exclusion of other causes of hemolytic anemias (e. g., infections, toxins, drugs) | | |
| AND | | |
| 2 - Prescribed by or in consultation with a hematologist | | |

Г

| Product Name: Pyrukynd | |
|------------------------|------------------|
| Diagnosis | Hemolytic Anemia |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |

| Guideline Type | Prior Authorization |
|--|--|
| | |
| Approval Criteria | |
| clinical response to reduction in transfus transfused during th burden, improveme | edical records (e.g., chart notes, lab work, imaging) documenting positive therapy [e.g., hemoglobin greater than or equal to 1.5g/dL from baseline, sions of greater than or equal to 33% in the number of red blood cell units e fixed dose period compared with the patient's historical transfusion nt in markers of hemolysis from baseline (e.g., bilirubin, lactated H], haptoglobin, reticulocyte count)] |
| | AND |
| 2 - Prescribed by or | in consultation with a hematologist |
| Notes | If the member does not meet the medical necessity reauthorization cri |

| Notes | If the member does not meet the medical necessity reauthorization cri |
|-------|---|
| | teria requirements, a denial should be issued and a 1-month authoriza |
| | tion should be issued one time for Pyrukynd gradual therapy discontin |
| | uation. |

| Date | Notes |
|-----------|-------------|
| 5/24/2022 | New Program |

Radicava (edaravone)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-112929 | Radicava (e | daravone) |
|-----------|-------------|-----------|
|-----------|-------------|-----------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 9/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Radicava IV, Radicava ORS | |
|---|-------------------------------------|
| Diagnosis | Amyotrophic Lateral Sclerosis (ALS) |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of documentation (e.g., chart notes) confirming diagnosis of "definite" or "probable" amyotrophic lateral sclerosis (ALS) per the revised EL Escorial and Airlie House diagnostic criteria

AND

2 - Prescribed by or in consultation with a neurologist with expertise in the diagnosis of ALS

AND

3 - Patient has scores greater than or equal to 2 in all items of the ALS Functional Rating Scale-Revised (ALSFRS-R) criteria at the start of treatment

AND

4 - Patient has a percent (%) forced vital capacity (%FVC) greater than or equal to 80% at the start of treatment

| Product Name: Radica | va IV, Radicava ORS |
|----------------------|-------------------------------------|
| Diagnosis | Amyotrophic Lateral Sclerosis (ALS) |
| Approval Length | 6 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of documentation (e.g., chart notes) confirming positive clinical response to therapy (e.g., slowing in the decline of functional abilities)

AND

2 - Patient is not dependent on invasive ventilation or tracheostomy

| | Date No | Notes |
|--|---------|-------|
|--|---------|-------|

| New rogram | 8/29/2022 | New Program |
|------------|-----------|-------------|
|------------|-----------|-------------|

Ranolazine products

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-110773 Ranolazine products

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 8/15/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Ranexa, generic ranolazine | |
|---|--------------|
| Approval Length | 12 month(s) |
| Guideline Type | Step Therapy |
| | |
| Approval Criteria | |
| History of ONE of the following standard anti-angina treatments: 1.1 One beta-blocker [e.g. Lopressor (metoprolol), Inderal (propranolol)] | |

OR

1.2 One calcium channel blocker [e.g. Procardia XL (nifedipine ER), Cardizem LA/Cardizem CD (diltiazemER)]

OR

1.3 One long acting nitrate therapy [e.g. Imdur (isosorbide mononitrate), Isordil (isosorbide dinitrate), Nitro-Time/Nitro-Dur/Nitro-Bid (nitroglycerin ER)]

AND

2 - For Brand Ranexa requests ONLY: Trial and failure to generic ranolazine (verified via paid pharmacy claims or submission of medical records/chart notes)

| Product Name: Aspruz | yo Sprinkle |
|----------------------|--------------|
| Approval Length | 12 month(s) |
| Guideline Type | Step Therapy |

Approval Criteria

1 - History of ONE of the following standard anti-angina treatments:

1.1 One beta-blocker [e.g. Lopressor (metoprolol), Inderal (propranolol)]

OR

1.2 One calcium channel blocker [e.g. Procardia XL (nifedipine ER), Cardizem LA/Cardizem CD (diltiazemER)]

OR

1.3 One long acting nitrate therapy [e.g. Imdur (isosorbide mononitrate), Isordil (isosorbide dinitrate), Nitro-Time/Nitro-Dur/Nitro-Bid (nitroglycerin ER)]

AND

2 - One of the following:

2.1 Trial and failure to generic ranolazine (verified via paid pharmacy claims or submission of medical records/chart notes)

OR

2.2 One of the following:

- •
- Patient is 8 years of age or younger Patient is unable to swallow the oral tablet (solid formulation) due to swallowing • difficulties

| Date | Notes |
|----------|---|
| 8/4/2022 | Added Aspruzyo Sprinkle as target. Updated guideline name to Rano lazine Products |

Ravicti

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99671 Ravicti

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Ravicti | |
|--|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of urea cycle disorders (UCDs) | |

AND

2 - Inadequate response to ONE of the following:

- Dietary protein restriction
- Amino acid supplementation

AND

3 - Will be used concomitantly with dietary protein restriction and, in some cases, dietary supplements (e.g., essential amino acids, arginine, citrulline, protein-free calorie supplements)

AND

4 - History of failure, contraindication, or intolerance to sodium phenylbutyrate [Buphenyl] *

| Notes | *Note: AZM generally does not consider frequency of dosing and or la |
|-------|--|
| | ck of compliance to dosing regimens an indication of medical necessi |
| | ty |

| Product Name: Ravicti | |
|-----------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Ravicti therapy

AND

2 - Patient is actively on dietary protein restriction and, in some cases, dietary supplements (e.g., essential amino acids, arginine, citrulline, protein-free calorie supplements)

| Date | Notes |
|-----------|-------------------------------------|
| 5/21/2021 | Arizona Medicaid 7.1 Implementation |

Rayos

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99523 Rayos

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Rayos | | |
|---|---------------------|--|
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - ONE of the following: | | |
| 1.1 The requested drug must be used for a Food and Drug Administration (FDA)-approved indication | | |

OR

1.2 The use of this drug is supported by information from ONE of the following appropriate compendia of current literature:

- American Hospital Formulary Service Drug Information
- National Comprehensive Cancer Network Drugs and Biologics Compendium
- Thomson Micromedex DrugDex
- Clinical pharmacology
- United States Pharmacopoeia-National Formulary (USP-NF)

AND

2 - The drug is being prescribed for a medically accepted indication that is recognized as a covered benefit by the applicable health plan's program*

AND

3 - Submission of medical records (e.g. chart notes, laboratory values) or claims history documenting an intolerance to generic prednisone tablets which is unable to be resolved with attempts to minimize the adverse effects where appropriate

AND

4 - History of failure, contraindication, or intolerance to TWO the following:

- Dexamethasone tablet, oral solution
- Hydrocortisone tablet
- Methylprednisolone tablet
- Prednisolone tablet, oral solution

| *Note: Medications used solely for anti-obesity/weight loss, cosmetic (e.g., alopecia, actinic keratosis, vitiligo), erectile dysfunction, and sexu al dysfunction purposes are NOT medically accepted indications and are NOT recognized as a covered benefit. Erectile dysfunction drugs (Cialis/Tadalafil) are covered for clinical diagnoses other than ED. |
|---|
| Clairs/Tadalalli) are covered for clinical diagnoses other than ED. |

| Date | Notes |
|-----------|-------------------------------------|
| 5/14/2021 | Arizona Medicaid 7.1 Implementation |

Reblozyl (luspatercept-aamt)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-113533 Reblozyl (luspatercept-aamt)

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 9/8/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Reblozyl | |
|------------------------|-----------------------|
| Diagnosis | Beta Thalassemia |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - One of the following:

1.1 Both of the following:

1.1.1 Diagnosis of beta thalassemia major

AND

1.1.2 Patient requires regular red blood cell (RBC) transfusions

OR

1.2 Diagnosis of transfusion-dependent beta thalassemia

AND

2 - Prescribed by or in consultation with one of the following:

- Hematologist
- Oncologist

| Product Name: Reblozyl | |
|------------------------|---------------------|
| Diagnosis | Beta Thalassemia |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of a positive clinical response to therapy (e.g., reduction in RBC transfusion burden)

| Product Name: Reblozyl | |
|------------------------|--|
| Diagnosis | Myelodysplastic Syndromes, Myelodysplastic/Myeloproliferative Neoplasm (MDS-RS, MDS/MPN-RS-T) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |

| Guideline Type | Prior Authorization |
|---|--|
| | |
| Approval Criteria | |
| 1 - One of the following | diagnoses: |
| 1.1 Very low-to interm | ediate-risk myelodysplastic syndrome with ring sideroblasts (MDS-RS) |
| | OR |
| 1.2 Myelodysplastic of thrombocytosis (MDS/N | r myeloproliferative neoplasm with ring sideroblasts and /IPN-RS-T) |
| | AND |
| 2 - Patient has failed ar (darbepoetin)] | n erythropoiesis stimulating agent [e.g., Epogen (epoetin alfa), Aranesp |
| | AND |
| 3 - Patient requires trar | nsfusions of 2 or more red blood cell (RBC) units over 8 weeks |
| AND | |
| 4 - Prescribed by or in consultation with one of the following: | |
| HematologistOncologist | |

| Product Name: Reblozyl | |
|------------------------|--|
| Diagnosis | Myelodysplastic Syndromes, Myelodysplastic/Myeloproliferative Neoplasm (MDS-RS, MDS/MPN-RS-T) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of a positive clinical response to therapy (e.g., RBC transfusion independence, improvement in hemoglobin levels)

| Date | Notes |
|----------|---|
| 9/8/2022 | Updated guideline name, removed reference and end notes. No clini cal criteria changes. |

Recorlev (levoketoconazole)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-102891 Recorlev (levoketoconazole)

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: |
|-----------------|
|-----------------|

1. Criteria

| Product Name: Recorlev | |
|------------------------|-----------------------|
| Diagnosis | Cushing's Disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | · |

Approval Criteria

1 - Both of the following:

1.1 Diagnosis of Cushing's disease

AND

1.2 ONE of the following:

- Patient is not a candidate for pituitary surgery
- Pituitary surgery has not been curative

| Product Name: Recorlev | |
|------------------------|---------------------|
| Diagnosis | Cushing's Disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| | |

Approval Criteria

1 - Documentation of positive response to therapy

| Product Name: Recorlev | |
|------------------------|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Recorlev will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Recorlev | |
|------------------------|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |

| Therapy Stage | Reauthorization |
|--|---------------------|
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Documentation of positive clinical response to therapy | |

| Date | Notes |
|----------|--|
| 2/3/2022 | New Program (mirrors Isturisa PA criteria) |

Rectiv

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99492 Rectiv

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Rectiv | | |
|----------------------|--|--|
| Diagnosis | Pain Associated with Chronic Anal Fissures | |
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |

1 - Diagnosis of moderate to severe pain associated with chronic anal fissures

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Standard to Medicaid Arizona Standard for 7 /1 go live |

Regranex

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-102898 Regranex

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 2/3/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Regranex | |
|--|---------------------|
| Approval Length | 6 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Patient has a lower extremity diabetic neuropathic ulcer | |

2. Revision History

| Date | Notes |
|----------|---------------------------------|
| 2/3/2022 | Removed t/f Santyl prerequisite |

Relistor

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99493 Relistor

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Relistor Injection | |
|----------------------------------|--|
| 12 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Documentation (e.g. chart notes) demonstrating ONE of the following:

1.1 Diagnosis of opioid induced constipation in a patient with advanced illness receiving palliative care

OR

1.2 BOTH of the following:

1.2.1 ONE of the following:

1.2.1.1 Diagnosis of opioid induced constipation with chronic, non-cancer pain

OR

1.2.1.2 Diagnosis of opioid induced constipation in patients with chronic pain related to prior cancer diagnosis or cancer treatment who do not require frequent (e.g., weekly) opioid dosage escalation

AND

1.2.2 ONE of the following:

1.2.2.1 The patient is not able to swallow oral medications

OR

1.2.2.2 BOTH of the following:

- History of failure, contraindication or intolerance to an over-the-counter (OTC) laxative ٠ (document name and date tried)
- History of failure, contraindication or intolerance to Movantik •

| Product Name: Relistor Injection | |
|----------------------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Documentation of positive clinical response to Relistor Injection therapy

| Product Name: Relistor tabs | |
|-----------------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - ONE of the following:

1.1 Diagnosis of opioid induced constipation with chronic, non-cancer pain

OR

1.2 Diagnosis of opioid induced constipation in patients with chronic pain related to prior cancer diagnosis or cancer treatment who do not require frequent (e.g., weekly) opioid dosage escalation

AND

2 - BOTH of the following:

2.1 History of failure, contraindication or intolerance to an over-the-counter (OTC) laxative (document name and date tried)

AND

2.2 History of failure, contraindication or intolerance to Movantik

| Product Name: Relistor tabs | |
|-----------------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Documentation of positive clinical response to Relistor Tablet therapy

2. Revision History

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Standard to Medicaid Arizona Standard for 7 /1 go live |

Repatha

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-103331 Repatha

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 2/3/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Repatha | |
|-----------------------|---|
| Diagnosis | Heterozygous familial hypercholesterolemia (HeFH), Atherosclerotic cardiovascular disease (ASCVD) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - ONE of the following diagnoses:

1.1 Heterozygous familial hypercholesterolemia (HeFH) as confirmed by ONE of the following*:

1.1.1 BOTH of the following:

1.1.1.1 Pre-treatment LDL-C (low-density lipoprotein cholesterol) greater than 190 milligrams per deciliter (mg/dL) (greater than 155 mg/dL if less than 16 years of age)

AND

1.1.1.2 ONE of the following:

- Family history of myocardial infarction in first degree relative less than 60 years of age
- Family history of myocardial infarction in second degree relative less than 50 years of age
- Family history of LDL-C greater than 190 mg/dL in first or second degree relative
- Family history of heterozygous or homozygous familial hypercholesterolemia in first or second degree relative
- Family history of tendinous xanthomata and or arcus cornealis in first or second degree relative

OR

1.1.2 BOTH of the following:

1.1.2.1 Pre-treatment LDL-C greater than 190 mg/dL (greater than 155 mg/dL if less than 16 years of age)

AND

1.1.2.2 Submission of medical records (e.g., chart notes, laboratory values) documenting ONE of the following:

- Functional mutation in LDL (low-density lipoprotein), apoB (Apolipoprotein B), or PCSK9 (Proprotein convertase subtilisin/kexin type 9) gene*
- Tendinous xanthomata
- Arcus cornealis before age 45

1.2 Atherosclerotic cardiovascular disease (ASCVD) as confirmed by ONE of the following:

- Acute coronary syndromes
- History of myocardial infarction
- Stable or unstable angina
- Coronary or other arterial revascularization
- Stroke
- Transient ischemic attack
- Peripheral arterial disease presumed to be of atherosclerotic origin

AND

2 - Submission of medical records (e.g., chart notes, laboratory values) documenting ONE of the following (prescription claims history may be used in conjunction as documentation of medication use, dose, and duration):

2.1 Patient has been receiving at least 12 consecutive weeks of high-intensity statin therapy (i.e. atorvastatin 40-80 mg, rosuvastatin 20-40 mg) and will continue to receive high-intensity statin at maximally tolerated dose

OR

2.2 BOTH of the following:

2.2.1 Patient is unable to tolerate high-intensity statin as evidenced by ONE of the following intolerable and persistent (i.e. more than 2 weeks) symptoms:

- Myalgia (muscle symptoms without CK elevations)
- Myositis (muscle symptoms with CK elevations less than 10 times upper limit of normal [ULN])

AND

2.2.2 ONE of the following:

2.2.2.1 Patient has been receiving at least 12 consecutive weeks of moderate-intensity statin [i.e. atorvastatin 10-20 mg, rosuvastatin 5-10 mg, simvastatin greater than or equal to 20 mg, pravastatin greater than or equal to 40 mg, lovastatin 40 mg, Lescol XL (fluvastatin XL) 80 mg, fluvastatin 40 mg twice daily or Livalo (pitavastatin) greater than or equal to 2 mg] and will continue to receive a moderate-intensity statin at maximally tolerated dose

OR

2.2.2.2 Patient has been receiving at least 12 consecutive weeks of low-intensity statin [i.e. simvastatin 10 mg, pravastatin 10-20 mg, lovastatin 20 mg, fluvastatin 20-40 mg, or Livalo (pitavastatin) 1 mg] therapy and will continue to receive a low-intensity statin at maximally tolerated dose

OR

2.3 Patient is unable to tolerate low or moderate, and high intensity statins as evidenced by ONE of the following:

2.3.1 ONE of the following intolerable and persistent (i.e. more than 2 weeks) symptoms for low or moderate, and high intensity statins:

- Myalgia (muscle symptoms without CK elevations)
- Myositis (muscle symptoms with CK elevations less than 10 times upper limit of normal [ULN])

OR

2.3.2 Patient has a labeled contraindication to all statins as documented in medical records

OR

2.3.3 Patient has experienced rhabdomyolysis or muscle symptoms with statin treatment with CK elevations greater than 10 times ULN

AND

3 - ONE of the following:

3.1 Submission of medical records (e.g., laboratory values) documenting ONE of the following LDL-C values while on maximally tolerated lipid lowering therapy for a minimum of at least 12 weeks within the last 120 days:

• LDL-C greater than or equal to 100 mg/dL with ASCVD

• LDL-C greater than or equal to 130 mg/dL without ASCVD

OR

3.2 BOTH of the following:

3.2.1 Submission of medical records (e.g., laboratory values) documenting ONE of the following LDL-C values while on maximally tolerated lipid lowering therapy for a minimum of at least 12 weeks within the last 120 days:

- LDL-C between 70 mg/dL and 99 mg/dL with ASCVD
- LDL-C between 100 mg/dL and 129 mg/dL without ASCVD

AND

3.2.2 Submission of medical records (e.g., chart notes, laboratory values) documenting ONE of the following [prescription claims history may be used in conjunction as documentation of medication use, dose, and duration]:

- Patient has been receiving at least 12 consecutive weeks of ezetimibe (Zetia) therapy as adjunct to maximally tolerated statin therapy
- Patient has a history of contraindication or intolerance to ezetimibe

AND

4 - Used as an adjunct to a low-fat diet and exercise

AND

5 - Prescribed by ONE of the following:

- Cardiologist
- Endocrinologist
- Lipid specialist

AND

6 - Not used in combination with another proprotein convertase subtilisin/kexin type 9 (PCSK9) inhibitor (e.g., Praluent (alirocumab))

| *Results of prior genetic testing can be submitted as confirmation of di agnosis of HeFH . |
|--|
| |

| Product Name: Repatha | |
|-----------------------|---|
| Diagnosis | Heterozygous familial hypercholesterolemia (HeFH), Atherosclerotic cardiovascular disease (ASCVD) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Patient continues to receive statin at maximally tolerated dose (unless patient has documented inability to take statins)

AND

2 - Patient is continuing a low-fat diet and exercise regimen

AND

- 3 Prescribed by ONE of the following:
 - Cardiologist
 - Endocrinologist
 - Lipid specialist

AND

4 - Submission of medical records (e.g. chart notes, laboratory values) documenting LDL-C (low-density lipoprotein cholesterol) reduction while on Repatha therapy

AND

5 - Not used in combination with another proprotein convertase subtilisin/kexin type 9 (PCSK9) inhibitor (e.g., Praluent (alirocumab))

| Product Name: Repatha | |
|-----------------------|---|
| Diagnosis | Homozygous Familial Hypercholesterolemia (HoFH) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

1 - Diagnosis of homozygous familial hypercholesterolemia (HoFH) as confirmed by submission of medical records (e.g., chart notes, laboratory values) documenting BOTH of the following:*

1.1 ONE of the following:

- Pre-treatment LDL-C (low-density lipoprotein cholesterol) greater than 500 mg/dL (milligrams per deciliter)
- Treated LDL-C greater than 300 mg/dL

AND

1.2 ONE of the following:

- Xanthoma before 10 years of age
- Evidence of heterozygous familial hypercholesterolemia (HeFH) in both parents

AND

2 - Used as an adjunct to a low-fat diet and exercise

AND

3 - Patient is receiving other lipid-lowering therapy (e.g., statin, ezetimibe, LDL [low-density lipoprotein] apheresis)

AND

4 - Prescribed by ONE of the following:

- Cardiologist
- Endocrinologist
- Lipid specialist

AND

5 - Not used in combination with another proprotein convertase subtilisin/kexin type 9 (PCSK9) inhibitor (e.g., Praluent (alirocumab))

| *Results of prior genetic testing can be submitted as confirmation of di |
|--|
| agnosis of HoFH. |

| Product Name: Repatha | |
|-----------------------|--|
| Diagnosis | Homozygous Familial Hypercholesterolemia |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient is continuing a low-fat diet and exercise regimen

AND

2 - Patient continues to receive other lipid-lowering therapy (e.g., statin, LDL apheresis)

AND

3 - Submission of medical records (e.g. chart notes, laboratory values) documenting LDL-C (low-density lipoprotein cholesterol) reduction while on Repatha therapy

AND

4 - Prescribed by ONE of the following:
Cardiologist
Endocrinologist
Lipid Specialist

AND
5 - Not used in combination with another proprotein convertase subtilisin/kexin type 9 (PCSK9) inhibitor (e.g., Praluent (alirocumab))

2. Revision History

| Date | Notes |
|----------|--|
| 2/3/2022 | Added effective date to guideline details, no change to criteria |

Retinal Vascular Disease Agents

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-114852 Retinal Vascular Disease Agents

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/3/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Beovu | |
|---------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting one of the following diagnoses:

• Neovascular (wet) age-related macular degeneration (nAMD)

Diabetic macular edema (DME)

AND

2 - Trial and failure, contraindication, or intolerance to compounded Avastin* prepared by a 503(B) Outsourcing Facility [B]

AND

3 - Prescribed by or in consultation with an ophthalmologist experienced in the treatment of retinal diseases

Notes *Note: Trial and failure of compounded bevacizumab can be accepted as meeting the trial and failure of compounded Avastin requirement

| Product Name: Eylea | |
|---------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting one of the following diagnoses:

- Neovascular (wet) age-related macular degeneration (nAMD)
- Macular edema following retinal vein occlusion (RVO)
- Diabetic macular edema (DME)
- Diabetic retinopathy (DR)

AND

2 - Trial and failure, contraindication, or intolerance to compounded Avastin* prepared by a 503(B) Outsourcing Facility [B]

AND

3 - Prescribed by or in consultation with an ophthalmologist experienced in the treatment of retinal diseases

| Notes | *Note: Trial and failure of compounded bevacizumab can be accepted |
|-------|--|
| | as meeting the trial and failure of compounded Avastin requirement |

| Product Name: Lucentis 0.5mg, Byooviz | |
|---------------------------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting one of the following diagnoses:

- Neovascular (wet) age-related macular degeneration (nAMD)
- Macular edema following retinal vein occlusion (RVO)
- Myopic choroidal neovascularization (mCNV)

AND

2 - Trial and failure, contraindication, or intolerance to compounded Avastin* prepared by a 503(B) Outsourcing Facility [B]

AND

3 - Prescribed by or in consultation with an ophthalmologist experienced in the treatment of retinal diseases

| Product Name: Lucentis 0.3mg | |
|------------------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting one of the following diagnoses:

- Diabetic macular edema (DME)
- Diabetic retinopathy (DR)

AND

2 - Trial and failure, contraindication, or intolerance to compounded Avastin* prepared by a 503(B) Outsourcing Facility [B]

AND

3 - Prescribed by or in consultation with an ophthalmologist experienced in the treatment of retinal diseases

| Product Name: Susvimo | |
|-----------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of neovascular (wet) age-related macular degeneration (nAMD)

AND

2 - Trial and failure, contraindication, or intolerance to compounded Avastin* prepared by a 503(B) Outsourcing Facility [B]

AND

3 - Prescribed by or in consultation with an ophthalmologist experienced in the treatment of retinal diseases

| Notes | *Note: Trial of compounded bevacizumab can be accepted as meetin |
|-------|--|
| | g the trial of compounded Avastin requirement |

| Product Name: Vabysmo | |
|-----------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting one of the following diagnoses:

- Neovascular (wet) age-related macular degeneration (nAMD)
- Diabetic macular edema (DME)

AND

2 - Trial and failure, contraindication, or intolerance to compounded Avastin* prepared by a 503(B) Outsourcing Facility [B]

AND

3 - Prescribed by or in consultation with an ophthalmologist experienced in the treatment of retinal diseases

| Notes | *Note: Trial and failure of compounded bevacizumab can be accepted |
|-------|--|
| | as meeting the trial and failure of compounded Avastin requirement |

| Product Name: Beovu, Byooviz, Eylea, Lucentis, Susvimo, Vabysmo | | |
|---|----------------------------|--|
| Approval Length | oproval Length 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting positive

clinical response to therapy (e.g., Improvement in Best Corrected Visual Acuity (BCVA) compared to baseline, stable vision)

2. Endnotes

- A. Neovascular Age-Related Macular Degeneration (nAMD) may also be referred to as wet or exudative AMD. [1]
- B. Congress established the 503(B) facilities to provide compounded pharmaceuticals for office use without a prescription. 503(B) Outsourcing Facilities are compounding pharmacies that must meet higher federal safety, sterility, and quality control standards. [5,6]

3. Revision History

| Date | Notes |
|-----------|------------------------------------|
| 10/3/2022 | Clarified t/f criteria for Susvimo |

Revcovi - AZ

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99639 Revcovi - AZ

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Revcovi | |
|-----------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| | |

1 - Diagnosis of severe combined immunodeficiency disease (SCID)

AND

2 - Deficiency of adenosine deaminase is confirmed by one of the following:

- Deficiency or absence of adenosine deaminase (ADA) in plasma, lysed erythrocytes, fibroblasts (cultured from amniotic fluid), or chorionic villus
- Increase in deoxyadenosine triphosphate (dATP) levels in erythrocyte lysates compared to laboratory standard
- Decrease in ATP (Adenosine triphosphate) concentration in erythrocytes
- Molecular genetic confirmation of mutations in both alleles of the ADA1 gene
- Positive screening by T cell receptor excision circles (TRECs)

AND

3 - One of the following:

- Patient is not a suitable candidate for hematopoietic cell transplantation (HCT)
- Patient has failed HCT
- Patient is awaiting HCT

AND

4 - Dosing is in accordance with the United States Food and Drug Administration approved labeling

| Product Name: Revcovi | |
|-----------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient has previously received treatment with Revcovi (elapegademase) therapy

AND

2 - Patient has experienced a positive clinical response to therapy (e.g., normalization of plasma ADA activity, erythrocyte dATP levels, improvement of disease symptoms, etc.)

AND

3 - Dosing is in accordance with the United States Food and Drug Administration approved labeling

2. Revision History

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Revlimid (lenalidomide)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-105276 I | Revlimid (lenalidomide) |
|-------------|-------------------------|
|-------------|-------------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 4/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: generic lenalidomide | |
|--|--|
| Diagnosis All indications, Non-Preferred | |
| 12 month(s) | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Trial and failure to Brand Revlimid (verified via paid pharmacy claims or submission of medical records)

| Product Name: Brand Revlimid | |
|------------------------------|--|
| Multiple Myeloma | |
| 12 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting patient has a diagnosis of multiple myeloma

| Product Name: Brand Revlimid | |
|------------------------------|---------------------------------|
| Diagnosis | Myelodysplastic Syndromes (MDS) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting patient has a diagnosis of symptomatic anemia due to myelodysplastic syndrome (MDS) associated with a deletion 5q

OR

2 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting BOTH of the following:

2.1 Patient has a diagnosis of anemia due to myelodysplastic syndrome WITHOUT deletion 5q

AND

2.2 ONE of the following:

2.2.1 Serum erythropoetin levels greater than 500 mU (milliunits)/mL (milliliter)

OR

2.2.2 ALL of the following:

2.2.2.1 Serum erythropoetin levels less than or equal to 500 mU/mL

AND

2.2.2.2 Ring sideroblasts less than 15%

AND

2.2.2.3 ONE of the following:

- Revlimid therapy is in combination with an erythropoietin [e.g., Epogen, Procrit, Retacrit (epoetin alfa)]
- History of failure, contraindication, or intolerance to erythropoietins [e.g., Epogen, Procrit, Retacrit (epoetin alfa)]

OR

2.2.3 ALL of the following:

2.2.3.1 Serum erythropoetin levels less than or equal to 500 mU/mL

AND

2.2.3.2 Ring sideroblasts greater than or equal to 15%

AND

2.2.3.3 No response to an erythropoietin in combination with a granulocyte-colony stimulating factor (G-CSF)

OR

3 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting BOTH of the following:

3.1 Diagnosis of myelodysplastic/myeloproliferative neoplasms (MDS/MPN) overlap neoplasm

AND

3.2 Patient has ring sideroblasts and thrombocytosis (MDS/MPN-RS-T)

| Product Name: Brand Revlimid | |
|------------------------------|-----------------------|
| Diagnosis | B-Cell Lymphomas |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting ONE of the following diagnoses:

- Mantle cell lymphoma (MCL)
- Diffuse large B-cell lymphoma (patients 60 to 80 years old)
- Follicular lymphoma
- Gastric mucosa-associated lymphoid tissue (MALT) lymphoma
- Nodal marginal zone lymphoma
- Non-gastric MALT lymphoma
- Splenic marginal zone lymphoma

OR

2 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting BOTH of the following:

2.1 ONE of the following diagnoses:

- Acquired immunodeficiency syndrome (AIDS)-related B-cell lymphoma
- Castleman's Disease (CD)
- Diffuse large B-cell lymphoma (patients who are less than 60 years old)
- High-grade B-cell lymphoma
- Histologic transformation of marginal zone lymphoma to diffuse large B-cell lymphoma
- Post-transplant lymphoproliferative disorders

AND

2.2 NOT used as first line therapy

| Product Name: Brand Revlimid | |
|------------------------------|-----------------------|
| Diagnosis | Hodgkin Lymphoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting patient has a diagnosis of Hodgkin lymphoma

AND

- 2 Disease is ONE of the following:
 - Relapsed
 - Refractory

AND

3 - Used as third-line or subsequent therapy

| Product Name: Brand Revlimid | |
|------------------------------|----------------------------------|
| Diagnosis | Systemic Light Chain Amyloidosis |

| Approval Length | 12 month(s) |
|-----------------|-----------------------|
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting patient has a diagnosis of systemic light chain amyloidosis

AND

2 - ONE of the following:

2.1 Used in combination with dexamethasone

OR

2.2 Used in combination with dexamethasone and cyclophosphamide

| Product Name: Brand Revlimid | | |
|------------------------------|---|--|
| Diagnosis | Chronic Lymphocytic Leukemia/Small Lymphocytic Lymphoma | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting patient has a diagnosis of chronic lymphocytic leukemia (CLL)/small lymphocytic lymphoma (SLL)

AND

2 - ONE of the following:

• Used post first-line chemoimmunotherapy maintenance therapy

- Used post second-line maintenance therapy
- Used for relapsed or refractory disease

| Product Name: Brand Revlimid | |
|------------------------------|-----------------------------|
| Diagnosis | Primary Cutaneous Lymphomas |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting patient has a diagnosis of ONE of the following:

- Mycosis Fungoides (MF) / Sezary Syndrome (SS)
- Primary cutaneous CD30+ T-cell lymphoproliferative disorders

| Product Name: Brand Revlimid | |
|------------------------------|-----------------------|
| Diagnosis | T-Cell Lymphomas |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting patient has ONE of the following diagnoses:

- Peripheral T-cell lymphoma
- T-cell leukemia/lymphoma
- Hepatosplenic gamma-delta T-cell lymphoma

AND

2 - NOT used as first line therapy

| Product Name: Brand Revlimid | |
|------------------------------|-----------------------|
| Diagnosis | Primary CNS Lymphoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting patient has a diagnosis of primary central nervous system lymphoma

| Product Name: Brand Revlimid | |
|------------------------------|-----------------------------|
| Diagnosis | AIDS-Related Kaposi Sarcoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting patient has diagnosis of AIDS-related Kaposi Sarcoma

AND

2 - Patient is currently being treated with antiretroviral therapy (ART)

AND

3 - NOT used as first line therapy

| Product Name: Brand Revlimid | |
|------------------------------|---|
| Diagnosis | * |

| Approval Length | 12 month(s) |
|-----------------|---------------------|
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting patient does not show evidence of progressive disease while on Revlimid therapy

| omas, Hodgkin Lymphoma, Systemic Light Chain Amyloidosis, Chro c Lymphocytic Leukemia/Small Lymphocytic Lymphoma, Primary Cu | | *Multiple Myeloma, Myelodysplastic Syndromes (MDS), B-Cell Lymph omas, Hodgkin Lymphoma, Systemic Light Chain Amyloidosis, Chroni c Lymphocytic Leukemia/Small Lymphocytic Lymphoma, Primary Cut aneous Lymphomas, T-Cell Lymphomas, Primary CNS Lymphomas, AIDS-Related Kaposi Sarcoma |
|---|--|---|
|---|--|---|

| Product Name: Brand Revlimid | |
|------------------------------|---------------------------------|
| Diagnosis | Myelofibrosis-Associated Anemia |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting patient has a diagnosis of myelofibrosis

AND

2 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting ONE of the following:

2.1 BOTH of the following:

2.1.1 Serum erythropoietin levels less than 500 mU (milliunits)/mL (milliliter)

AND

2.1.2 History of failure, contraindication, or intolerance to erythropoietins [e.g., Procrit (epoetin alfa)]

OR

2.2 Serum erythropoietin levels greater than or equal to 500 mU/mL

| Product Name: Brand Revlimid | |
|------------------------------|---------------------------------|
| Diagnosis | Myelofibrosis-Associated Anemia |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting the patient has evidence of symptom improvement or reduction in spleen/liver volume while on Revlimid

| Product Name: Brand Revlimid | |
|------------------------------|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Revlimid will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Brand Revlimid | |
|------------------------------|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |

| Therapy Stage | Reauthorization |
|-------------------|---------------------|
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Documentation of positive clinical response to Revlimid therapy

2. Revision History

| Date | Notes |
|-----------|---|
| 3/28/2022 | Added criteria for Non-Preferred generic lenalidomide. Added submis sion of records where applicable. |

Reyvow - Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99548 | Reyvow - Arizona |
|----------|------------------|
|----------|------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Reyvow | |
|----------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of moderate to severe migraine headaches with or without aura

2 - Used for acute treatment of migraine

AND

3 - Patient is 18 years of age or older

AND

4 - Documentation of a one month trial resulting in therapeutic failure, contraindication, or intolerance to THREE of the following:

- naratriptan tablets
- rizatriptan tablets/ODT (oral disintegrating tablets)
- sumatriptan tablets/auto injection/cartridge or Imitrex nasal spray (Brand only)
- zolmitriptan tablets/ODT

AND

5 - Prescribed by or in consultation with one of the following specialists with expertise in the acute treatment of migraine:

- Neurologist
- Pain Specialist
- Headache Specialist*

AND

6 - Prescriber attests to ALL of the following:

- Patient has been informed the use of Reyvow may result in significant CNS impairment, and may impact the patient's ability to drive or operate machinery for 8 hours after each dose
- If used concurrently with a benzodiazepine or other drugs that could potentially cause central nervous system (CNS) depression, the prescriber has acknowledged that they have completed an assessment of increased risk for sedation and other cognitive and/or neuropsychiatric adverse events

| • | The information provided is true and accurate to the best of their knowledge and they |
|---|---|
| | understand that OptumRx may perform a routine audit and request the medical |
| | information necessary to verify the accuracy of the information provided |

7 - Both of the following:

7.1 One of the following

7.1.1 The patient must have a history of therapeutic failure, contraindication, or intolerance to THREE of the following:

- Amitriptyline (Elavil)**
- A beta-blocker (i.e., atenolol, metoprolol, nadolol, propranolol, or timolol)**
- Divalproex sodium [Depakote/Depakote ER (extended-release)]**
- Topiramate (Topamax)**
- VENLAFAXINE [EFFEXOR/EFFEXOR XR (EXTENDED-RELEASE)]**

OR

7.1.2 The patient must be currently treated with one of the following prophylactic therapies unless there is a contraindication or intolerance to ALL:

- Amitriptyline (Elavil)**
- A beta-blocker (i.e., atenolol, metoprolol, nadolol, propranolol, or timolol)**
- Divalproex sodium [Depakote/Depakote ER (extended-release)]**
- Topiramate (Topamax)**
- Venlafaxine [Effexor/Effexor XR (extended-release)]**

AND

7.2 Both of the Following

7.2.1 History of a therapeutic failure after 3 month trial, contraindication, or intolerance to two of the following biologic calcitonin gene-related peptide receptor (CGRP) antagonists for preventive treatment of migraine

- Ajovy (fremanezumab)
- Emgality (galcanezumab)
- Aimovig (erenumab)

7.2.2 History of a therapeutic failure, contraindication, or intolerance to Ubrelvy

| Notes | *Headache specialists are physicians certified by the United Council f |
|-------|--|
| | or Neurologic Subspecialties (UCNS) **Drugs may require PA |

| Product Name: Reyvow | |
|----------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to therapy

AND

2 - Prescribed by or in consultation with one of the following specialists with expertise in the acute treatment of migraine:

- Neurologist
- Pain Specialist
- Headache Specialist*

| Notes | *Headache specialists are physicians certified by the United Council f or Neurologic Subspecialties (UCNS) |
|-------|---|
| | |

| Date | Notes |
|-----------|-------------------|
| 7/13/2021 | Updated Guideline |

Rezurock (belumosudil)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-103329 | Rezurock (belumosudil) |
|-----------|------------------------|
|-----------|------------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 4/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Rezurock | |
|------------------------|-----------------------------------|
| Diagnosis | Chronic graft-versus-host disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of chronic graft-versus-host disease

2 - Trial and failure of two or more lines of systemic therapy (e.g., corticosteroids, mycophenolate, etc.)

AND

3 - Prescribed by or in consultation with one of the following:

- Hematologist
- Oncologist
- Physician experienced in the management of transplant patients

| Product Name: Rezurock | |
|-----------------------------------|--|
| Chronic graft-versus-host disease | |
| 12 month(s) | |
| Reauthorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on therapy

| Product Name: Rezurock | |
|------------------------|---|
| Diagnosis | Chronic graft-versus-host disease - Twice daily (BID) Therapy |
| Approval Length | 12 month(s) |
| Guideline Type | Quantity Limit |

Approval Criteria

- **1** Patient is using medication concomitantly with one of the following:
 - Strong CYP3A inducer (e.g., carbamazepine, phenobarbital, phenytoin, rifampin)

• Proton pump inhibitor (e.g., omeprazole, pantoprazole, lansoprazole)

| Date | Notes |
|----------|-------------|
| 2/3/2022 | New Program |

Rhofade

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99494 Rhofade

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Rhof | ict Name: Rhofade | |
|--------------------|---|--|
| Diagnosis | Persistent erythema associated with rosacea | |
| Approval Length | 3 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |

1 - Diagnosis of persistent erythema associated with rosacea

2 - ONE of the following:

2.1 History of a 30 day or longer trial and failure of one of the following:

- metronidazole cream, gel, or lotion
- azelaic acid gel

OR

2.2 Contraindication or intolerance to both of the following:

- metronidazole cream, gel, or lotion
- azelaic acid gel

| sistent erythema associated with rosacea |
|--|
| |
| month(s) |
| authorization |
| or Authorization |
| |

Approval Criteria

1 - Documentation of a positive clinical response to Rhofade therapy

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona Standard to Medicaid Arizona Standard for 7 /1 go live |

Rinvoq (upadacitinib)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 6/24/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Rinvoo | 1 |
|----------------------|---------------------------|
| Diagnosis | Rheumatoid Arthritis (RA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of moderately to severely active rheumatoid arthritis

2 - Prescribed by or in consultation with a rheumatologist

AND

3 - Submission of medical records (e.g., chart notes, lab work, imaging, paid claims history) documenting one of the following:

3.1 Both of the following:

3.1.1 History of failure to a 3 month trial of one non-biologic disease modifying antirheumatic drug (DMARD) (e.g., methotrexate, leflunomide, sulfasalazine, hydroxychloroquine) at maximally indicated doses within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*

AND

3.1.2 History of failure, contraindication or intolerance to ALL of the following preferred drugs**:

- 2 preferred TNF inhibitors [e.g., Avsola (infliximab), Enbrel (etanercept), Humira (adalimumab)]
- Orencia (abatacept)
- Xeljanz (tofacitinib)

OR

3.2 Patient is currently on Rinvoq therapy as documented by claims history or medical records (document drug, date, and duration of trial)***

AND

4 - Not used in combination with other Janus kinase (JAK) inhibitors, biologic DMARDs, or potent immunosuppressants (e.g., azathioprine or cyclosporine)*

| Notes | *Rinvoq may be used with concomitant methotrexate, topical or inhale |
|-------|--|
| | d corticosteroids, and/or low stable dosages of oral corticosteroids (eq |

| | uivalent to 10 mg or less of prednisone daily). **PA may be required ** *Patients requesting initial authorization who were established on ther apy via the receipt of a manufacturer supplied sample at no cost in the prescriber's office or any form of assistance from manufacturer spons ored programs shall be required to meet initial authorization criteria as if patient were new to therapy. |
|--|--|
|--|--|

| Product Name: Rinvoq | |
|----------------------|---------------------------|
| Diagnosis | Rheumatoid Arthritis (RA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting positive clinical response to therapy

AND

2 - Not used in combination with other JAK inhibitors, biologic DMARDs, or potent immunosuppressants (e.g., azathioprine or cyclosporine)*

AND

3 - Prescribed by or in consultation with a rheumatologist

| Notes | *Rinvoq may be used with concomitant methotrexate, topical or inhale |
|-------|--|
| | d corticosteroids, and/or low stable dosages of oral corticosteroids (eq |
| | uivalent to 10 mg or less of prednisone daily). |

| Product Name: Rinvoq | |
|----------------------|---------------------------|
| Diagnosis | Psoriatic Arthritis (PsA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

1 - Diagnosis of active psoriatic arthritis

AND

2 - Prescribed by or in consultation with one of the following:

- Dermatologist
- Rheumatologist

AND

3 - Submission of medical records (e.g., chart notes, lab work, imaging, paid claims history) documenting one of the following:

3.1 History of failure, contraindication or intolerance to ALL of the following preferred drugs**:

- 2 preferred TNF inhibitors [e.g., Avsola (infliximab), Enbrel (etanercept), Humira (adalimumab)]
- Orencia (abatacept)
- Otezla (apremilast)
- Xeljanz (tofacitinib)

OR

3.2 Patient is currently on Rinvoq therapy as documented by claims history or medical records (document drug, date, and duration of trial)***

AND

4 - Not used in combination with other JAK inhibitors, biologic DMARDs, or potent immunosuppressants (e.g., azathioprine or cyclosporine)*

| Notes | *Rinvoq may be used with concomitant methotrexate, topical or inhale |
|-------|--|
| | d corticosteroids, and/or low stable dosages of oral corticosteroids (eq |
| | uivalent to 10 mg or less of prednisone daily). **PA may be required ** |
| | *Patients requesting initial authorization who were established on ther |

| apy via the receipt of a manufacturer supplied sample at no cost in the prescriber's office or any form of assistance from manufacturer spons |
|---|
| ored programs shall be required to meet initial authorization criteria as if patient were new to therapy. |

| Product Name: Rinvoq | |
|----------------------|---------------------------|
| Diagnosis | Psoriatic Arthritis (PsA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting positive clinical response to therapy

AND

2 - Not used in combination with other JAK inhibitors, biologic DMARDs, or potent immunosuppressants (e.g., azathioprine or cyclosporine)*

AND

3 - Prescribed by or in consultation with one of the following:

- Dermatologist
- Rheumatologist

| *Rinvoq may be used with concomitant methotrexate, topical or inhale d corticosteroids, and/or low stable dosages of oral corticosteroids (eq | |
|---|--|
| uivalent to 10 mg or less of prednisone daily). | |

| Product Name: Rinvoq | |
|----------------------|-----------------------------|
| Diagnosis | Ankylosing Spondylitis (AS) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |

| Guideline Type | Prior Authorization | |
|---|---|--|
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of active | ankylosing spondylitis | |
| | | |
| | AND | |
| 2 - Prescribed by or in (| consultation with a rheumatologist | |
| | AND | |
| 3 - Submission of medi documenting one of the | cal records (e.g., chart notes, lab work, imaging, paid claims history) e following: | |
| 3.1 Both of the following: | | |
| 3.1.1 Trial and failure, contraindication, or intolerance to TWO nonsteroidal anti- inflammatory drugs (NSAIDs) (e.g., ibuprofen, naproxen) | | |
| | AND | |
| 3.1.2 History of failure drugs**: | e, contraindication or intolerance to ALL of the following preferred | |
| 2 preferred TNF (adalimumab)] Xeljanz (tofacitin) | ⁻ inhibitors [e.g., Avsola (infliximab), Enbrel (etanercept), Humira nib) | |
| OR | | |
| | y on Rinvoq therapy as documented by claims history or medical g, date, and duration of trial)*** | |
| | AND | |

| 4 - Not used in combination with other JAK inhibitors, biologic DMARDs, or potent immunosuppressants (e.g., azathioprine or cyclosporine)* | |
|---|--|
| Notes | *Rinvoq may be used with concomitant methotrexate, topical or inhale d corticosteroids, and/or low stable dosages of oral corticosteroids (eq uivalent to 10 mg or less of prednisone daily). **PA may be required ** *Patients requesting initial authorization who were established on ther apy via the receipt of a manufacturer supplied sample at no cost in the prescriber's office or any form of assistance from manufacturer spons ored programs shall be required to meet initial authorization criteria as if patient were new to therapy. |

| Product Name: Rinvoq | |
|----------------------|-----------------------------|
| Diagnosis | Ankylosing Spondylitis (AS) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting positive clinical response to therapy

AND

2 - Not used in combination with other JAK inhibitors, biologic DMARDs, or potent immunosuppressants (e.g., azathioprine or cyclosporine)*

AND

3 - Prescribed by or in consultation with a rheumatologist

| Notes | *Rinvoq may be used with concomitant methotrexate, topical or inhale |
|-------|--|
| | d corticosteroids, and/or low stable dosages of oral corticosteroids (eq |
| | uivalent to 10 mg or less of prednisone daily). |

| Product Name: Rinvoq | |
|----------------------|------------------------|
| Diagnosis | Atopic Dermatitis (AD) |

| Approval Length | 6 month(s) | |
|---|--|--|
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| Approval Criteria | | |
| 1 - Diagnosis of modera | ate to severe atopic dermatitis | |
| | AND | |
| 2 - Patient is 12 years of | of age or older | |
| | AND | |
| 3 - Submission of medi | cal records documenting one of the following: | |
| | Involvement of at least 10% body surface area (BSA) SCORing Atopic Dermatitis (SCORAD) index value of at least 25 [A] | |
| | AND | |
| 4 - Prescribed by or in a | consultation with one of the following: | |
| DermatologistAllergist/Immunologist | | |
| AND | | |
| 5 - Submission of medical records (e.g., chart notes, lab work, imaging, paid claims history) documenting one of the following: | | |
| 5.1 Both of the following: | | |
| 5.1.1 Trial and failure of a minimum 30-day supply (14-day supply for topical corticosteroids), contraindication, or intolerance to at least ONE of the following**: | | |
| Medium or highPimecrolimus cr | er potency topical corticosteroid ream | |

| • | Tacrolimus | ointment |
|---|--------------------|------------|
| | 1 4 6 1 6 11 1 4 6 | 0111110110 |

• Eucrisa (crisaborole) ointment

AND

5.1.2 Submission of medical records (e.g., chart notes, lab work, imaging, paid claims history) documenting trial and failure of a minimum 12-week supply of at least one systemic drug product for the treatment of atopic dermatitis (examples include, but are not limited to, Adbry [tralokinumab-ldrm], Dupixent [dupilumab], etc.)**

OR

5.2 Patient is currently on Rinvoq therapy as documented by claims history or medical records (document drug, date, and duration of trial)***

AND

6 - Not used in combination with other JAK inhibitors, biologic immunomodulators (e.g., Dupixent, Adbry), or other immunosuppressants (e.g., azathioprine, cyclosporine)*

Notes *Rinvoq may be used with concomitant methotrexate, topical or inhale d corticosteroids, and/or low stable dosages of oral corticosteroids (eq uivalent to 10 mg or less of prednisone daily). ** PA may be required. ***Patients requesting initial authorization who were established on th erapy via the receipt of a manufacturer supplied sample at no cost in t he prescriber's office or any form of assistance from manufacturer spo nsored programs shall be required to meet initial authorization criteria as if patient were new to therapy.

| Product Name: Rinvoq | |
|----------------------|------------------------|
| Diagnosis | Atopic Dermatitis (AD) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records documenting positive clinical response to therapy as evidenced by at least ONE of the following:

| • | Reduction | in b | odv s | urface | area | involv | /ement | from | baseli | ne |
|---|-----------|-------|-------|--------|------|--------|----------|------|--------|-----|
| • | recucion | 111 D | ouy J | unacc | aica | 111001 | Chilonic | nom | Dascii | 110 |

• Reduction in SCORing Atopic Dermatitis (SCORAD) index value from baseline [A]

AND

2 - Prescribed by or in consultation with one of the following:

- Dermatologist
- Allergist/Immunologist

AND

3 - Not used in combination with other JAK inhibitors, biologic immunomodulators (e.g., Dupixent, Adbry), or other immunosuppressants (e.g., azathioprine, cyclosporine)*

| Notes | *Rinvoq may be used with concomitant methotrexate, topical or inhale |
|-------|--|
| | d corticosteroids, and/or low stable dosages of oral corticosteroids (eq |
| | uivalent to 10 mg or less of prednisone daily). |

| Product Name: Rinvoq | | | |
|----------------------|-------------------------|--|--|
| Diagnosis | Ulcerative Colitis (UC) | | |
| Approval Length | 12 month(s) | | |
| Therapy Stage | Initial Authorization | | |
| Guideline Type | Prior Authorization | | |

Approval Criteria

1 - Diagnosis of moderately to severely active ulcerative colitis

AND

2 - Prescribed by or in consultation with a gastroenterologist

AND

3 - Submission of medical records (e.g., chart notes, lab work, imaging, paid claims history) documenting one of the following:

3.1 Both of the following:

3.1.1 Trial and failure, contraindication, or intolerance to ONE of the following conventional therapies:**

- 6-mercaptopurine
- Aminosalicylate (e.g., mesalamine, olsalazine, sulfasalazine)
- Azathioprine
- Corticosteroids (e.g., prednisone)

AND

3.1.2 History of failure to ALL of the following preferred drugs**:

- 2 preferred TNF inhibitors [e.g., Avsola (infliximab), Enbrel (etanercept), Humira (adalimumab)]
- Xeljanz (tofacitinib)

OR

3.2 Patient is currently on Rinvoq therapy as documented by claims history or medical records (document drug, date, and duration of trial)***

AND

4 - Not used in combination with other JAK inhibitors, biological therapies for UC, or with potent immunosuppressants (e.g., azathioprine, cyclosporine)*

Notes *Rinvoq may be used with concomitant methotrexate, topical or inhale d corticosteroids, and/or low stable dosages of oral corticosteroids (eq uivalent to 10 mg or less of prednisone daily). **PA may be required ** *Patients requesting initial authorization who were established on ther apy via the receipt of a manufacturer supplied sample at no cost in the prescriber's office or any form of assistance from manufacturer spons ored programs shall be required to meet initial authorization criteria as if patient were new to therapy.

Product Name: Rinvoq

| Diagnosis | Ulcerative Colitis (UC) | | | |
|--|---|--|--|--|
| Approval Length | 12 month(s) | | | |
| Therapy Stage | Reauthorization | | | |
| Guideline Type | Prior Authorization | | | |
| Approval Criteria 1 - Submission of medical records documenting positive clinical response to therapy | | | | |
| | | | | |
| AND | | | | |
| 2 - Prescribed by or in consultation with a gastroenterologist | | | | |
| AND | | | | |
| 3 - Not used in combination with other JAK inhibitors, biological therapies for UC, or with potent immunosuppressants (e.g., azathioprine, cyclosporine)* | | | | |
| Notes | *Rinvoq may be used with concomitant methotrexate, topical or inhale d corticosteroids, and/or low stable dosages of oral corticosteroids (eq uivalent to 10 mg or less of prednisone daily). | | | |

2. Background

| | ative potencies of topical cort | icosteroids [5] | |
|-------------------|--------------------------------------|-----------------------|-----------------|
| Class | Drug | Dosage Form | Strength (%) |
| Very high potency | Augmented betamethasone dipropionate | Ointment, gel | 0.05 |
| | Clobetasol propionate | Cream, foam, ointment | 0.05 |
| | Diflorasone diacetate | Ointment | 0.05 |

| | Halobetasol propionate | Cream, ointment | 0.05 |
|-------------------|--------------------------------------|---------------------------------|-------|
| High | Amcinonide | Cream, lotion, ointment | 0.1 |
| Potency | Augmented betamethasone dipropionate | Cream, lotion | 0.05 |
| | Betamethasone dipropionate | Cream, foam, ointment, solution | 0.05 |
| | Desoximetasone | Cream, ointment | 0.25 |
| | Desoximetasone | Gel | 0.05 |
| | Diflorasone diacetate | Cream | 0.05 |
| | Fluocinonide | Cream, gel, ointment, solution | 0.05 |
| | Halcinonide | Cream, ointment | 0.1 |
| | Mometasone furoate | Ointment | 0.1 |
| | Triamcinolone acetonide | Cream, ointment | 0.5 |
| Medium | Betamethasone valerate | Cream, foam, lotion, ointment | 0.1 |
| potency | Clocortolone pivalate | Cream | 0.1 |
| | Desoximetasone | Cream | 0.05 |
| | Fluocinolone acetonide | Cream, ointment | 0.025 |
| | Flurandrenolide | Cream, ointment, lotion | 0.05 |
| | Fluticasone propionate | Cream | 0.05 |
| | Fluticasone propionate | Ointment | 0.005 |
| | Mometasone furoate | Cream, lotion | 0.1 |
| | Triamcinolone acetonide | Cream, ointment, lotion | 0.1 |
| Lower- | Hydrocortisone butyrate | Cream, ointment, solution | 0.1 |
| medium potency | Hydrocortisone probutate | Cream | 0.1 |
| | Hydrocortisone valerate | Cream, ointment | 0.2 |
| | Prednicarbate | Cream | 0.1 |
| | Alclometasone dipropionate | Cream, ointment | 0.05 |

| Low | Desonide | Cream, gel, foam, ointment | 0.05 |
|-------------------|------------------------|-----------------------------------|--------------|
| potency | Fluocinolone acetonide | Cream, solution | 0.01 |
| Lowest potency | Dexamethasone | Cream | 0.1 |
| potency | Hydrocortisone | Cream, lotion, ointment, solution | 0.25, 0.5, 1 |
| | Hydrocortisone acetate | Cream, ointment | 0.5-1 |

| Date | Notes |
|-----------|---|
| 6/23/2022 | Updated criteria to include all approved indications. |

Ruconest-Arizona

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Ruconest | | | |
|------------------------|-----------------------------|--|--|
| Diagnosis | Hereditary Angioedema (HAE) | | |
| Approval Length | 12 month(s) | | |
| Therapy Stage | Initial Authorization | | |
| Guideline Type | Prior Authorization | | |
| | • | | |

Approval Criteria

1 - Diagnosis of hereditary angioedema (HAE) as confirmed by ONE of the following:

| 1.1 C1 inhibitor (C1-INH) deficiency or dysfunction (Type I or II HAE) as documented by one of the following (per laboratory standard): |
|---|
| C1-INH antigenic level below the lower limit of normal C1-INH functional level below the lower limit of normal |
| OR |
| 1.2 HAE with normal C1 inhibitor levels and one of the following: |
| Confirmed presence of a FXII, angiopoietin-1 or plasminogen gene mutation Recurring angioedema attacks that are refractory to high-dose antihistamines with confirmed family history of angioedema |
| AND |
| 2 - Prescribed for the acute treatment of HAE attacks |
| AND |
| 3 - Not used in combination with other products indicated for the acute treatment of HAE attacks (e.g. Berinert, Firazyr) |
| AND |
| 4 - Prescribed by ONE of the following: |
| ImmunologistAllergist |

| Product Name: Rucone | est |
|----------------------|-----------------------------|
| Diagnosis | Hereditary Angioedema (HAE) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

| Approval Criteria |
|---|
| 1 - Documentation of positive clinical response |
| |
| AND |
| |
| 2 - Prescribed for the acute treatment of HAE attacks |
| |
| AND |
| |
| 3 - Not used in combination with other products indicated for the acute treatment of HAE attacks (e.g., Berinert, Firazyr) |
| |
| AND |
| |
| 4 - Prescribed by ONE of the following: |
| Immunologist |
| Allergist |

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Samsca

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99642 Samsca

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Samsca | a, generic tolvaptan |
|--------------------------|---|
| Approval Length | 30 Day(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - One of the following | : |
| | nically significant euvolemic hyponatremia nically significant hypervolemic hyponatremia |

| AND |
|---|
| 2 - Patient has not responded to fluid restriction |
| AND |
| 3 - Treatment has been initiated or re-initiated in a hospital setting prior to discharge |

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Sandostatin

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99705 Sandostatin

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brar | nd Sandostatin, generic octreotide |
|--------------------|------------------------------------|
| Diagnosis | Acromegaly |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of acromegaly

2 - One of the following:

2.1 Inadequate response to one of the following:

- Surgery
- Radiotherapy
- Dopamine agonist (e.g., bromocriptine, cabergoline) therapy

OR

2.2 Not a candidate for all of the following:

- Surgery
- Radiotherapy
- Dopamine agonist (e.g., bromocriptine, cabergoline) therapy

| Product Name: Brand S | Sandostatin, generic octreotide |
|-----------------------|---------------------------------|
| Diagnosis | Acromegaly |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to the requested therapy

| Product Name: Brand S | Sandostatin, generic octreotide |
|-----------------------|---------------------------------|
| Diagnosis | Meningioma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

| Approval Criteria | | |
|--|-----|--|
| 1 - Diagnosis of meningioma | | |
| | | |
| | AND | |
| 2 - Disease is surgically inaccessible | | |
| | | |
| | AND | |
| | | |
| 3 - One of the following: | | |
| Disease is recurrent Disease is programive | | |
| Disease is progressive | | |
| | AND | |
| | | |
| 4 - Additional radiation is not possible | | |

| Product Name: Brand Sandostatin, generic octreotide | |
|---|--|
| Meningioma | |
| 12 month(s) | |
| Reauthorization | |
| Prior Authorization | |
| | |

1 - Patient does not show evidence of progressive disease while on the requested therapy

| Product Name: Brand Sandostatin, generic octreotide | |
|---|--|
| Diagnosis | Neuroendocrine tumors, Pheochromocytoma, Paraganglioma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |

| Guideline Type | Prior Authorization | |
|---|---------------------|--|
| | | |
| Approval Criteria | | |
| 1 - Patient has neuroendocrine tumors [e.g., carcinoid tumors, Islet cell tumors, gastrinomas, glucagonomas, insulinomas, lung tumors, somatostatinomas, tumors of the pancreas, GI (gastrointestinal) tract, lung and thymus, adrenal glands, and vasoactive intestinal polypeptidomas (VIPomas)] | | |
| OR | | |
| 2 - All of the following: | | |
| Diagnosis of Pheochromocytoma or Paraganglioma Disease is locally unresectable or distant metastases Disease is somatostatin receptor positive Presence of symptomatic disease | | |

| Product Name: Brand Sandostatin, generic octreotide | |
|---|--|
| Diagnosis | Neuroendocrine tumors, Pheochromocytoma, Paraganglioma |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Patient does not show evidence of progressive disease while on the requested therapy

OR

2 - Documentation of positive clinical response (e.g. suppression of severe diarrhea, flushing, etc.) to the requested therapy

| Product Name: Brand Sandostatin, generic octreotide | |
|---|-----------------------------|
| Diagnosis | Thymoma or Thymic Carcinoma |

| Approval Length | 12 month(s) | |
|---|--------------------------------------|--|
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of thymoma or thymic carcinoma | | |
| | | |
| AND | | |
| 2 - Used as a second-li | ne therapy for one of the following: | |
| 2.1 Unresectable disease following first-line chemotherapy for potentially resectable locally advanced disease, solitary metastasis, or ipsilateral pleural metastasis | | |
| OR | | |

2.2 Extrathoracic metastatic disease

| Product Name: Brand Sandostatin, generic octreotide | |
|---|-----------------------------|
| Diagnosis | Thymoma or Thymic Carcinoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on the requested therapy

| Product Name: Brand Sandostatin, generic octreotide | |
|---|-----------------------------|
| Diagnosis | Malignant Bowel Obstruction |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

1 - Diagnosis of malignant bowel obstruction

AND

2 - Gut function cannot be maintained

| Product Name: Brand Sandostatin, generic octreotide | |
|---|-----------------------------|
| Diagnosis | Malignant Bowel Obstruction |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to the requested therapy

| Product Name: Brand Sandostatin, generic octreotide | | |
|---|---|--|
| Diagnosis | Diarrhea due to concurrent cancer chemotherapy and/or radiation | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Diagnosis of diarrhea due to concurrent cancer chemotherapy and/or radiation

AND

2 - One of the following:

2.1 Presence of Grade 3 or 4 severe diarrhea

2.2 Patient is in palliative or end of life care

| Product Name: Brand Sandostatin, generic octreotide | |
|---|---|
| Diagnosis | Diarrhea due to concurrent cancer chemotherapy and/or radiation |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to the requested therapy

| Product Name: Brand Sandostatin, generic octreotide | |
|---|---------------------------|
| Diagnosis | HIV/AIDS-Related Diarrhea |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of HIV (human immunodeficiency virus)/AIDS (acquired immunodeficiency syndrome)-related diarrhea

| Product Name: Brand Sandostatin, generic octreotide | | |
|---|---------------------------|--|
| Diagnosis | HIV/AIDS-Related Diarrhea | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |
| | | |
| | | |

1 - Documentation of positive clinical response to the requested therapy

| Product Name: Brand Sandostatin, generic octreotide | | |
|---|--|--|
| Bleeding Gastroesophageal Varices | | |
| 12 month(s) | | |
| Initial Authorization | | |
| Prior Authorization | | |
| | | |

Approval Criteria

1 - Diagnosis of bleeding gastroesophageal varices associated with liver disease

| Product Name: Brand Sandostatin, generic octreotide | | |
|---|-----------------------------------|--|
| Diagnosis | Bleeding Gastroesophageal Varices | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |
| | | |
| | | |

Approval Criteria

1 - Documentation of positive clinical response to the requested therapy

| Product Name: Brand Sandostatin, generic octreotide | |
|---|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Use supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Brand Sandostatin, generic octreotide | | |
|---|---------------------------|--|
| Diagnosis | NCCN Recommended Regimens | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |
| | | |

1 - Documentation of positive clinical response to the requested therapy

| Date | Notes |
|-----------|--------------------|
| 5/11/2021 | 7/1 Implementation |

Sedative Hypnotics - AZM

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-112728 Sedative Hypnotics - AZM

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 8/27/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

Product Name: Non-Preferred Drugs: Brand Ambien, Brand Ambien CR, generic zolpidem ER, Edluar, Brand Intermezzo, generic zolpidem SL tablets, Zolpimist, Belsomra, Dayvigo, estazolam, flurazepam, Brand Halcion, generic triazolam, Brand Lunesta, Quvivic, Brand Restoril, generic temazepam 7.5 mg and 22.5 mg capsules, Brand Rozerem, generic ramelteon, Brand Silenor, generic doxepin, generic zaleplon

| Diagnosis | Non-Preferred |
|-------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - History of failure, contraindication, or intolerance to a trial of at least two of the following preferred agents:*

- Eszopiclone (Generic Lunesta)
- Zolpidem (Generic Ambien)
- Temazepam 15/30mg capsules (Generic Restoril)

AND

2 - For generic ramelteon requests ONLY, patient must have tried and failed Brand Rozerem

Product Name: Brand Ambien, generic zolpidem, Brand Ambien CR, generic zolpidem ER, Edluar, Brand Intermezzo, generic zolpidem SL tablets, Zolpimist, Belsomra, Dayvigo, estazolam, flurazepam, Brand Halcion, generic triazolam, Brand Lunesta, generic eszopiclone, Quvivic, Brand Restoril, generic temazepam, Brand Rozerem, generic ramelteon, Brand Silenor, generic doxepin, generic zaleplon

| Diagnosis | Reject 75: Greater than 1 hypnotic in 30 days |
|-----------------|---|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - The requested medication is being used to adjust the dose of the drug

OR

2 - The requested medication will be used in place of the previously prescribed drug, and not in addition to it

OR

3 - The requested medication dosage form will be used in place of the previously prescribed medication dosage form, and not in addition to it

4 - The physician attests they are aware of the multiple sedative hypnotics prescribed to the patient and feels treatment with both medications is medically necessary (Document rationale for use)

Product Name: Brand Ambien, generic zolpidem, Brand Ambien CR, generic zolpidem ER, Edluar, Brand Intermezzo, generic zolpidem SL tablets, Zolpimist, Belsomra, Dayvigo, estazolam, flurazepam, Brand Halcion, generic triazolam, Brand Lunesta, generic eszopiclone, Quvivic, Brand Restoril, generic temazepam, Brand Rozerem, generic ramelteon, Brand Silenor, generic doxepin, generic zaleplon

| Diagnosis | Requests for Patients less than 6 years of age |
|-----------------|--|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - The patient is unresponsive to other treatment modalities, unless contraindicated (i.e. other medications or behavioral modification attempted)

AND

2 - The physician attests that the requested medication is medically necessary. (Document rationale for use)

2. Revision History

| Date | Notes |
|-----------|---|
| 8/26/2022 | Updated NP product list, removed preferred agents listed incorrectly. Added Dayvigo as NP target. Combined criteria for all NP drugs into one category. |

Serevent Diskus - Arizona

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Serevent Diskus | |
|-------------------------------|---------------------|
| Diagnosis | Asthma |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of asthma | |

AND 2 - Patient is 4 years of age or older AND

3 - Patient is also receiving treatment with an inhaled corticosteroid

| Product Name: Sere | event Diskus |
|---|----------------------------------|
| Diagnosis | Exercise-Induced Bronchospasm |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| Approval Criteria 1 - Diagnosis of exe | rcise-induced bronchospasm (EIB) |
| | AND |
| 2 - Being used for p | revention |
| | AND |
| 3 - Patient is 4 years | s of age or older |

| Product Name: Serevent Diskus | |
|---|--|
| Bronchospasm associated with chronic obstructive pulmonary disease (COPD) | |
| 12 month(s) | |
| Prior Authorization | |
| | |
| | |

Approval Criteria

1 - Diagnosis of bronchospasm associated with chronic obstructive pulmonary disease (COPD)

2. Revision History

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona Standard to Medicaid Arizona Standard for 7 /1 go live |

SGLT-2 Inhibitors - AZM

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-114544 SGLT-2 Inhibitors - AZM

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Steglatro, Segluromet, Invokamet, Invokamet XR, Synjardy, Synjardy XR | | |
|---|---------------------|--|
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - The patient has a diagnosis of type 2 diabetes mellitus | | |
| | | |
| AND | | |
| | | |

2 - History of failure to metformin at a minimum dose of 1500mg daily for 90 days, or contraindication or intolerance to metformin.

AND

3 - History of failure, intolerance, or contraindication to ALL of the following:

- •
- Farxiga Jardiance •
- Invokana •

| Product Name: Xigduo XR | | |
|--|---------------------|--|
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - One of the following | r. | |
| 1.1 Both of the followi | ng: | |
| Diagnosis of type 2 diabetes mellitus History of failure to metformin at a minimum dose of 1500mg daily for 90 days, or contraindication or intolerance to metformin. | | |
| | OR | |
| 1.2 Diagnosis of heart failure (NYHA class II-IV) with reduced ejection fraction | | |
| | OR | |
| 1.3 Diagnosis of chronic kidney disease (CKD) | | |

2. Revision History

| Date | Notes |
|-----------|--|
| 9/26/2022 | Added criteria for Xigduo XR (preferred) |

Short-Acting Opioid Products - AZM

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-112965 Short-Acting Opioid Products - AZM

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 8/30/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

Product Name: Generic butorphanol, generic carisoprodol-aspirin-codeine, generic codeine, generic acetaminophen w/codeine, Brand Tylenol/Codeine. generic butalbital-acetaminophencaffeine w/codeine, Brand Fioricet/codeine, generic butalbital-aspirin-caffeine-codeine, Brand Fiorinal/Codeine, generic morphine, generic hydrocodone/acetaminophen, Lortab, Vicodin HP, Norco, Vicodin ES, Lorcet Plus, Lorcet, Lorcet HD, Brand Xodol, generic hydrocodone ibuprofen, Brand Dilaudid, generic hydromorphone, generic oxycodone, Brand Ro xicodone, Brand Oxaydo, generic oxycodone/acetaminophen, Brand Percocet, Brand Primlev, Brand Prolate, Brand Nalocet, Endocet, generic oxycodone-aspirin, oxycodone-ibuprofen, Brand Opana, generic oxymorphone, generic tramadol-acetaminophen, Nucynta, generic meperidine, Fortigan, generic levorphanol, generic acetaminophen -caffeine-dihydrocodeine, Trezix, Dvorah, generic belladonna alkaloids-opium, opium, Apadaz, benzhydrocodoneacetaminophen*

| Diagnosis | PA REQUIRED for use of MAT and other Opioids (Reject 88) |
|-----------|--|
|-----------|--|

| Guideline Type | DUR |
|--|---|
| | |
| Approval Criteria | |
| | notify the prescriber of the MAT therapy and the prescriber of the MAT concurrent opioid therapy. |
| | AND |
| 2 - The days supply do | pes not exceed 14 days for a surgical procedure. |
| | AND |
| 3 - The days supply do | bes not exceed 5 days for all other requests. |
| | AND |
| 4 - There has not been a previous approval in the last 6 months. | |
| Notes | Approval Length: 14 Days for surgical procedure, 5 Days for all other r equests |

| Product Name: Generic butorphanol, generic carisoprodol-aspirin-codeine, generic codeine, | | | |
|--|---|--|--|
| Brand Tylenol/Codeine, Brand Fioricet/codeine, Brand Fiorinal/Codeine, Lortab, Vicodin HP, | | | |
| | cet Plus, Lorcet, Lorcet HD, Brand Xodol, Brand Dilaudid, , Brand | | |
| | aydo, Brand Percocet, Brand Primlev, Brand Prolate, Brand Nalocet, | | |
| | odone-aspirin, Brand Opana, generic oxymorphone, generic | | |
| | e, Brand Ultram, Synapryn, Brand Ultracet, generic tramadol- | | |
| | , Nucynta, Fortigan, generic levorphanol, generic acetaminophen - | | |
| | caffeine-dihydrocodeine, Trezix, Dvorah, generic belladonna alkaloids-opium, opium, Apadaz, | | |
| benzhydrocodone- ace | benzhydrocodone- acetaminophen* | | |
| Diagnosis | Non-Preferred Reviews ** | | |
| Approval Length | 12 month(s) | | |
| Guideline Type | Prior Authorization | | |
| | | | |
| | | | |
| Approval Criteria | | | |

1 - If the request is for a non-preferred medication the patient must have a history of failure, contraindication or intolerance to a trial of at least FIVE preferred short -acting opioids **.

- hydromorphone (generic Dilaudid)
- meperidine
- morphine sulfate
- oxycodone (generic Roxicodone)
- tramadol (generic Ultram)
- oxycodone w/ acetaminophen (generic Percocet)
- oxycodone-ibuprofen
- acetaminophen w/ codeine
- butalbital-acetaminophen-caffeine w/ codeine (Generic Fioricet)
- butalbital-aspirin-caffeine w/cod (generic Fiorinal)
- hydrocodone-acetaminophen (generic Norco)
- hydrocodone-ibuprofen

| Notes *This section does NOT apply to cough and cold products. | |
|--|--|
|--|--|

Product Name: generic acetaminophen w/codeine, generic butalbital-acetaminophen-caffeine w/codeine, generic butalbital-aspirin-caffeine-codeine, generic morphine, generic hydrocodone/acetaminophen, generic hydrocodone-ibuprofen, generic hydromorphone, generic oxycodone, generic oxycodone/acetaminophen, generic tramadol, generic meperidine

| Diagnosis | PA Required for > 2 Short Acting Opioids |
|----------------|--|
| Guideline Type | Prior Authorization |

Approval Criteria

1 - One of the following:

1.1 The requested medication is being used to adjust the dose of the

OR

1.2 The requested medication will be used in place of the previously prescribed drug, and not in addition to it

OR

1.3 The requested medication dosage form will be used in place of the previously prescribed medication dosage form, and not in addition to it

1.4 The physician attests they are aware of the multiple short-acting opioids prescribed to the patient and feels treatment with all medications is medically necessary (Document rationale for use)

OR

| *This section does NOT apply to cough and cold products. ** Authoriz |
|--|
| ation will be issued for the requested duration, not to exceed 12 mont |
| hs. |

Product Name: Generic butorphanol, generic carisoprodol-aspirin-codeine, generic codeine, generic acetaminophen w/codeine, generic butalbital-acetaminophen-caffeine w/codeine, Brand Fioricet/codeine, generic butalbital-aspirin-caffeine-codeine, Brand Fiorinal/Codeine, generic morphine, generic hydrocodone/acetaminophen, Lortab, Norco, Brand Xodol, generic hydrocodone-ibuprofen, Brand Dilaudid, generic hydromorphone, generic oxycodone, Brand Roxicodone, Brand Oxaydo, generic oxycodone/acetaminophen, Brand Percocet, Brand Primlev, Brand Prolate, Brand Nalocet, Endocet, generic oxycodone-aspirin, Brand Opana, generic oxymorphone, generic pentazocine w/naloxone, Brand Ultram, Synapryn, generic tramadol, Brand Ultracet, generic tramadol-acetaminophen, Nucynta, generic meperidine, Fortigan, generic levorphanol, generic acetaminophen-caffeine-dihydrocodeine, Trezix, generic belladonna alkaloids-opium, opium, Apadaz, benzhydrocodone-acetaminophen

| Diagnosis | Quantity Limit |
|-----------------|----------------|
| Approval Length | 12 month(s) |
| Guideline Type | Quantity Limit |

Approval Criteria

1 - The requested dose cannot be achieved by moving to a higher strength of the product

AND

2 - The requested dose is within FDA (Food and Drug Administration) approved maximum dose per day, where an FDA maximum dose per day exists (See table in background section)

| Notes | *This section does NOT apply to cough and cold products. |
|-------|--|
| | |

Product Name: Generic butorphanol, generic carisoprodol-aspirin-codeine, generic codeine, generic acetaminophen w/codeine, generic butalbital-acetaminophen-caffeine w/codeine, Brand Fioricet/codeine, generic butalbital-aspirin-caffeine-codeine, Brand Fiorinal/Codeine, generic morphine, generic hydrocodone/acetaminophen, Lortab, Norco, Brand Xodol, generic

hydrocodone-ibuprofen, Brand Dilaudid, generic hydromorphone, generic oxycodone, Brand Roxicodone, Brand Oxaydo, generic oxycodone/acetaminophen, Brand Percocet, Brand Primlev, Brand Prolate, Brand Nalocet, Endocet, generic oxycodone-aspirin, Brand Opana, generic oxymorphone, generic pentazocine w/naloxone, Brand Ultram, Synapryn, generic tramadol, Brand Ultracet, generic tramadol-acetaminophen, Nucynta, generic meperidine, Fortigan, generic levorphanol, generic acetaminophen-caffeine-dihydrocodeine, Trezix, generic belladonna alkaloids-opium, opium, Apadaz, benzhydrocodone-acetaminophen

| 0 | Greater than 5 day supply requests for patients 18 years of age and older ** | |
|----------------|--|--|
| Guideline Type | Quantity Limit | |

Approval Criteria

- **1** ONE of the following conditions or care instances:
 - Active oncology diagnosis
 - Hospice care
 - End-of-life care (other than hospice)
 - Palliative care
 - Skilled nursing facility care
 - Traumatic injury, excluding post-surgical procedures
 - Chronic conditions for which the provider has received PA approval
 - Post-surgical procedures

| Approvals are for 6 months for all of the above with the exception of p ost-surgical procedures which can be approved for a 14 day supply. A dults may obtain additional fills without PA if the refill is requested with |
|---|
| in 60 days from the initial fill. |

Product Name: Generic butorphanol, generic carisoprodol-aspirin-codeine, generic codeine, generic acetaminophen w/codeine, generic butalbital-acetaminophen-caffeine w/codeine, Brand Fioricet/codeine, generic butalbital-aspirin-caffeine-codeine, Brand Fiorinal/Codeine, generic morphine, generic hydrocodone/acetaminophen, Lortab, Norco, Brand Xodol, generic hydrocodone-ibuprofen, Brand Dilaudid, generic hydromorphone, generic oxycodone, Brand Roxicodone, Brand Oxaydo, generic oxycodone/acetaminophen, Brand Percocet, Brand Primlev, Brand Prolate, Brand Nalocet, Endocet, generic oxycodone-aspirin, Opana, generic oxymorphone, generic pentazocine w/naloxone, Brand Ultram, Synapryn, generic tramadol, Brand Ultracet, generic tramadol-acetaminophen, Nucynta, generic meperidine, Fortigan, generic levorphanol, generic acetaminophen-caffeine-dihydrocodeine, Trezix, generic belladonna alkaloids-opium, opium, Apadaz, benzhydrocodone-acetaminophen*

| Diagnosis | Greater than 5 day supply requests for patients under 18 years of age** |
|----------------|---|
| Guideline Type | Quantity Limit |

Approval Criteria

1 - ONE of the following conditions or care instances:

- Active oncology diagnosis
- Hospice care
- End-of-life care (other than hospice)
- Palliative care
- Children on opioid wean at time of hospital discharge
- Skilled nursing facility care
- Traumatic injury, excluding post-surgical procedures
- Chronic conditions for which the provider has received PA approval
- Post-surgical procedures

| Approvals are for 6 months for all of the above with the exception of p ost-surgical procedures which can be approved for a 14 day supply. C |
|--|
| hildren and adolescents may obtain additional fills without PA for 5 da ys supply unless the submitted PA supports a longer duration for use. |

Product Name: Generic butorphanol, generic carisoprodol-aspirin-codeine, generic codeine, generic acetaminophen w/codeine, generic butalbital-acetaminophen-caffeine w/codeine, Brand Fioricet/codeine, generic butalbital-aspirin-caffeine-codeine, Brand Fiorinal/Codeine, generic morphine, generic hydrocodone/acetaminophen, Lortab, Norco, Brand Xodol, generic hydrocodone-ibuprofen, Brand Dilaudid, generic hydromorphone, generic oxycodone, Brand Roxicodone, Brand Oxaydo, generic oxycodone/acetaminophen, Brand Percocet, Brand Primlev, Brand Prolate, Brand Nalocet, Endocet, generic oxycodone-aspirin, Opana, generic oxymorphone, generic pentazocine w/naloxone, Brand Ultram, Synapryn, generic tramadol, Brand Ultracet, generic tramadol-acetaminophen, Nucynta, generic meperidine, Fortigan, generic levorphanol, generic acetaminophen-caffeine-dihydrocodeine, Trezix, generic belladonna alkaloids-opium, opium, Apadaz, benzhydrocodone-acetaminophen*

| Diagnosis | Opioid Naïve (Not having filled an opioid in the past 120 days)* |
|-----------|--|
| | Morphine Milligram Equivalents (MME)** MME 50.00 exceeded; PA Required for dosage above 50 MEDD |

Approval Criteria

1 - Opioid naïve members may receive greater than 50 morphine milligram equivalent (MME) based on the following:

1.1 If the request is for 50 MME to 90 MME, ONE of the following (NOTE: If the request exceeds 90 MME please skip this section and proceed to the Exceeding the 90 MME Cumulative Threshold Reviews section):

1.1.1 Diagnosis of ONE of the following:

- Cancer
- End of life pain (including hospice care)
- Palliative care
- Sickle cell anemia

OR

1.1.2 Patient is currently exceeding 50 MME and prescriber attests patient has been on a short-acting opioid in the past 120 days

OR

1.1.3 Document ALL of the following:

- The diagnosis associated with the need for pain management with opioid
- If used in patients with medical comorbidities or if used concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression
- The prescriber has acknowledged that they have completed an addiction risk and risk of overdose assessment
- Prescriber attests the member requires more than 50 MME per day to adequately control pain

| Notes | *This section does NOT apply to cough and cold products. **Approval length for cancer, end of life, palliative care, or sickle cell pain will be i ssued for 12 months. All other approvals will be issued for one month. |
|-------|---|
| | |

Product Name: Generic butorphanol, generic carisoprodol-aspirin-codeine, generic codeine, generic acetaminophen w/codeine, generic butalbital-acetaminophen-caffeine w/codeine, Brand Fioricet/codeine, generic butalbital-aspirin-caffeine-codeine, Brand Fiorinal/Codeine, generic morphine, generic hydrocodone/acetaminophen, Lortab, Norco, Brand Xodol, generic hydrocodone-ibuprofen, Brand Dilaudid, generic hydromorphone, generic oxycodone, Brand Roxicodone, Brand Oxaydo, generic oxycodone/acetaminophen, Brand Percocet, Brand Primlev, Brand Prolate, Brand Nalocet, Endocet, generic oxycodone-aspirin, Opana, generic oxymorphone, generic pentazocine w/naloxone, Brand Ultram, Synapryn, generic tramadol, Brand Ultracet, generic tramadol-acetaminophen, Nucynta, generic meperidine, Fortigan, generic levorphanol, generic acetaminophen-caffeine-dihydrocodeine, Trezix, generic belladonna alkaloids-opium, opium, Apadaz, benzhydrocodone-acetaminophen*

| Diagnosis | Cancer/Hospice/End of Life/ Palliative Care/Skilled Nursing Facility/Traumatic Injury Related Pain Exceeding the 90 MME Cumulative Threshold* | |
|--|---|--|
| Approval Length | 12 month(s) | |
| Guideline Type | Morphine Milligram Equivalents (MME) Reviews** (MME 90.00 exceeded; PA REQUIRED; Dosage Above MEDD Limit) | |
| Approval Criteria 1 - ONE of the followin Active oncology Hospice End-of-life care Palliative care Skilled nursing | / diagnosis (other than hospice) | |
| | y, including burns and excluding post-surgical procedures | |
| Notes | *This section does NOT apply to cough and cold products. ** The auth orization should be entered for an MME of 9999 so as to prevent futur e disruptions in therapy if the patient's dose is increased. | |

Product Name: Generic butorphanol, generic carisoprodol-aspirin-codeine, generic codeine, generic acetaminophen w/codeine, generic butalbital-acetaminophen-caffeine w/codeine, Brand Fioricet/codeine, generic butalbital-aspirin-caffeine-codeine, Brand Fiorinal/Codeine, generic morphine, generic hydrocodone/acetaminophen, Lortab, Norco, Brand Xodol, generic hydrocodone-ibuprofen, Brand Dilaudid, generic hydromorphone, generic oxycodone, Brand Primlev, Brand Oxaydo, generic oxycodone/acetaminophen, Brand Percocet, Brand Primlev, Brand Prolate, Brand Nalocet, Endocet, generic oxycodone-aspirin, Opana, generic oxymorphone, generic pentazocine w/naloxone, Brand Ultram, Synapryn, generic tramadol, Brand Ultracet, generic tramadol-acetaminophen, Nucynta, generic meperidine, Fortigan, generic levorphanol, generic acetaminophen-caffeine-dihydrocodeine, Trezix, generic belladonna alkaloids-opium, opium, Apadaz, benzhydrocodone-acetaminophen*

| Diagnosis | Non-cancer/non-hospice/non-end of life/non-palliative care/non-skilled nursing facility/non-traumatic injury related pain Exceeding the 90 MME Cumulative Threshold* |
|-------------------|--|
| Therapy Stage | Initial Authorization |
| Guideline Type | Morphine Milligram Equivalents (MME) Reviews** (MME 90.00 exceeded; PA REQUIRED; Dosage Above MEDD Limit) |
| Approval Criteria | |

- **1** Prescriber attests to ALL of the following:
 - The information provided is true and accurate to the best of their knowledge and they understand that OptumRx may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided
 - Treatment goals are defined, including estimated duration of treatment
 - Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic intervention
 - Patient has been screened for substance abuse/opioid dependence
 - If used in patients with medical comorbidities or if used concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression

2 - BOTH of the following:

- Patient has tried and failed non-opioid pain medication (document drug name and date of trial)
- Opioid medication doses of less than 90 morphine milligram equivalent (MME) have been tried and did not adequately control pain (document drug regimen or MME and dates of therapy)***

| Notes | *This section does NOT apply to cough and cold products. ** Authoriz ation will be issued for 6 months for non-cancer/non-hospice/non-end- of-life/non-palliative care/non-skilled nursing facility/non-traumatic inju ry related pain related pain up to the current requested MME plus 90 M ME. ***If the member has been established on the requested MME do se for at least 30 days and does not meet the medical necessity au thorization criteria requirements, a denial should be issued and a maxi mum 30 -day authorization may be authorized one time for the reques ted MME dose. |
|-------|--|
|-------|--|

Product Name: Generic butorphanol, generic carisoprodol-aspirin-codeine, generic codeine, generic acetaminophen w/codeine, generic butalbital-acetaminophen-caffeine w/codeine, Brand Fioricet/codeine, generic butalbital-aspirin-caffeine-codeine, Brand Fiorinal/Codeine, generic morphine, generic hydrocodone/acetaminophen, Lortab, Norco, Brand Xodol, generic hydrocodone-ibuprofen, Brand Dilaudid, generic hydromorphone, generic oxycodone, Brand Roxicodone, Brand Oxaydo, generic oxycodone/acetaminophen, Brand Percocet, Brand Primlev, Brand Prolate, Brand Nalocet, Endocet, generic oxycodone-aspirin, Opana, generic oxymorphone, generic pentazocine w/naloxone, Brand Ultram, Synapryn, generic tramadol, Brand Ultracet, generic tramadol-acetaminophen, Nucynta, generic meperidine, Fortigan, generic levorphanol, generic acetaminophen-caffeine-dihydrocodeine, Trezix, generic belladonna alkaloids-opium, opium, Apadaz, benzhydrocodone-acetaminophen*

| Diagnosis | Non-cancer/non-hospice/non-end of life/non-palliative care/non-skilled nursing facility/non-traumatic injury related pain Exceeding the 90 MME Cumulative Threshold* |
|--|---|
| Therapy Stage | Reauthorization |
| Guideline Type | Morphine Milligram Equivalents (MME) Reviews** (MME 90.00 exceeded; PA REQUIRED; Dosage Above MEDD Limit) |
| Annual Criteria | |
| Approval Criteria 1 - Prescriber attests to | o ALL of the following: |
| understand that information nec • Treatment goal | n provided is true and accurate to the best of their knowledge and they t OptumRx may perform a routine audit and request the medical cessary to verify the accuracy of the information provided s are defined, including estimated duration of treatment includes the use of a non-opioid analgesic and/or non-pharmacologic |
| Patient has bee If used in patient benzodiazepine | en screened for substance abuse/opioid dependence nts with medical comorbidities or if used concurrently with a e or other drugs that could potentially cause drug-drug interactions, the acknowledged that they have completed an assessment of increased ory depression |
| | AND |
| 2 - Identify rationale for | r not tapering and discontinuing opioid (Document rationale) |
| | OR |
| | es meaningful improvement in pain and function (Document on or pain score improvement)*** |
| Notes | *This section does NOT apply to cough and cold products. ** Authoriz ation will be issued for 6 months for non-cancer/non-hospice/non-end- of-life/non-palliative care/non-skilled nursing facility/non-traumatic inju ry related pain related pain up to the current requested MME plus 90 |

ry related pain related pain up to the current requested MME plus 90 M ME. *** If the member has been established on the requested MME dose for at least 30 days and does not meet the medical necessity au thorization criteria requirements, a denial should be issued and a maxi m um 30 -day authorization may be authorized one time for the reque sted MME dose.

2. Background

| Active Ingredient | FDA Label Max Daily Doses |
|---|--|
| | |
| Morphine | None |
| Hydromorphone | None |
| Hydrocodone | None |
| Tapentadol | 600mg IR products |
| Oxymorphone | None |
| Oxycodone | None |
| Codeine | 360mg |
| Pentazocine | None |
| Tramadol | 400mg IR products |
| Meperidine | 600mg |
| Butorphanol nasal | None |
| Opium | 4 suppositories/day |
| | Deodorized tincture: 24mg/day Camphorate tincture: 16mg/day |
| Acetaminophen | 4g/day |
| Aspirin | 2080mg/day |
| lbuprofen | 3200mg/day |
| Benzhydrocodone** | None |
| Levorphanol | None |
| *Doses are not considered equia chart. | analgesic and table does not represent a dose conversion |

3. Revision History

| Date | Notes |
|-----------|---|
| 8/29/2022 | Updated Rej 88 section to indicate pt must meet all criteria. |

Signifor

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99643 Signifor

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Signifor | |
|------------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Both of the following:

1.1 Diagnosis of endogenous Cushing's disease (i.e., hypercortisolism is not a result of chronic administration of high dose glucocorticoids)

1.2 One of the following:

- Pituitary surgery has not been curative for the patient Patient is not a candidate for pituitary surgery •
- •

| Product Name: Signifor | |
|------------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| Approval Criteria | |

1 - Documentation of positive clinical response to Signifor therapy

2. Revision History

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Siliq- Arizona

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99706 Siliq- Arizona

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Siliq | |
|---------------------------|-----------------------|
| Diagnosis | Plaque Psoriasis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - One of the following: | |

1.1 Submission of medical records (e.g., chart notes, laboratory values, prescription claims history) documenting ALL of the following:

1.1.1 Diagnosis of chronic moderate to severe plaque psoriasis

AND

1.1.2 Greater than or equal to 3 percent body surface area involvement, palmoplantar, facial, or genital involvement, or severe scalp psoriasis

AND

1.1.3 Both of the following:

1.1.3.1 History of failure to ONE of the following topical therapies, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*:

- Corticosteroids (e.g., betamethasone, clobetasol, desonide)
- Vitamin D analogs (e.g., calcitriol, calcipotriene)
- Tazarotene
- Calcineurin inhibitors (e.g., tacrolimus, pimecrolimus)
- Anthralin
- Coal tar

AND

1.1.3.2 History of failure to a 3 month trial of methotrexate at the maximally indicated dose within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*

AND

1.1.4 History of failure, contraindication, or intolerance to ALL of the following preferred biologic products (document drug, date, and duration of trial)*:

- Humira (adalimumab)
- Enbrel (etanercept)
- Otezla (apremilast)

1.1.5 Patient is not receiving Siliq in combination with ONE of the following:

- Biologic Disease-modifying anti-rheumatic drugs (DMARD) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Cosentyx (secukinumab), Orencia (abatacept)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

1.1.6 Prescribed by or in consultation with a dermatologist

OR

1.2 All of the following:

1.2.1 Patient is currently on Siliq therapy as documented by claims history or medical records (document drug, date, and duration of therapy)

AND

1.2.2 Diagnosis of chronic moderate to severe plaque psoriasis

AND

1.2.3 Patient is not receiving Siliq in combination with ONE of the following:

- Biologic DMARD [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Cosentyx (secukinumab), Orencia (abatacept)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

1.2.4 Prescribed by or in consultation with a dermatologist

| Claims history may be used in conjunction as documentation of drug, date, and duration of trial |
|---|
| |

| Product Name: Siliq | |
|---------------------|---------------------|
| Diagnosis | Plaque Psoriasis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Siliq therapy

AND

- 2 Patient is not receiving Siliq in combination with one of the following:
 - Biologic Disease-modifying anti-rheumatic drugs (DMARD) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Cosentyx (secukinumab), Orencia (abatacept)]
 - Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
 - Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with a dermatologist

2. Revision History

| Date | Notes |
|-----------|--------------------|
| 5/11/2021 | 7/1 Implementation |

Simponi- Arizona

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99707 Simponi- Arizona

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Simponi | |
|-----------------------|---------------------------|
| Diagnosis | Rheumatoid Arthritis (RA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - All of the following:

1.1 Diagnosis of moderately to severely active rheumatoid arthritis (RA)

1.2 Patient is NOT receiving Simponi in combination with ONE of the following:

- Biologic disease modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

1.3 One of the following:

1.3.1 Patient is receiving concurrent therapy with methotrexate (e.g., Rheumatrex, Trexall)

OR

1.3.2 History of failure to a 3 month trial of methotrexate at the maximally indicated dose within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*

AND

1.4 History of failure, contraindication, or intolerance to all of the following:

- Humira (adalimumab)
- Enbrel (etanercept)
- Xeljanz (tofacitinib)

AND

1.5 Prescribed by or in consultation with a rheumatologist

OR

2 - All of the following:

| 2.1 Patient is currently on Simponi therapy as documented by claims history or medical records (document drug, date, and duration of therapy) | |
|---|--|
| | AND |
| 2.2 Diagnosis of mode | erately to severely active RA |
| | AND |
| 2.3 Patient is NOT rec | ceiving Simponi in combination with ONE of the following: |
| Biologic DMARD [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab)] Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)] Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)] | |
| AND | |
| 2.4 Prescribed by or in | n consultation with a rheumatologist |
| Notes | *Claims history may be used in conjunction as documentation of drug date, and duration of trial |
| | |

| Product Name: Simponi | |
|-----------------------|------------------------|
| Diagnosis | Ankylosing Spondylitis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- **1** All of the following:
- **1.1** Diagnosis of active ankylosing spondylitis

1.2 History of failure to two NSAIDs (non-steroidal anti-inflammatory drugs) (e.g., ibuprofen, naproxen) at maximally indicated doses, each used for at least 4 weeks within the last 3 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trials)*

AND

1.3 Patient is NOT receiving Simponi in combination with ONE of the following:

- Biologic disease modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

1.4 History of failure, contraindication, or intolerance to BOTH of the following:

- Humira (adalimumab)
- Enbrel (etanercept)

AND

1.5 Prescribed by or in consultation with a rheumatologist

OR

2 - All of the following:

2.1 Patient is currently on Simponi therapy as documented by claims history or medical records (document drug, date, and duration of therapy)*

AND

2.2 Diagnosis of active ankylosing spondylitis

AND

2.3 Patient is NOT receiving Simponi in combination with ONE of the following:

- Biologic DMARD [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

2.4 Prescribed by or in consultation with a rheumatologist

| Notes | *Claims history may be used in conjunction as documentation of drug, |
|-------|--|
| | date, and duration of trial |

| Product Name: Simponi | |
|-----------------------|--|
| Diagnosis | Rheumatoid Arthritis, Ankylosing Spondylitis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Simponi therapy

AND

2 - Patient is NOT receiving Simponi in combination with ONE of the following:

- Biologic disease modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

3 - Prescribed by or in consultation with a rheumatologist

| Product Name: Simponi | |
|-----------------------|-----------------------|
| Diagnosis | Psoriatic Arthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - All of the following:

1.1 Diagnosis of active psoriatic arthritis

AND

1.2 History of failure to a 3 month trial of methotrexate at the maximally indicated dose within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*

AND

1.3 Patient is NOT receiving Simponi in combination with ONE of the following:

- Biologic disease modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

1.4 History of failure, contraindication, or intolerance to three of the following:

• Humira (adalimumab)

| • | Enbrel | (etanercept) |
|---|--------|--------------|
|---|--------|--------------|

- Otezla (apremilast)
- Xeljanz (tofacitinib)

1.5 Prescribed by or in consultation with one of the following:

- Rheumatologist
- Dermatologist

OR

2 - All of the following:

2.1 Patient is currently on Simponi therapy as documented by claims history or medical records (document drug, date, and duration of therapy)*

AND

2.2 Diagnosis of active psoriatic arthritis

AND

2.3 Patient is NOT receiving Simponi in combination with ONE of the following:

- Biologic DMARD [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

2.4 Prescribed by or in consultation with one of the following:

- Rheumatologist
- Dermatologist

| Notes | *Claims history may be used in conjunction as documentation of drug, date, and duration of trials |
|-------|---|
| | |

| Product Name: Simponi | |
|-----------------------|---------------------|
| Diagnosis | Psoriatic Arthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Simponi therapy

AND

- **2** Patient is NOT receiving Simponi in combination with ONE of the following:
 - Biologic disease modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab)]
 - Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
 - Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with one of the following:

- Rheumatologist
- Dermatologist

| Product Name: Simponi | |
|-----------------------|-----------------------|
| Diagnosis | Ulcerative Colitis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - All of the following:

1.1 Diagnosis of moderately to severely active ulcerative colitis

AND

1.2 One of the following:

1.2.1 Patient is corticosteroid dependent (i.e., an inability to successfully taper corticosteroids without a return of the symptoms of UC)

OR

1.2.2 History of failure to ONE of the following conventional therapies at maximally indicated doses within the last 3 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*:

- Corticosteroids (e.g., prednisone, methylprednisolone, budesonide)
- 6-mercaptopurine (Purinethol)
- Azathioprine (Imuran)
- Aminosalicylates (e.g., mesalamine, sulfasalazine)

AND

1.3 Patient is NOT receiving Simponi in combination with ONE of the following:

- Biologic disease modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

1.4 History of failure, contraindication, or intolerance to Humira (adalimumab)

1.5 Prescribed by or in consultation with a gastroenterologist

OR

2 - All of the following:

2.1 Patient is currently on Simponi therapy as documented by claims history or medical records (document drug, date, and duration of therapy)*

AND

2.2 Diagnosis of moderately to severely active ulcerative colitis

AND

2.3 Patient is NOT receiving Simponi in combination with ONE of the following:

- Biologic DMARD [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

2.4 Prescribed by or in consultation with a gastroenterologist

| Notes | *Claims history may be used in conjunction as documentation of drug, |
|-------|--|
| | date, and duration of trials |

| Product Name: Simponi | |
|-----------------------|--------------------|
| Diagnosis | Ulcerative Colitis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |

| Guideline Type | Prior Authorization | | |
|---|--|--|--|
| | | | |
| Approval Criteria | | | |
| 1 - Documentation of p | 1 - Documentation of positive clinical response to Simponi therapy | | |
| | | | |
| | AND | | |
| 2 - Patient is NOT receiving Simponi in combination with ONE of the following: Biologic disease modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab)] Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)] Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)] | | | |
| | | | |
| AND | | | |
| 3 - Prescribed by or in | consultation with a gastroenterologist | | |

| Date | Notes |
|-----------|--------------------|
| 5/11/2021 | 7/1 Implementation |

Sivextro

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99592 Sivextro

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Sivextro | |
|---|------------------------------------|
| Diagnosis | Skin and Skin Structure Infections |
| Approval Length | 6 Day(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - ONE of the following: | |
| 1.1 For continuation of therapy upon hospital discharge | |

OR

1.2 As continuation of therapy when transitioning from intravenous antibiotics that are shown to be sensitive to the cultured organism for the requested indication.

OR

1.3 ALL of the following:

1.3.1 Diagnosis of acute bacterial skin and skin structure infection (including diabetic foot infections)

AND

1.3.2 ONE of the following diagnoses:

1.3.2.1 BOTH of the following:

- Acute bacterial skin and skin structure infections
- Infection caused by methicillin-resistant Staphylococcus aureus (MRSA) documented by culture and sensitivity report

OR

1.3.2.2 BOTH of the following:

- Empirical treatment of patients with acute bacterial skin and skin structure infections
- Presence of MRSA infection is likely

AND

1.3.3 History of failure, contraindication, or intolerance to linezolid (generic Zyvox)

AND

1.3.4 History of failure, contraindication, or intolerance to ONE of the following antibiotics:

• Sulfamethoxazole-trimethoprim (SMX-TMP)

- A tetracycline
- Clindamycin

OR

1.4 ALL of the following:

1.4.1 Diagnosis of acute bacterial skin and skin structure infection(including diabetic foot infections)

AND

1.4.2 Infection caused by an organism that is confirmed to be or likely to be susceptible to treatment with Sivextro

AND

1.4.3 History of failure, contraindication, or intolerance to linezolid (generic Zyvox)

AND

1.4.4 History of failure, contraindication, or intolerance to TWO of the following antibiotics:

- Dicloxacillin
- A cephalosporin
- A tetracycline
- Amoxicillin/clavulanate
- Clindamycin
- Sulfamethoxazole-trimethoprim (SMX-TMP)
- A fluoroquinolone

| Product Name: Sivextro | |
|------------------------|---------------------|
| Diagnosis | Off-Label Uses |
| Approval Length | 60 Day(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - One of the following:

1.1 For continuation of therapy upon hospital discharge

OR

1.2 As continuation of therapy when transitioning from intravenous antibiotics that are shown to be sensitive to the cultured organism for the requested indication

OR

1.3 BOTH of the following:

1.3.1 The medication is being prescribed by or in consultation with an infectious disease specialist

AND

1.3.2 History of failure, contraindication, or intolerance to linezolid (generic Zyvox), if culture and susceptibility confirm susceptibility.

| Date | Notes |
|------------|---|
| 11/11/2021 | Updated off-label approval duration to 60 days. |

Skyrizi (risankizumab-rzaa)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-114498 | Skyrizi (risankizumab-rzaa) | |
|-----------|-----------------------------|--|
|-----------|-----------------------------|--|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Skyrizi SC 75 mg, 150 mg | |
|--|-----------------------|
| Diagnosis | Plaque Psoriasis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - One of the following: | |

1.1 Submission of medical records (e.g., chart notes, laboratory values) documenting ALL of the following:

1.1.1 Diagnosis of moderate to severe plaque psoriasis

AND

1.1.2 Greater than or equal to 3 percent body surface area involvement, palmoplantar, facial, or genital involvement, or severe scalp psoriasis

AND

1.1.3 BOTH of the following:

1.1.3.1 History of failure to ONE of the following topical therapies, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial):*

- Corticosteroids (e.g., betamethasone, clobetasol, desonide)
- Vitamin D analogs (e.g., calcitriol, calcipotriene)
- Tazarotene
- Calcineurin inhibitors (e.g., tacrolimus, pimecrolimus)
- Anthralin
- Coal tar

AND

1.1.3.2 History of failure to a 3 month trial of methotrexate at the maximally indicated dose within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*

AND

1.1.4 History of failure, contraindication, or intolerance to ALL of the following preferred biologic products (document drug, date, and duration of trial):*

- Humira (adalimumab)
- Enbrel (etanercept)
- Otezla (apremilast)

1.1.5 Prescribed by or in consultation with a dermatologist

OR

1.2 All of the following:

1.2.1 Patient is currently on Skyrizi therapy as documented by claims history or medical records (document drug, date, and duration of therapy)

AND

1.2.2 Diagnosis of chronic moderate to severe plaque psoriasis

AND

1.2.3 Prescribed by or in consultation with a dermatologist

| Notes | *Note: Claims history may be used in conjunction as documentation of |
|-------|--|
| | drug, date, and duration of trial |

| Product Name: Skyrizi SC 75 mg, 150 mg | |
|--|---------------------|
| Diagnosis | Plaque Psoriasis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting positive clinical response to Skyrizi therapy

2 - Prescribed by or in consultation with a dermatologist

| Product Name: Skyrizi SC 75 mg, 150 mg | |
|--|---------------------------|
| Diagnosis | Psoriatic Arthritis (PsA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - One of the following:

1.1 Submission of medical records (e.g., chart notes, laboratory values) documenting ALL of the following:

1.1.1 Diagnosis of active psoriatic arthritis (PsA)

AND

1.1.2 History of failure to a 3 month trial of methotrexate at the maximally indicated dose within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*

AND

1.1.3 History of failure, contraindication, or intolerance to ALL of the following preferred biologic products (document drug, date, and duration of trial):*

- Enbrel (etanercept)
- Humira (adalimumab)
- Xeljanz oral tablet (tofacitinib)

OR

1.2 Both of the following:

| | ntly on Skyrizi therapy as documented by claims history or medical g, date, and duration of therapy) |
|--|--|
| | AND |
| 1.2.2 Diagnosis of ac | tive psoriatic arthritis (PsA) |
| | AND |
| 2 - Prescribed by or in | consultation with one of the following: |
| DermatologistRheumatologist | |
| Notes | *Note: Claims history may be used in conjunction as documentation of drug, date, and duration of trial |

| Product Name: Skyrizi SC 75 mg, 150 mg | |
|--|---------------------------|
| Diagnosis | Psoriatic Arthritis (PsA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

Г

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting positive clinical response to Skyrizi therapy

AND

2 - Prescribed by or in consultation with one of the following:

- •
- Dermatologist Rheumatologist •

| Product Name: Skyrizi | IV | |
|---|----------------------|--|
| Diagnosis | Crohn's Disease (CD) | |
| Approval Length | 3 month(s) | |
| Guideline Type | Prior Authorization | |
| Approval Criteria 1 - Submission of medical records (e.g., chart notes, laboratory values) confirming diagnosis of moderately to severely active Crohn's disease (CD) | | |
| | AND | |
| 2 - Trial and failure, contraindication, or intolerance to one of the following conventional therapies 6-mercaptopurine Azathioprine Methotrexate Corticosteroid (e.g., prednisone) | | |
| | AND | |
| 3 - Will be administered as an intravenous induction dose | | |
| AND | | |
| 4 - Prescribed by or in consultation with a gastroenterologist | | |
| | | |
| Product Name: Skyrizi | | |
| Diagnosis | Crohn's Disease (CD) | |
| Approval Length | 12 month(s) | |

| Approval Length | 12 monun(s) |
|-----------------|-----------------------|
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, laboratory values) confirming diagnosis of moderately to severely active Crohn's disease (CD)

AND

2 - Will be used as a maintenance dose following the intravenous induction doses

AND

3 - Prescribed by or in consultation with a gastroenterologist

| Product Name: Skyrizi SC 360 mg | |
|---------------------------------|----------------------|
| Diagnosis | Crohn's Disease (CD) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting positive clinical response to Skyrizi therapy

AND

2 - Prescribed by or in consultation with a gastroenterologist

| Date | Notes |
|-----------|--|
| 9/26/2022 | Added criteria for CD. Updated product list for all indications. |

Soliris- AZ Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99727 Soliris- AZ

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Soliris | |
|-----------------------|---|
| Diagnosis | Atypical hemolytic uremic syndrome (aHUS) |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation supporting the diagnosis of atypical hemolytic uremic syndrome (aHUS) by ruling out BOTH of the following:

• Shiga toxin E. coli-related hemolytic uremic syndrome (STEC-HUS)*

• Thrombotic thrombocytopenia purpura (TTP) (e.g., rule out ADAMTS13 deficiency)

AND

2 - Laboratory results, signs, and/or symptoms attributed to aHUS (e.g., thrombocytopenia, microangiopathic hemolysis, thrombotic microangiopathy, acute renal failure, etc.)

AND

3 - Patient is treatment naïve with Soliris

AND

 ${\bf 4}$ - Soliris is dosed according to the Food and Drug Administration (FDA) labeled dosing for aHUS

AND

5 - Prescribed by, or in consultation with, a hematologist or nephrologist

| Product Name: Soliris | |
|-----------------------|---|
| Diagnosis | Atypical hemolytic uremic syndrome (aHUS) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient has previously been treated with Soliris

AND

2 - Documentation demonstrating a positive clinical response from baseline (e.g., reduction of plasma exchanges, reduction of dialysis, increased platelet count, reduction of hemolysis)

AND

3 - Soliris is dosed according to the United States Food and Drug Administration (FDA) labeled dosing for atypical hemolytic uremic syndrome (aHUS)

AND

4 - Prescribed by, or in consultation with, a hematologist or nephrologist

| Product Name: Soliris | |
|-----------------------|---|
| Diagnosis | Paroxysmal nocturnal hemoglobinuria (PNH) |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation supporting the diagnosis of paroxysmal nocturnal hemoglobinuria (PNH) that includes BOTH of the following:

- Flow cytometry analysis confirming presence of PNH clones
- Laboratory results, signs, and/or symptoms attributed to PNH (e.g., abdominal pain, anemia, dyspnea, extreme fatigue, smooth muscle dystonia, unexplained/unusual thrombosis, hemolysis/hemoglobinuria, kidney disease, pulmonary hypertension, etc.)

AND

2 - Patient is treatment naïve with Soliris

AND

3 - Soliris is dosed according to the United States Food and Drug Administration (FDA) labeled dosing for PNH

AND

4 - Prescribed by, or in consultation with, ONE of the following:

- Hematologist
- Oncologist

| Product Name: Soliris | |
|-----------------------|---|
| Diagnosis | Paroxysmal nocturnal hemoglobinuria (PNH) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient has previously been treated with Soliris

AND

2 - Documentation demonstrating a positive clinical response from baseline (e.g., increased or stabilization of hemoglobin levels, reduction in transfusions, improvement in hemolysis, decrease in lactate dehydrogenase [LDH], increased reticulocyte count, etc.)

AND

3 - Soliris is dosed according to the United States Food and Drug Administration (FDA) labeled dosing for paroxysmal nocturnal hemoglobinuria (PNH)

AND

4 - Prescribed by, or in consultation with, ONE of the following:

- Hematologist
- Oncologist

| Product Name: Soliris | |
|-----------------------|-------------------------------|
| Diagnosis | Generalized myasthenia gravis |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, laboratory values, etc.) to support the diagnosis of generalized myasthenia gravis (gMG) confirming ALL of the following:

1.1 Patient has not failed a previous course of Soliris therapy

AND

1.2 Positive serologic test for anti-acetylcholine receptor (AChR) antibodies

AND

1.3 ONE of the following:

- History of abnormal neuromuscular transmission test demonstrated by single-fiber electromyography (SFEMG) or repetitive nerve stimulation
- History of positive anticholinesterase test, e.g., edrophonium chloride test
- Patient has demonstrated improvement in myasthenia gravis (MG) signs on oral cholinesterase inhibitors, as assessed by the treating neurologist

AND

1.4 Patient has a Myasthenia Gravis Foundation of America (MGFA) Clinical Classification of class II, III, or IV at initiation of therapy

1.5 Patient has a Myasthenia Gravis-specific Activities of Daily Living scale (MG-ADL) total score greater than or equal to 6 at initiation of therapy

AND

2 - BOTH of the following:

2.1 History of failure of at least TWO immunosuppressive agents over the course of at least 12 months [e.g., azathioprine, methotrexate, cyclosporine, mycophenolate, etc.]

AND

2.2 Patient has required TWO or more courses of plasmapheresis/plasma exchanges and/or intravenous immune globulin for at least the previous 12 months without symptom control

AND

3 - Patient is currently on a stable therapeutic dose (at least 3 to 6 months) of immunosuppressive therapy

AND

4 - Soliris is initiated and titrated according to the United States Food and Drug Administration (FDA) labeled dosing for gMG: up to a maximum of 1200 milligrams every 2 weeks

AND

5 - Prescribed by, or in consultation, with a neurologist

| Product Name: Soliris | |
|-----------------------|-------------------------------|
| Diagnosis | Generalized myasthenia gravis |

| Approval Length | 12 month(s) |
|---|---------------------|
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| Approval Criteria 1 - Patient has previously been treated with Soliris | |
| | AND |
| 2 - Submission of medical records (e.g., chart notes, laboratory tests) to demonstrate a positive clinical response from baseline as demonstrated by ALL of the following: Improvement and/or maintenance of at least a 3 point improvement (reduction in score) in the Myasthenia Gravis Activities of Daily Living (MG-ADL) score from pretreatment baseline Reduction in signs and symptoms of myasthenia gravis Maintenance, reduction, or discontinuation of dose(s) of baseline immunosuppressive therapy (IST) prior to starting Soliris (Note: Add on, dose escalation of IST, or additional rescue therapy from baseline to treat myasthenia gravis or exacerbation of symptoms while on Soliris therapy will be considered as treatment failure) | |
| | AND |
| 3 - Soliris is dosed according to the United States Food and Drug Administration (FDA) labeled dosing for generalized myasthenia gravis (gMG): up to a maximum of 1200 milligrams every 2 weeks | |
| AND | |
| 4 - Prescribed by, or in consultation, with a neurologist | |

| Product Name: Soliris | |
|-----------------------|--|
| Diagnosis | Neuromyelitis optica spectrum disorder (NMOSD) |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, laboratory values, etc.) to support the diagnosis of neuromyelitis optica spectrum disorder (NMOSD) confirming ALL of the following:

1.1 Past medical history of ONE of the following:

- Optic neuritis
- Acute myelitis
- Area postrema syndrome: Episode of otherwise unexplained hiccups or nausea and vomiting
- Acute brainstem syndrome
- Symptomatic narcolepsy or acute diencephalic clinical syndrome with NMOSD-typical diencephalic MRI lesions
- Symptomatic cerebral syndrome with NMOSD-typical brain lesions

AND

1.2 Positive serologic test for anti-aquaporin-4 immunoglobulin G (AQP4-IgG)/NMO-IgG antibodies

AND

1.3 Diagnosis of multiple sclerosis or other diagnoses have been ruled out

AND

2 - Patient has not failed a previous course of Soliris therapy

AND

3 - History of failure of, contraindication, or intolerance to rituximab (Rituxan, Ruxience, Truxima) therapy

AND

4 - One of the following:

4.1 History of at least two relapses during the previous 12 months prior to initiating Soliris

OR

4.2 History of at least three relapses during the previous 24 months, at least one relapse occurring within the past 12 months prior to initiating Soliris

AND

5 - Soliris is initiated and titrated according to the U.S. FDA labeled dosing for NMOSD, up to a maximum of 1200 mg every 2 weeks

AND

6 - Prescribed by, or in consultation with, a neurologist

AND

7 - Patient is NOT receiving Soliris in combination with one of the following:

- Disease modifying therapies for the treatment of multiple sclerosis [e.g., Gilenya (fingolimod), Tecfidera (dimethyl fumarate), Ocrevus (ocrelizumab), etc.]
- Anti-IL6 (interleukin 6) therapy [e.g., Actemra (tocilizumab)]

| Product Name: Soliris | |
|-----------------------|--|
| Diagnosis | Neuromyelitis optica spectrum disorder (NMOSD) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Patient has previously been treated with Soliris

2 - Submission of medical records (e.g., chart notes, laboratory tests) to demonstrate a positive clinical response from baseline as demonstrated by BOTH of the following:

2.1 Reduction in the number and/or severity of relapses or signs and symptoms of neuromyelitis optica spectrum disorder (NMOSD)

AND

2.2 Maintenance, reduction, or discontinuation of dose(s) of any baseline immunosuppressive therapy (IST) prior to starting Soliris. (Note: Add on, dose escalation of IST, or additional rescue therapy from baseline to treat NMOSD or exacerbation of symptoms while on Soliris therapy will be considered as treatment failure)

AND

3 - Soliris is dosed according to the U.S. FDA (Food and Drug Administration) labeled dosing for NMOSD: up to a maximum of 1200 mg every 2 weeks

AND

4 - Prescribed by, or in consultation with, a neurologist

AND

5 - Patient is not receiving Soliris in combination with one of the following:

- Disease modifying therapies for the treatment of multiple sclerosis [e.g., Gilenya (fingolimod), Tecfidera (dimethyl fumarate), Ocrevus (ocrelizumab), etc.]
- Anti-IL6 (interleukin 6) therapy [e.g., Actemra (tocilizumab)]

| Date | Notes |
|----------|-------------------------------------|
| 6/8/2021 | Arizona Medicaid 7.1 Implementation |

Somatuline Depot (lanreotide)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-107414 Somatuline Depot (lanreotide)

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

Effective Date: 5/25/2022

1. Indications

Drug Name: Somatuline Depot (lanreotide)

Acromegaly Indicated for the long-term treatment of acromegalic patients who have had an inadequate response to surgery and/or radiotherapy, or for whom surgery and/or radiotherapy is not an option. The goal of treatment in acromegaly is to reduce growth hormone (GH) and insulin growth factor-1 (IGF-1) levels to normal.

Gastroenteropancreatic Neuroendocrine Tumors (GEP-NET) Indicated for the treatment of adult patients with unresectable, well or moderately differentiated, locally advanced or metastatic gastroenteropancreatic neuroendocrine tumors (GEP-NETs) to improve progression-free survival.

Carcinoid Syndrome Indicated for the treatment of adults with carcinoid syndrome; when used, it reduces the frequency of short-acting somatostatin analog rescue therapy.

Drug Name: Lanreotide Injection

Acromegaly Indicated for the long-term treatment of acromegalic patients who have had an inadequate response to surgery and/or radiotherapy, or for whom surgery and/or radiotherapy is not an option. The goal of treatment in acromegaly is to reduce growth hormone (GH) and

insulin growth factor-1 (IGF-1) levels to normal.

Gastroenteropancreatic Neuroendocrine Tumors (GEP-NETs) Indicated for the treatment of adult patients with unresectable, well or moderately differentiated, locally advanced or metastatic gastroenteropancreatic neuroendocrine tumors (GEP-NETs) to improve progression-free survival.

<u>Off Label Uses:</u> Carcinoid Syndrome [3] Indicated for the treatment of adults with carcinoid syndrome; when used, it reduces the frequency of short-acting somatostatin analog rescue therapy.

2. Criteria

| Product Name: Somatuline Depot, Brand Lanreotide | |
|--|-----------------------|
| Diagnosis | Acromegaly |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of acromegaly

AND

2 - One of the following:

2.1 Inadequate response to one of the following:

- Surgery
- Radiotherapy

OR

2.2 Not a candidate for one of the following:

- Surgery
- Radiotherapy

3 - Prescribed by or in consultation with an endocrinologist

| Product Name: Somatuline Depot, Brand Lanreotide | |
|--|--|
| Acromegaly | |
| 12 month(s) | |
| Reauthorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting positive clinical response to therapy, such as a reduction or normalization of IGF-1/GH level for same age and sex

| Product Name: Somatuline Depot 120mg/0.5mL, Brand Lanreotide 120mg/0.5ml | |
|--|---|
| Diagnosis | Advanced or metastatic gastroenteropancreatic neuroendocrine tumors (GEP-NET) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of gastroenteropancreatic neuroendocrine tumor (GEP-NET)

AND

2 - Disease is one of the following:

- Unresectable, locally advanced
- Metastatic

3 - Prescribed by or in consultation with an oncologist

| Product Name: Somatuline Depot 120mg/0.5mL, Brand Lanreotide 120mg/0.5ml | |
|--|---|
| Diagnosis | Advanced or metastatic gastroenteropancreatic neuroendocrine tumors (GEP-NET) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting patient does not show evidence of progressive disease while on therapy

| Product Name: Somatuline Depot 120mg/0.5mL, Brand Lanreotide 120mg/0.5ml [off-label] | |
|--|-----------------------|
| Diagnosis | Carcinoid Syndrome |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of carcinoid syndrome

AND

2 - Used to reduce the frequency of short-acting somatostatin analog rescue therapy

3 - Prescribed by or in consultation with one of the following:

- Endocrinologist
- Oncologist

| Product Name: Somatuline Depot 120mg/0.5mL, Brand Lanreotide 120mg/0.5ml [off-label] | |
|--|---------------------|
| Diagnosis | Carcinoid Syndrome |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting positive clinical response to therapy

| Date | Notes |
|-----------|--|
| 5/23/2022 | New guideline, mirrors ORx with addition of submission of MR req for both initial and reauth in all sections |

Somavert

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99644 Somavert

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Somavert | |
|------------------------|--|
| Acromegaly | |
| 12 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - All of the following:

1.1 Diagnosis of acromegaly by ONE of the following:

| Serum GH (growth hormone) level greater than 1 ng/mL (nanograms per milliliter) after a 2 hour oral glucose tolerance test (OGTT) at time of diagnosis Elevated serum IGF-1 (Insulin-like growth factor-1) levels (above the age and gender |
|--|
| adjusted normal range as provided by the physician's lab) at time of diagnosis |
| AND |
| 1.2 One of the following: |
| 1.2.1 Inadequate response to one of the following: |
| Surgery Radiation therapy Dopamine agonist (e.g., bromocriptine, cabergoline) therapy |
| OR |
| 1.2.2 Not a candidate for all of the following: |
| Surgery Radiation therapy Dopamine agonist (e.g., bromocriptine, cabergoline) therapy |
| AND |
| 1.3 Inadequate response, intolerance, or contraindication to one of the following somatostatin analogs: |
| Sandostatin (octreotide) or Sandostatin LAR Somatuline Depot (lanreotide) |
| OR |
| 2 - Patient is currently on Somavert therapy for acromegaly |

| Product Name: Somavert | |
|------------------------|------------|
| Diagnosis | Acromegaly |

| Approval Length | 12 month(s) |
|-----------------|---------------------|
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Somavert therapy

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Soriatane

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99496 Soriatane

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Soriatane, Generic acitretin | |
|--|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of severe psoriasis | |

2 - Prescribed or recommended by a dermatologist

AND

3 - One of the following:

3.1 Patient is unresponsive to other therapies (e.g., topical corticosteroids, topical vitamin D analogs, tazarotene, methotrexate)

OR

3.2 Other therapies are contraindicated based on the patient's clinical condition

AND

- **4** One of the following:
 - Greater than or equal to 10% body surface area involvement
 - Palmoplantar, facial, or genital involvement
 - Severe scalp psoriasis

| Product Name: Brand Soriatane, Generic acitretin | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Soriatane therapy

| AND | |
|--|--|
| 2 - Prescribed or recommended by a dermatologist | |

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona Standard to Medicaid Arizona Standard for 7 /1 go live |

Spinraza- Arizona

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99729 Spinraza- Arizona

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Spinraza | |
|------------------------|--|
| 3 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Diagnosis of spinal muscular atrophy (SMA) type I, II, or III made by, or in consultation with, a neurologist with expertise in the diagnosis of SMA

2 - Submission of medical records (e.g., chart notes, laboratory values) confirming both of the following:

2.1 The mutation or deletion of genes in chromosome 5q resulting in one of the following:

- Homozygous gene deletion or mutation (e.g., homozygous deletion of exon 7 at locus 5q13)
- Compound heterozygous mutation (e.g., deletion of SMN1 exon 7 [allele 1] and mutation of SMN1 [allele 2])

AND

2.2 Patient has at least 2 copies of SMN2

AND

3 - Patient is not dependent on invasive ventilation or tracheostomy

AND

4 - Patient is not dependent on use of non-invasive ventilation beyond use for naps and nightime sleep

AND

5 - Submission of medical records (e.g., chart notes, laboratory values) or claims history of the baseline exam of one of the following exams (based on patient age and motor ability) to establish baseline motor ability:

- Hammersmith Infant Neurological Exam Part 2 (HINE-2) (infant to early childhood)
- Hammersmith Functional Motor Scale Expanded (HFMSE)
- Upper Limb Module (ULM) Test (Non ambulatory)
- Children's Hospital of Philadelphia Infant Test of Neuromuscular Disorders (CHOP INTEND)

6 - Prescribed by, or in consultation with, a neurologist with expertise in the treatment of SMA

AND

7 - One of the following:

7.1 Patient has not previously received gene replacement therapy for the treatment of SMA

OR

7.2 One of the following:

7.2.1 Both of the following:

7.2.1.1 Patient recently received gene replacement therapy within the previous 6 months

AND

7.2.1.2 Patient has experienced a declination in clinical status since receipt of gene replacement therapy

OR

7.2.2 Both of the following:

7.2.2.1 Patient has previously received gene replacement therapy

AND

7.2.2.2 Patient has experienced a declination in clinical status that represents a potential abatement of gene therapy efficacy

AND

8 - Spinraza is to be administered intrathecally by, or under the direction of, healthcare professionals experienced in performing lumbar punctures

AND

9 - Spinraza dosing for SMA is within accordance with the United States Food and Drug Administration approved labeling: maximum dosing of 12 milligrams for each loading dose

| Product Name: Spinraza | |
|------------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of spinal muscular atrophy (SMA) type I, II, or III made by, or in consultation with, a neurologist with expertise in the diagnosis of SMA

AND

2 - Submission of medical records (e.g., chart notes, laboratory values) or claims history confirming both of the following:

2.1 The mutation or deletion of genes in chromosome 5q resulting in one of the following:

- Homozygous gene deletion or mutation (e.g., homozygous deletion of exon 7 at locus 5q13)
- Compound heterozygous mutation (e.g., deletion of SMN1 exon 7 [allele 1] and mutation of SMN1 [allele 2])

AND

2.2 Patient has at least 2 copies of SMN2

AND

3 - Patient is not dependent on invasive ventilation or tracheostomy

AND

4 - Patient is not dependent on use of non-invasive ventilation beyond use for naps and nightime sleep

AND

5 - One of the following:

5.1 Patient has not previously received gene replacement therapy for the treatment of SMA

OR

5.2 Both of the following:

5.2.1 Patient has previously received gene replacement therapy

AND

5.2.2 Patient has experienced a declination in clinical status that represented a potential failure or abatement of gene therapy efficacy

AND

6 - Submission of medical records (e.g., chart notes, laboratory values) or claims history with the most recent results (less than 1 month prior to request) documenting a positive clinical response from pretreatment baseline status to Spinraza therapy as demonstrated by one of the following exams:

6.1 Both of the following for Hammersmith Infant Neurological Exam Part 2 (HINE-2) milestones:

6.1.1 One of the following:

 Improvement or maintenance of previous improvement of at least 2 point (or maximal score) increase in ability to kick

| Improvement or maintenance of previous improvement of at least 1 point increase in any other HINE-2 milestone (e.g., head control, rolling, sitting, crawling, etc.), excluding voluntary grasp |
|--|
| AND |
| 6.1.2 One of the following: |
| • The patient exhibited improvement or maintenance of previous improvement in more HINE motor milestones than worsening, from pretreatment baseline (net positive improvement) |
| Achieved and maintained any new motor milestones when they would otherwise be unexpected to do so (e.g., sit unassisted, stand, walk) |
| OR |
| 6.2 One of the following for Hammersmith Functional Motor Scale Expanded (HFMSE): |
| Improvement or maintenance of previous improvement of at least a 3 point increase in score from pretreatment baseline Patient has achieved and maintained any new motor milestone from pretreatment baseline when they would otherwise be unexpected to do so |
| OR |
| 6.3 One of the following for Upper Limb Module (ULM): |
| Improvement or maintenance of previous improvement of at least a 2 point increase in score from pretreatment baseline Patient has achieved and maintained any new motor milestone from pretreatment baseline when they would otherwise be unexpected to do so |
| OR |
| 6.4 One of the following for Children's Hospital of Philadelphia Infant Test of Neuromuscular Disorders (CHOP INTEND): |
| 6.4.1 Improvement or maintenance of previous improvement of at least a 4 point increase in score from pretreatment baseline |

OR

6.4.2 Patient has achieved and maintained any new motor milestone from pretreatment baseline when they would otherwise be unexpected to do so

OR

6.4.3 Both of the following:

- Patient was prescribed Spinraza due to clinical declination after receipt of gene therapy
- Patients clinical status has stabilized after receipt of Spinraza therapy

AND

7 - Prescribed by, or in consultation with, a neurologist with expertise in the treatment of SMA

AND

8 - Spinraza is to be administered intrathecally by, or under the direction of, healthcare professionals experienced in performing lumbar punctures

AND

9 - Spinraza dosing for SMA is within accordance with the United States Food and Drug Administration approved labeling: maximum dosing of 12 milligrams every 4 months, starting 4 months after the last loading dose

2. Revision History

| Date | Notes |
|-----------|--------------------|
| 5/25/2021 | 7/1 Implementation |

Spiriva Respimat

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99569 Spiriva Respimat

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Spiriva Respimat | |
|---|---------------------|
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Requests for Spiriva Respimat should be denied. The plan's preferred product is Spiriva Handihaler | |

Spravato - AZ Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99645 Spravato - AZ

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Spravato | |
|------------------------|---|
| Diagnosis | Major Depressive Disorder (Treatment-Resistant) |
| Approval Length | 3 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient has a confirmed diagnosis of major depressive disorder as defined by the DSM-V (Diagnostic and Statistical Manual of Mental Disorders) criteria and is treatment resistant

2 - Patient is 18 years of age or older

AND

3 - Spravato is prescribed by, or in consultation with, a psychiatric provider

AND

4 - ONE of the following:

4.1 Patient does not have an active substance use disorder (SUD)

OR

4.2 BOTH of the following:

- Patient has an active substance use disorder
- Patient is currently receiving treatment

AND

5 - ONE of the following:

5.1 Patient has experienced an inadequate response during the current depressive episode with BOTH of the following therapies:

5.1.1 Two antidepressants from at least two different classes [must include one of each AHCCCS (Arizona Health Care Cost Containment System) preferred agents: SSRI (selective serotonin reuptake inhibitor), SNRI (serotonin-norepinephrine reuptake inhibitor), or bupropion] having different mechanisms of action at the maximally tolerated labeled dose, each used for at least 4-6 weeks

AND

5.1.2 At least TWO augmentation therapies below for at least 4 weeks: SSRI or SNRI, and a second-generation antipsychotic used concomitantly • (aripiprazole, quetiapine, risperidone, olanzapine) SSRI or SNRI, and lithium used concomitantly • SSRI or SNRI, and liothyronine (T3) used concomitantly • SSRI or SNRI, and mirtazapine • SSRI and bupropion and buspirone • OR 5.2 Patient has active suicidal ideation and urgent symptom control is necessary AND 6 - Spravato is used in combination with an oral antidepressant (e.g., duloxetine, escitalopram, sertraline, venlafaxine) AND 7 - Spravato is administered under the direct supervision of a healthcare provider AND 8 - Provider is certified in the Spravato REMS (risk evaluation and mitigation strategy) program AND 9 - Patient must be monitored by a health care provider for at least 2 hours after administration

| Product Name: Spravato | |
|------------------------|---|
| Diagnosis | Major Depressive Disorder (Treatment-Resistant) |
| Approval Length | 6 month(s) |

| Therapy Stage | Reauthorization | | |
|--|---|--|--|
| Guideline Type | Prior Authorization | | |
| | | | |
| Approval Criteria | | | |
| | 1 - Provider attests that the patient has documented improvement or sustained improvement in depressive symptoms from baseline | | |
| | AND | | |
| 2 - Patient use of Sprav | 2 - Patient use of Spravato is in combination with an oral antidepressant | | |
| | AND | | |
| 3 - Patient administers | 3 - Patient administers Spravato under the direct supervision of a healthcare provider | | |
| | AND | | |
| 4 - Provider is certified in the Spravato REMS (risk evaluation and mitigation strategy) program | | | |
| AND | | | |
| | ue to be monitored by a health care provider certified by the Spravato east 2 hours after administration | | |

| Product Name: Spravato | |
|------------------------|--|
| Diagnosis | Requests for Patients less than 6 years of age |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - The patient is unresponsive to other treatment modalities, unless contraindicated (i.e. other medications or behavioral modification attempted)

AND

2 - The physician attests that the requested medication is medically necessary. (Document rationale for use)

| Product Name: Sprava | Product Name: Spravato | |
|--|---|--|
| Diagnosis | Depressive symptoms in an adult with major depressive disorder (MDD) with acute suicidal ideation or behavior | |
| Approval Length | 1 month(s) | |
| Guideline Type | Prior Authorization | |
| Approval Criteria 1 - Diagnosis of major depressive disorder according to the current Diagnostic and Statistical Manual of Mental Disorders (DSM) (i.e., DSM-5) criteria | | |
| | AND | |
| 2 - Patient is experiencing an acute suicidal ideation or behavior | | |
| AND | | |
| 3 - Patient is receiving newly initiated or optimized oral antidepressant | | |
| | AND | |
| | | |

4 - Provider and/or the provider's healthcare setting is certified in the Spravato REMS (Risk Evaluation and Mitigation Strategy) program

2. Revision History

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Sprycel (dasatinib)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-109039 | Sprycel (| (dasatinib) |
|-----------|-----------|-------------|
|-----------|-----------|-------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 7/6/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Sprycel | |
|-----------------------|--|
| Diagnosis | Philadelphia chromosome-positive or BCR-ABL1-positive chronic myeloid leukemia |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of Philadelphia chromosome-positive or BCR-ABL1-positive chronic myeloid leukemia

2 - One of the following:

2.1 Patient is not a candidate for imatinib as attested by physician

OR

2.2 Patient is currently on Sprycel therapy

| Product Name: Sprycel | |
|-----------------------|--|
| Diagnosis | Philadelphia chromosome-positive or BCR-ABL1-positive chronic myeloid leukemia |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Patient does not show evidence of progressive disease while on Sprycel therapy

| Product Name: Sprycel | |
|-----------------------|--|
| Diagnosis | Philadelphia chromosome-positive acute lymphoblastic leukemia (Ph+ALL) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of Philadelphia chromosome-positive acute lymphoblastic leukemia (Ph+ALL)

| Product | Name: | Sprvcel |
|---------|-------|---------|
| | | 001,000 |

| Diagnosis | Philadelphia chromosome-positive acute lymphoblastic leukemia (Ph+ALL) |
|-----------------|--|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

1 - Patient does not show evidence of progressive disease while on Sprycel therapy

| Product Name: Sprycel | |
|-----------------------|---------------------------------------|
| Diagnosis | Gastrointestinal stromal tumor (GIST) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of gastrointestinal stromal tumor (GIST) with PDGFRA D842V mutation

| Product Name: Sprycel | |
|---------------------------------------|--|
| Gastrointestinal stromal tumor (GIST) | |
| 12 month(s) | |
| Reauthorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Sprycel therapy

| Product Name: Sprycel | |
|-----------------------|----------------|
| Diagnosis | Chondrosarcoma |
| Approval Length | 12 month(s) |

| Therapy Stage | Initial Authorization |
|----------------|-----------------------|
| Guideline Type | Prior Authorization |
| | |

1 - Diagnosis of metastatic chondrosarcoma

| Product Name: Sprycel | |
|-----------------------|--|
| Chondrosarcoma | |
| 12 month(s) | |
| Reauthorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Sprycel therapy

| Product Name: Sprycel | |
|-----------------------|-----------------------|
| Diagnosis | Chordoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| | |

Approval Criteria

1 - Diagnosis of recurrent chordoma

| Product Name: Sprycel | |
|-----------------------|---------------------|
| Diagnosis | Chordoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Patient does not show evidence of progressive disease while on Sprycel therapy

| Product Name: Sprycel | |
|-----------------------|---|
| Diagnosis | Lymphoid, myeloid, or mixed lineage neoplasms with eosinophilia |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of lymphoid, myeloid, or mixed lineage neoplasms with eosinophilia

AND

2 - Patient has an ABL1 (gene) rearrangement

| Product Name: Sprycel | |
|-----------------------|---|
| Diagnosis | Lymphoid, myeloid, or mixed lineage neoplasms with eosinophilia |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Sprycel therapy

| Product Name: Sprycel | |
|-----------------------|--|
| Diagnosis | National Comprehensive Cancer Network (NCCN) Recommended Regimen |
| Approval Length | 12 month(s) |

| Therapy Stage | Initial Authorization |
|----------------|-----------------------|
| Guideline Type | Prior Authorization |

1 - Uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Sprycel | |
|-----------------------|--|
| Diagnosis | National Comprehensive Cancer Network (NCCN) Recommended Regimen |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | • |

Approval Criteria

1 - Documentation of positive clinical response to Sprycel therapy

2. Revision History

| Date | Notes |
|----------|--------------|
| 7/6/2022 | Updated GPIs |

Stelara (ustekinumab)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-114494 | Stelara | (ustekinumab) |
|-----------|---------|---------------|
|-----------|---------|---------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Stelara SC | |
|---------------------------|-----------------------|
| Diagnosis | Plaque Psoriasis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - ONE of the following: | |

1.1 Submission of medical records (e.g., chart notes, laboratory values, prescription claims history) documenting ALL of the following:

1.1.1 Diagnosis of chronic moderate to severe plaque psoriasis

AND

1.1.2 Greater than or equal to 3 percent body surface area involvement, palmoplantar, facial, or genital involvement, or severe scalp psoriasis

AND

1.1.3 History of failure to ONE of the following topical therapies, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*:

- Corticosteroids (e.g., betamethasone, clobetasol, desonide)
- Vitamin D analogs (e.g., calcitriol, calcipotriene)
- Tazarotene
- Calcineurin inhibitors (e.g., tacrolimus, pimecrolimus)
- Anthralin
- Coal tar

AND

1.1.4 History of failure to a 3 month trial of methotrexate at the maximally indicated dose within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*

AND

1.1.5 History of failure, contraindication, or intolerance to ALL of the following preferred biologic products (document drug, date, and duration of trial)*:

- Humira (adalimumab)
- Enbrel (etanercept)
- Otezla (apremilast)

1.1.6 Patient is NOT receiving Stelara in combination with ANY of the following: Biologic disease modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), • Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)] Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)] • Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)] • AND **1.1.7** ONE of the following: **1.1.7.1** Requested medication is Stelara 45 mg (milligrams) per 0.5 mL (milliliter) OR **1.1.7.2** BOTH of the following: Requested medication is Stelara 90 mg per 1 mL Patient's weight is greater than 100 kg (kilograms) (220 pounds) AND **1.1.8** Prescribed by or in consultation with a dermatologist OR **1.2** All of the following: **1.2.1** Patient is currently on Stelara therapy as documented by claims history or medical records (document drug, date, and duration of therapy) AND 1.2.2 Diagnosis of chronic moderate to severe plaque psoriasis

| AND | |
|---|--|
| | |
| 1.2.3 Patient is NOT | receiving Stelara in combination with ANY of the following: |
| (certolizumab), • Janus kinase in | D [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia Simponi (golimumab)] hibitor [e.g., Xeljanz (tofacitinib)] rase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)] |
| | AND |
| 1.2.4 Prescribed by or in consultation with a dermatologist | |
| | AND |
| 2 - Patient is 6 years of age or older | |
| Notes | *Claims history may be used in conjunction as documentation of drug, date, and duration of trial |

| Product Name: Stelara SC | |
|--------------------------|---------------------|
| Diagnosis | Plaque Psoriasis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Documentation of positive clinical response to Stelara therapy

AND

2 - Patient is NOT receiving Stelara in combination with ANY of the following:

- Biologic disease modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

3 - Prescribed by or in consultation with a dermatologist

| Product Name: Stelara SC | |
|--------------------------|-----------------------|
| Diagnosis | Psoriatic Arthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- **1** ONE of the following:
- **1.1** ALL of the following:
- **1.1.1** ONE of the following
- **1.1.1.1** BOTH of the following:
- Requested medication is Stelara 45 mg (milligrams) per 0.5 mL (milliliter)
- Diagnosis of active psoriatic arthritis

OR

1.1.1.2 ALL of the following:

- Diagnosis of active psoriatic arthritis
- Diagnosis of co-existent moderate to severe plaque psoriasis

AND

1.1.2 Patient is NOT receiving Stelara in combination with ANY of the following: Biologic disease modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), • Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)] Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)] Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)] • AND 1.1.3 History of failure to a 3 month trial of methotrexate at the maximally indicated dose within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)* AND **1.1.4** History of failure, contraindication, or intolerance to three of the following: Humira (adalimumab) Enbrel (etanercept) ٠ Otezla (apremilast) • Xeljanz (tofacitinib) AND **1.1.5** Prescribed by or in consultation with one of the following: Rheumatologist . Dermatologist OR **1.2** All of the following: **1.2.1** Patient is currently on Stelara therapy as documented by claims history or medical records (document drug, date, and duration of therapy) AND

| 1.2.2 Diagnosis of active psoriatic arthritis | |
|--|--|
| | AND |
| 1.2.3 Patient is NOT | receiving Stelara in combination with ANY of the following: |
| (certolizumab), Janus kinase in | D [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia Simponi (golimumab)] hibitor [e.g., Xeljanz (tofacitinib)] rase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)] |
| | AND |
| 1.2.4 Prescribed by c | or in consultation with one of the following: |
| RheumatologistDermatologist | t |
| | AND |
| 2 - Patient is 6 years of | f age or older |
| Notes | *Claims history may be used in conjunction as documentation of drug, date, and duration of trial |

| Product Name: Stelara SC | |
|--------------------------|---------------------|
| Diagnosis | Psoriatic Arthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Documentation of positive clinical response to Stelara therapy

2 - Patient is NOT receiving Stelara in combination with ANY of the following:

- Biologic disease modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with one of the following:

- Rheumatologist
- Dermatologist

| Product Name: Stelara SC, Stelara IV | |
|--------------------------------------|-----------------------|
| Diagnosis | Crohn's Disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of moderately to severely active Crohn's disease

AND

2 - One of the following:

2.1 Both of the following

2.1.1 History of failure to one of the following conventional therapies at maximally indicated doses within the last 3 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*:

| 6-mercaptopurir | |
|---|--|
| Azathioprine (InMethotrexate (F | nuran) Rheumatrex, Trexall) |
| | AND |
| 2.1.2 History of failure | e, contraindication or intolerance to Humira (adalimumab) |
| | OR |
| | y on Stelara therapy as documented by claims history or medical g, date, and duration of therapy) |
| | AND |
| 3 - Patient is NOT rece | iving Stelara in combination with ANY of the following: |
| Humira (adalim | e modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), umab), Cimzia (certolizumab), Simponi (golimumab)] hibitor [e.g., Xeljanz (tofacitinib)] |
| Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)] | |
| AND | |
| 4 - Prescribed by or in a | consultation with a gastroenterologist |
| Notes | *Claims history may be used in conjunction as documentation of drug, date, and duration of trial |
| | date, and duration of trial |

| Product Name: Stelara SC, Stelara IV | |
|--------------------------------------|-----------------------|
| Diagnosis | Ulcerative Colitis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria 1 - Diagnosis of moderately to severely active ulcerative colitis AND 2 - One of the following: 2.1 Both of the following **2.1.1** History of failure to one of the following conventional therapies at maximally indicated doses within the last 3 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*: Corticosteroids (e.g., prednisone, methylprednisolone, budesonide) • 6-mercaptopurine (Purinethol) ٠ Azathioprine (Imuran) • Aminosalicylates (e.g., mesalamine, sulfasalazine) AND **2.1.2** History of failure, contraindication or intolerance to Humira (adalimumab) OR **2.2** Patient is currently on Stelara therapy as documented by claims history or medical records (document drug, date, and duration of therapy) AND **3** - Patient is NOT receiving Stelara in combination with ANY of the following: Biologic disease modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), • Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)] Janus Kinase Inhibitor [e.g., Xeljanz (tofacitinib)] • Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)] •

| | AND |
|--|--|
| 4 - Prescribed by or in consultation with a gastroenterologist | |
| Notes | *Claims history may be used in conjunction as documentation of drug, date, and duration of trial |

| Product Name: Stelara SC, Stelara IV | |
|--------------------------------------|-------------------------------------|
| Diagnosis | Crohn's Disease, Ulcerative Colitis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Documentation of positive clinical response to Stelara therapy

AND

2 - Patient is NOT receiving Stelara in combination with ANY of the following:

- Biologic disease modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab)]
- Janus Kinase Inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with a gastroenterologist

2. Revision History

| Date | Notes |
|-----------|--|
| 9/26/2022 | Added age criterion for PsA and PsO. Updated product list. |

Strensiq

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99646 Strensiq

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Strensiq | |
|---------------------------|--|
| Diagnosis | perinatal/infantile or juvenile-onset hypophosphatasia (HPP) |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - All of the following: | |

1.1 Diagnosis of perinatal/infantile or juvenile-onset hypophosphatasia based on all of the following:

1.1.1 One of the following:

- Onset of clinical signs and symptoms of hypophosphatasia prior to age 18 years (e.g., respiratory insufficiency, vitamin B6 responsive seizures, hypotonia, failure to thrive, delayed walking, waddling gait, dental abnormalities, low trauma fractures)
- Radiographic evidence supporting the diagnosis of hypophosphatasia at the age of onset prior to age 18 years (e.g., craniosynostosis, infantile rickets, non-traumatic fractures)

AND

1.1.2 One of the following:

1.1.2.1 Both of the following:

- Patient has low level activity of serum alkaline phosphatase (ALP) evidenced by an ALP level below the age-adjusted normal range
- Patient has an elevated level of tissue non-specific alkaline phosphatase (TNSALP) substrate (e.g. serum pyridoxal 5'-phosphate [PLP] level, serum or urine phosphoethanolamine [PEA] level, urinary inorganic pyrophosphate [PPi level])

OR

1.1.2.2 Confirmation of tissue-nonspecific alkaline phosphatase (TNSALP) gene mutation by ALPL genomic DNA testing*

AND

1.2 Prescribed by one of the following:

- Endocrinologist
- A specialist experienced in the treatment of metabolic bone disorders

AND

1.3 One of the following:

1.3.1 Both of the following:

- Diagnosis of perinatal/infantile-onset hypophosphatasia
- Coverage will be provided up to a maximum supply limit of 9 mg/kg/week

OR

1.3.2 Both of the following:

- Diagnosis of juvenile-onset hypophosphatasia
- Coverage will be provided up to a maximum supply limit of 6 mg/kg/week

AND

1.4 One of the following:

1.4.1 Patient is prescribed Strensiq 18 mg/0.45 mL, Strensiq 28 mg/0.7 mL, or Strensiq 40 mg/mL vials

OR

1.4.2 Both of the following:

- Patient is prescribed Strensiq 80 mg/0.8 mL vial
- Patient's weight is greater than or equal to 40 kg

AND

1.5 Prescriber attests to the following: the information provided is true and accurate to the best of their knowledge and they understand that UnitedHealthcare may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided

| *Results of prior genetic testing can be submitted as confirmation of di agnosis of HPP, however please note that the provider should confirm |
|---|
| coverage status of any new genetic testing under the patient's United Healthcare plan prior to ordering |

Product Name: Strensig

| Diagnosis | perinatal/infantile or juvenile-onset hypophosphatasia (HPP) |
|-----------------|--|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - All of the following:

1.1 Clinically relevant decrease from baseline in tissue non-specific alkaline phosphatase (TNSALP) substrate (e.g. serum pyridoxal 5'-phosphate [PLP] level, serum or urine phosphoethanolamine [PEA] level, urinary inorganic pyrophosphate [PPi level])

AND

1.2 Prescribed by one of the following:

- Endocrinologist
- A specialist experienced in the treatment of metabolic bone diseases

AND

- **1.3** One of the following:
- 1.3.1 Both of the following:
 - Diagnosis of perinatal/infantile-onset hypophosphatasia
 - Coverage will be provided up to a maximum supply limit of 9 mg/kg/week

OR

1.3.2 Both of the following:

- Diagnosis of juvenile-onset hypophosphatasia
- Coverage will be provided up to a maximum supply limit of 6 mg/kg/week

AND

1.4 One of the following:

1.4.1 Patient is prescribed Strensiq 18 mg/0.45 mL, Strensiq 28 mg/0.7 mL, or Strensiq 40 mg/mL vials

OR

1.4.2 Both of the following

- Patient is prescribed Strensiq 80 mg/0.8 mL vials
- Patient's weight is greater than or equal to 40 kg

AND

1.5 Prescriber attests to the following: the information provided is true and accurate to the best of their knowledge and they understand that UnitedHealthcare may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided

2. Revision History

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Sublingual Immunotherapy (SLIT)

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-105262 Sublingual Immunotherapy (SLIT)

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 4/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: All products | | |
|---|--|--|
| Diagnosis | Patients 21 years of age and older | |
| Approval Length | N/A - All requests for patients 21 years of age and older should be DENIED as benefit exclusion | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Requests for patients 21 years of age and older are not covered | | |
| Notes | Approval Length: N/A - All requests for patients 21 years of age and ol der should be denied as a benefit exclusion. | |

| Product Name: Grastek | |
|--|--|
| Diagnosis | Grass pollen-induced allergic rhinitis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| Approval Criteria 1 - Diagnosis of moderate to severe grass pollen-induced allergic rhinitis AND | |
| 2 - Diagnosis confirmed by one of the following: Positive skin test to Timothy grass or cross-reactive grass pollens (eg, Sweet Vernal, Orchard/Cocksfoot, Perennial Rye, Kentucky blue/June grass, Meadow Fescue, or Redtop) in vitro testing for pollen-specific IgE antibodies for Timothy grass or cross-reactive grass pollens (e.g., Sweet Vernal, Orchard/Cocksfoot, Perennial Rye, Kentucky blue/June grass, Meadow Fescue, or Redtop) | |

3 - Treatment is started or will be started at least 12 weeks before the beginning of the grass pollen season

AND

4 - History of failure, contraindication, or intolerance to two of the following:

- ٠
- oral antihistamine [e.g. cetirizine (Zyrtec)] intranasal antihistamine [e.g. azelastine (Astelin)] •
- intranasal corticosteroid [e.g. fluticasone (Flonase)] •
- leukotriene inhibitor [e.g. montelukast (Singulair)] •

5 - Not received in combination with similar cross-reactive grass pollen immunotherapy (e.g., Oralair)

AND

6 - Patient does not have unstable and/or uncontrolled asthma

AND

7 - Prescribed by or in consultation with a specialist in allergy and immunology

| Product Name: Grastek | |
|-----------------------|--|
| Diagnosis | Grass pollen-induced allergic rhinitis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Documentation of positive clinical response to Grastek therapy

| Product Name: Oralair | |
|--|--|
| Grass pollen-induced allergic rhinitis | |
| 12 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Diagnosis of moderate to severe grass pollen-induced allergic rhinitis

| AND |
|--|
| 2 - Diagnosis confirmed by one of the following: |
| Positive skin test to any of the five grass species contained in Oralair [(i.e., Sweet Vernal, Orchard, Perennial Rye, Timothy, and Kentucky Blue grass mixed pollens) or cross-reactive grass pollens (e.g., Cocksfoot, Meadow Fescue, or Redtop)] in vitro testing for pollen-specific IgE antibodies for any of the five grass species contained in Oralair [(i.e., Sweet Vernal, Orchard, Perennial Rye, Timothy, and Kentucky Blue grass mixed pollens) or cross-reactive grass pollens (e.g., Cocksfoot, Meadow Fescue, or Redtop)] |
| AND |
| 3 - Treatment is started or will be started at least 4 months before the beginning of the grass pollen season |
| AND |
| 4 - History of failure, contraindication, or intolerance to two of the following: |
| oral antihistamine [e.g. cetirizine (Zyrtec)] intranasal antihistamine [e.g. azelastine (Astelin)] intranasal corticosteroid [e.g. fluticasone (Flonase)] leukotriene inhibitor [e.g. montelukast (Singulair)] |
| AND |
| 5 - Not received in combination with similar cross-reactive grass pollen immunotherapy (e.g., Grastek) |
| AND |
| 6 - Patient does not have unstable and/or uncontrolled asthma |
| AND |

7 - Prescribed by or in consultation with a specialist in allergy and immunology

| Product Name: Oralair | |
|-----------------------|--|
| Diagnosis | Grass pollen-induced allergic rhinitis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Oralair therapy

| Product Name: Ragwitek | |
|------------------------|--|
| Diagnosis | Short ragweed pollen-induced allergic rhinitis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of moderate to severe short ragweed pollen-induced allergic rhinitis

AND

2 - Diagnosis confirmed by one of the following:

- Positive skin test to short ragweed pollen
- in vitro testing for pollen-specific IgE antibodies for short ragweed pollen

AND

3 - Treatment is started or will be started at least 12 weeks before the beginning of the short ragweed pollen season

4 - History of failure, contraindication, or intolerance to two of the following:

- oral antihistamine [e.g. cetirizine (Zyrtec)]
- intranasal antihistamine [e.g. azelastine (Astelin)]
- intranasal corticosteroid [e.g. fluticasone (Flonase)]
- leukotriene inhibitor [e.g. montelukast (Singulair)]

AND

5 - Patient does not have unstable and/or uncontrolled asthma

AND

6 - Prescribed by or in consultation with a specialist in allergy and immunology

| Product Name: Ragwitek | |
|--|--|
| Short ragweed pollen-induced allergic rhinitis | |
| 12 month(s) | |
| Reauthorization | |
| Prior Authorization | |
| | |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Ragwitek therapy

| Product Name: Odactra | |
|-----------------------|---|
| Diagnosis | House dust mite (HDM)-induced allergic rhinitis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

| Approval Criteria |
|---|
| 1 - Diagnosis of house dust mite (HDM)-induced allergic rhinitis. |
| |
| AND |
| |
| 2 - Diagnosis confirmed by one of the following: |
| Positive skin test to licensed house dust mite allergen extracts in vitro testing for IgE antibodies to Dermatophagoides farinae or Dermatophagoides pteronyssinus house dust mites |
| |
| AND |
| |
| 3 - History of failure, contraindication, or intolerance to two of the following: |
| oral antihistamine [e.g. cetirizine (Zyrtec)] intranasal antihistamine [e.g. azelastine (Astelin)] intranasal corticosteroid [e.g. fluticasone (Flonase)] leukotriene inhibitor [e.g. montelukast (Singulair)] |
| AND |
| 4 - Patient does not have unstable and/or uncontrolled asthma |
| AND |

Γ

5 - Prescribed by or in consultation with a specialist in allergy and immunology

| Product Name: Odactra | |
|-----------------------|---|
| Diagnosis | House dust mite (HDM)-induced allergic rhinitis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Documentation of positive clinical response to Odactra therapy

| Date | Notes |
|-----------|---|
| 3/28/2022 | Added box to deny as benefit exclusion for patients 21 years and old er |

Sublocade - Arizona

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99647 | Sublocade - Arizona |
|----------|---------------------|
|----------|---------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Sublocade | |
|-------------------------|--|
| 6 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Patient has severe Opioid Use Disorder (OUD) as defined by the DSM-5 (Diagnostic and Statistical Manual of Mental Disorders, Fifth Edition) OUD Diagnostic Tool and has a demonstrated history of non-adherence to oral medications

2 - Patient is currently maintained on 8mg to 24mg per day dose of oral, sublingual, or transmucosal buprenorphine product equivalent for at least 7 days prior to initiation of extended-release buprenorphine injection

AND

3 - Patient has not, nor will receive supplemental, oral, sublingual, or transmucosal buprenorphine

AND

4 - Patient is receiving psychosocial interventions as part of a comprehensive medication assisted treatment (MAT) program

AND

5 - Prescriber meets DATA 2000 (Drug Addiction Treatment Act of 2000) requirements and has been assigned a unique identification number specific to the prescription of medication assisted therapy (DEA-X)

AND

6 - Prescriber checks the Arizona State Board of Pharmacy Controlled Substance Prescription Monitoring Program (CSPMP) database prior to each monthly injection

AND

7 - Sublocade dosing is in accordance with the U. S. Food and Drug Administration approved labeling: 300mg (milligrams) subcutaneously monthly for the first 2 months, followed by a maintenance dose of 100mg or 300mg monthly

Product Name: Sublocade

| Approval Length | 12 month(s) |
|---|---|
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| Approval Criteria 1 - Physician documentation that the patient has experienced a positive clinical response to buprenorphine extended-release therapy, as defined by the provider | |
| | AND |
| 2 - Patient has not, n buprenorphine | or will receive supplemental, oral, sublingual, or transmucosal |
| | AND |
| 3 - Patient is receiving psychosocial interventions as part of a comprehensive medication assisted treatment (MAT) program | |
| | AND |
| | DATA 2000 (Drug Addiction Treatment Act of 2000) requirements and unique identification number specific to the prescription of medication EA-X) |
| AND | |
| 5 - Prescriber checks the Arizona State Board of Pharmacy Controlled Substance Prescription Monitoring Program (CSPMP) database prior to each monthly injection | |
| | AND |
| | g is in accordance with the U. S. Food and Drug Administration approved e dose of 100mg (milligrams) or 300mg monthly |

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Suboxone - AZM

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-114461 Suboxone - AZM

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 9/26/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Generic buprenorphine-naloxone film | | |
|---|--|--|
| Approval Length | N/A - Requests for generic buprenorphine hcl-naloxone film should not be approved | |
| Guideline Type | Prior Authorization | |
| Approval Criteria | | |
| | c buprenorphine-naloxone film are not authorized and will not be | |
| Notes | Approval Length: N/A - Requests for generic buprenorphine-naloxone film should not be approved. Patient need to use Brand Suboxone film or other preferred alternatives. | |

| Product Name: Zubsolv, Bunavail ** | |
|------------------------------------|-----------------------|
| Approval Length | 3 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

1 - The patient has a Diagnostic and Statistical Manual, Fifth Edition, Text Revision, (DSM-V-TR) diagnosis of opioid use disorder

AND

2 - The patient must have a reason or special circumstance that they cannot use the preferred products **

- brand Suboxone Film
- buprenorphine (generic Subutex)
- buprenorphine HCI/naloxone Tab (Generic Suboxone Tab)
- naloxone
- naltrexone
- Narcan (naloxone)
- Sublocade (buprenorphine)
- Vivitrol (naltrexone microspheres)

| *Up to 24 mg per day of Suboxone, or equivalent dosing of an alternat ive medication, will be authorized for the initial period. **Please refere nce current PDL: https://www.azahcccs.gov/Resources/Downloads/Ph armacyUpdates/AHCCCS_DRUG_LIST4.1.2022-updated.pdf |
|---|

| Product Name: Zubsolv, Bunavail ** | |
|------------------------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - The patient has been prescribed a buprenorphine product for the purpose of opioid use disorder maintenance therapy

AND

2 - The patient must have a reason or special circumstance that they cannot use the preferred products**

AND

3 - Patient must have tried Suboxone film or buprenorphine-naloxone ODT tablets

| Notes | * Up to 16 mg per day of Suboxone, or equivalent dosing of an alterna tive medication, will be authorized for the reauthorization period **Plea se reference current PDL: https://www.azahcccs.gov/Resources/Dow nloads/PharmacyUpdates/AHCCCS_DRUG_LIST4.1.2022-updated.p df |
|-------|---|
|-------|---|

| Product Name: Brand suboxone, generic buprenorphine hcl- | naloxone, |
|---|-----------|
| buprenorphine/naloxone sublingual tablet, Zubsolv, Bunavail | * |

| Approval Length | 3 month(s) |
|-----------------|-----------------------|
| Therapy Stage | Initial Authorization |
| Guideline Type | Quantity Limit |

Approval Criteria

1 - Physician has provided rationale for needing to exceed the buprenorphine daily limit

AND

2 - The requested dosage cannot be achieved using the plan accepted quantity limit of a different dose or formulation

| Notes | * This criteria applies to requests exceeding 24 mg of buprenorphine o |
|-------|--|
| | r equivalent |

| Product Name: Brand suboxone, generic buprenorphine hcl-naloxone, buprenorphine/naloxone sublingual tablet, Zubsolv, Bunavail * | |
|--|--|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Quantity Limit |
| | |
| Approval Criteria | |
| 1 - Physician has provided rationale for needing to exceed the buprenorphine daily limit | |
| AND | |
| 2 - The requested dosage cannot be achieved using the plan accepted quantity limit of a different dose or formulation | |
| Notes | *This criteria applies to requests exceeding 16 mg of buprenorphine o r equivalent |

| Date | Notes |
|-----------|--|
| 9/26/2022 | Added criteria to deny PA requests for generic suboxone film |

Sucraid

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99648 Sucraid

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Sucraid | |
|-----------------------|-----------------------|
| Approval Length | 3 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | • |

Approval Criteria

1 - Diagnosis of congenital sucrase-isomaltase deficiency (CSID) as confirmed by one of the following:

1.1 Duodenal biopsy showing low sucrose activity and normal amounts of other disaccharides

| OR |
|--|
| 1.2 All of the following: |
| Stool pH less than 6 Negative lactose breath test Increase in breath hydrogen greater than 10 ppm (parts per million) when challenged with sucrose after fasting |
| AND |
| 2 - Prescribed by or in consultation with a gastroenterologist or rare disease specialist |
| AND |
| 3 - Will be used with a sucrose-free, low starch diet |

| Product Name: Sucraid | |
|-----------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Prescribed by or in consultation with a gastroenterologist or rare disease specialist

AND

2 - Will be used with a sucrose-free, low starch diet

AND

3 - Provider attests that the patient has achieved a clinically meaningful response while on Sucraid therapy, defined as at least a 50 percent reduction in all of the following:

- Symptoms of abdominal pain, cramps, bloating, gas, vomiting •
- •
- Number of stools per day Watery, loose stool consistency •
- Number of symptomatic days •

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Sunosi

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99524 Sunosi

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Sunosi | |
|----------------------|-----------------------|
| Diagnosis | Narcolepsy |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g. chart notes, lab values) documenting a diagnosis of narcolepsy with BOTH of the following:

1.1 The patient has daily periods of irrepressible need to sleep or daytime lapses into sleep occurring for at least three months.

OR

1.2 A mean sleep latency of less than or equal to 8 minutes and two or more sleep onset rapid eye movement (REM) periods (SOREMPs) are found on a multiple sleep latency test (MSLT) performed according to standard techniques following a normal overnight polysomnogram. A SOREMP (within 15 minutes of sleep onset) on the preceding nocturnal polysomnogram may replace one of the SOREMPs on the MSLT.

AND

- 2 Physician attestation to the following:
 - Other causes of sleepiness have been ruled out or treated (including but not limited to obstructive sleep apnea, insufficient sleep syndrome, shift work, the effects of substances or medications or their withdrawal, sleep phase disorder, or other sleep disorders)

AND

- **3** History of failure, contraindication, or intolerance to BOTH of the following:
- **3.1** ONE of the following:
 - Amphetamine based stimulant (e.g., amphetamine, dextroamphetamine)
 - Methylphenidate based stimulant

AND

3.2 Armodafinil

AND

- **4** Prescribed by one of the following:
 - Neurologist

- Psychiatrist
- Sleep Medicine Specialist

| Product Name: Sunosi | |
|----------------------|---------------------|
| Diagnosis | Narcolepsy |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Reduction in symptoms of excessive daytime sleepiness associated with Sunosi therapy

| Product Name: Sunosi | |
|----------------------|-------------------------|
| Diagnosis | Obstructive Sleep Apnea |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g. chart notes, lab values) documenting a diagnosis of obstructive sleep apnea with ONE of the following:

1.1 Fifteen or more obstructive respiratory events per hour of sleep confirmed by a sleep study

OR

1.2 BOTH of the following:

1.2.1 Five or more obstructive respiratory events per hour of sleep confirmed by a sleep study

1.2.2 ONE or more of the following sign/symptoms are present:

- Daytime sleepiness
- Nonrestorative sleep
- Fatigue
- Insomnia
- Waking up with breath holding, gasping, or choking
- Habitual snoring noted by bed partner or other observer
- Observed apnea

AND

2 - BOTH of the following:

2.1 Standard treatments for the underlying airway obstruction (e.g., continuous positive airway pressure [CPAP], bi-level positive airway pressure [BiPAP]) have been used for one month or longer

AND

2.2 Patient is fully compliant with ongoing treatment(s) for the underlying airway obstruction

AND

3 - History of failure, contraindication, or intolerance to armodafinil

AND

- **4** Prescribed by one of the following:
 - Neurologist
 - Psychiatrist
 - Sleep Medicine Specialist

| Product Name: Sunosi | |
|----------------------|-------------------------|
| Diagnosis | Obstructive Sleep Apnea |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Reduction in symptoms of excessive daytime sleepiness associated with Sunosi therapy

AND

2 - Patient continues to be fully compliant with ongoing treatment(s) for the underlying airway obstruction (e.g. continuous positive airway pressure [CPAP], bi-level positive airway pressure [BiPAP])

| Date | Notes |
|-----------|--------------------|
| 5/27/2021 | 7/1 Implementation |

Sutent

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99767 Sutent

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Sutent | |
|--|---------------------------------------|
| Diagnosis | Gastrointestinal Stromal Tumor (GIST) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of gastrointestinal stromal tumor (GIST) | |

2 - History of failure, contraindication, or intolerance to Gleevec (imatinib)

| Product Name: Sutent | | |
|---|----------------------------|--|
| Diagnosis | Renal Cell Carcinoma (RCC) | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of renal c | ell carcinoma (RCC) | |
| | AND | |
| 2 - ONE of the following: | | |
| 2.1 Disease has relap | sed | |
| | | |
| | OR | |
| 2.2 Diagnosis of Stage IV disease | | |
| OR | | |
| 2.3 BOTH of the following: | | |
| 2.3.1 Used in adjuvant setting | | |
| AND | | |
| 2.3.2 Patient has a high risk of recurrence following nephrectomy | | |

| Product Name: Sutent | |
|----------------------|--|
| Diagnosis | Islet Cell Tumor / Progressive Pancreatic Neuroendocrine Tumors (pNET) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

1 - Diagnosis of islet cell tumor / progressive pancreatic neuroendocrine tumors (pNET)

AND

- **2** Disease is ONE of the following:
 - Unresectable, locally advanced
 - Metastatic

| Product Name: Sutent | |
|----------------------|-----------------------|
| Diagnosis | Soft Tissue Sarcoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- **1** Diagnosis of ONE of the following:
 - Alveolar soft part sarcoma (ASPS)
 - Angiosarcoma
 - Solitary fibrous tumor / hemangiopericytoma

Product Name: Sutent

| Diagnosis | Thyroid Carcinoma |
|-----------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

1 - ONE of the following:

- **1.1** ALL of the following:
- **1.1.1** Diagnosis of ONE of the following:
 - Follicular carcinoma
 - Hürthle cell carcinoma
 - Papillary carcinoma

AND

1.1.2 ONE of the following:

- Unresectable locoregional recurrent disease
- Persistent disease
- Metastatic disease

AND

1.1.3 ONE of the following:

- Patient has symptomatic disease
- Patient has progressive disease

AND

1.1.4 Disease is refractory to radioactive iodine treatment

1.2 ALL of the following:

1.2.1 Diagnosis of medullary thyroid carcinoma

AND

1.2.2 ONE of the following:

- •
- Patient has progressive disease Patient has symptomatic metastatic disease •

AND

1.2.3 History of failure, contraindication, or intolerance to ONE of the following:

- Caprelsa (vandetanib) •
- Cometriq (cabozantinib) •

| Product Name: Sutent | |
|----------------------|-----------------------|
| Diagnosis | Chordoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

Approval Criteria

1 - Diagnosis of recurrent chordoma

| Product Name: Sutent | |
|----------------------|-------------------------------|
| Diagnosis | Central Nervous System Cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

1 - Diagnosis of surgically inaccessible meningiomas

AND

2 - ONE of the following:

- Disease is recurrent
- Disease is progressive

AND

3 - Further radiation is not possible

| Product Name: Sutent | | |
|-----------------------------------|-----------------------|--|
| Diagnosis | Thymic Carcinoma | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of thymic carcinoma | | |
| | | |
| AND | | |
| | | |

2 - Used as second-line following a failure, contraindication, or intolerance to a first-line chemotherapy regimen (e.g., carboplatin/paclitaxel)

| Product Name: Sutent | |
|----------------------|---|
| 0 | Gastrointestinal Stromal Tumor (GIST), Renal Cell Carcinoma (RCC), Islet Cell Tumor / Progressive Pancreatic Neuroendocrine Tumors |

| | (pNET), Soft Tissue Sarcoma, Thyroid Carcinoma, Chordoma, Central Nervous System Cancer, Thymic Carcinoma | |
|-----------------|---|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Guideline Type Prior Authorization | |
| | | |

1 - Patient does not show evidence of progressive disease while on Sutent therapy

| Product Name: Sutent | | |
|----------------------|---------------------------|--|
| Diagnosis | NCCN Recommended Regimens | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |

Approval Criteria

1 - Sutent will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Sutent | | |
|----------------------|-----------------------------------|--|
| Diagnosis | NCCN Recommended Regimens | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | uideline Type Prior Authorization | |
| | | |

Approval Criteria

1 - Documentation of positive clinical response to Sutent therapy

| Date | Notes |
|----------|-------------------------------------|
| 6/2/2021 | Arizona Medicaid 7.1 Implementation |

Symdeko

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99649 Symdeko

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Symdeko | | |
|---------------------------------------|-----------------------|--|
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of cystic fibrosis (CF) | | |

2 - Submission of laboratory result documenting ONE of the following:

2.1 The patient is homozygous for the F508del mutation in the cystic fibrosis transmembrane conductance regulator (CFTR) gene

OR

2.2 The patient has at least ONE mutation in the CFTR gene that is responsive to Symdeko (See Table in Background Section)

AND

3 - The patient is greater than or equal to 6 years of age

AND

4 - Prescribed by or in consultation with a specialist affiliated with a CF care center

| Product Name: Symdeko | |
|-----------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Provider attests that the patient has achieved a clinically meaningful response while on Symdeko therapy to ONE of the following:

- Lung function as demonstrated by percent predicted expiratory volume in 1 second (ppFEV1)
- Body mass index (BMI)
- Pulmonary exacerbations

• Quality of life as demonstrated by Cystic Fibrosis Questionnaire-Revised (CFQ-R) respiratory domain score

AND

2 - Prescribed by, or in consultation with, a specialist affiliated with a cystic fibrosis (CF) care center

2. Background

| Benefit/Coverage/Program Information |
|--------------------------------------|
| Table 1 CFTR Gene Mutations |

| A1067T | D1270N | F1052V | R1070W | S945L | 3272-26A→G |
|--------|--------|--------|--------|-------|--------------|
| A455E | D579G | F1074L | R117C | S977F | 3849+10kbC→T |
| D110E | E193K | K1060T | R347H | | 711+3A→G |
| D110H | E56K | L206W | R352Q | | 2789+5G→A |
| D1152H | E831X | P67L | R74W | | |

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Symlin

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99499 Symlin

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Symlin | | | |
|---|---------------------|--|--|
| Approval Length | 12 month(s) | | |
| Guideline Type | Prior Authorization | | |
| | | | |
| Approval Criteria | | | |
| 1 - Patient must have ONE of the following diagnoses: | | | |
| Type 1 diabetesType 2 diabetes | | | |

| | AND |
|--|-----|
| 2 - Concurrent use of insulin therapy | |

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Standard to Medicaid Arizona Standard for 7 /1 go live |

Synagis

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99650 Synagis

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Synagis* | | | | |
|---|---------------------------------|--|--|--|
| Diagnosis | osis Prematurity | | | |
| Guideline Type | deline Type Prior Authorization | | | |
| | | | | |
| Approval Criteria | | | | |
| 1 - BOTH of the following: | | | | |
| 1.1 Patient is an infant born before 29 weeks, 0 days gestation | | | | |

AND

1.2 Patient is less than 12 months of age at the start of RSV "season"

AND

2 - Administered during RSV season**

AND

3 - Monthly dose of Synagis does not exceed 15 milligram per kilogram per dose

AND

4 - Monthly dose of Synagis does not exceed 5 doses per single RSV "season"***

AND

5 - The patient does not meet ONE of the following situations

- Infants and children with hemodynamically insignificant heart disease (e.g., secundum atrial septal defect, small ventricular septal defect, pulmonic stenosis, uncomplicated aortic stenosis, mild coarctation of the aorta, and patent ductus arteriosus)
- Infants with congenital heart disease and cardiac lesions adequately corrected by surgery, unless they continue to require medication for congestive heart failure
- Infants with cardiomyopathy sufficiently mild that they do not require pharmacotherapy
- Routine use of prophylaxis in children with Down syndrome [unless qualifying heart disease, CLD, airway clearance issues (the inability to clear secretions from the upper airway because of ineffective cough), or prematurity (less than 29 weeks, 0 days gestation) is present]
- Routine use of prophylaxis in children with cystic fibrosis (unless indications noted in proven indications above are present)
- Administration of monthly Synagis prophylaxis after an infant or child has experienced a breakthrough RSV hospitalization during the current season if child had met criteria for palivizumab
- Prophylaxis for primary asthma prevention or to reduce subsequent episodes of wheezing in infants and children
- Synagis prophylaxis for prevention of nosocomial disease

| Treatment of s | ymptomatic RSV disease |
|----------------|---|
| Notes | *NOTE: Approval for up to 5 doses per single RSV "season" ** Inform ation regarding RSV season may be found at: • Centers for Disease a nd Prevention (CDC) surveillance reports (http://www.cdc.gov/surveill ance/nrevss/rsv/index.html) • http://uhc-cs-10.uhc.com/sites/cspm/CS SP/Pages/Synagis.aspx ***NOTE: Infants in a neonatal intensive care unit who qualify for prophylaxis may receive the first dose 48 to 72 ho urs before discharge to home or promptly after discharge. If the first d ose is administered in the hospital, this dose will be considered the fir st dose of the maximum 5 dose series for the season. And any subse quent doses received in the hospital setting, are also considered as p art of the maximum 5 dose series. For infants born during the RSV "se ason," fewer than 5 monthly doses may be needed. |

| Product Name: Synagis* | | | | |
|------------------------------------|----------------------------|--|--|--|
| Diagnosis | Chronic Lung Disease (CLD) | | | |
| Guideline Type Prior Authorization | | | | |

1 - ONE of the following:

1.1 ALL of the following for patients age 0 to less than 12 months:

1.1.1 The patient is a preterm infant defined as gestational age less than 32 weeks, 0 days

AND

1.1.2 Patient has developed chronic lung disease (CLD) of prematurity

AND

1.1.3 There was a requirement for greater than 21% oxygen for at least the first 28 days after birth

1.2 ALL of the following for patients age greater than or equal to 12 months to less than 24 months:

1.2.1 The patient was born at less than 32 weeks, 0 days gestation

AND

1.2.2 The patient required at least 28 days of oxygen after birth

AND

1.2.3 The patient continues to require supplemental oxygen, diuretics, or chronic systemic corticosteroid therapy within 6 months of the start of the second RSV "season"

AND

2 - Administered during RSV season**

AND

3 - Monthly dose of Synagis does not exceed 15 milligram per kilogram per dose

AND

4 - Monthly dose of Synagis does not exceed 5 doses per single RSV "season"***

AND

5 - The patient does not meet ONE of the following situations

- Infants and children with hemodynamically insignificant heart disease (e.g., secundum atrial septal defect, small ventricular septal defect, pulmonic stenosis, uncomplicated aortic stenosis, mild coarctation of the aorta, and patent ductus arteriosus)
- Infants with congenital heart disease and cardiac lesions adequately corrected by surgery, unless they continue to require medication for congestive heart failure
- Infants with cardiomyopathy sufficiently mild that they do not require pharmacotherapy

| • • • • | Routine use of prophylaxis in children with Down syndrome [unless qualifying heart disease, CLD, airway clearance issues (the inability to clear secretions from the upper airway because of ineffective cough), or prematurity (less than 29 weeks, 0 days gestation) is present] Routine use of prophylaxis in children with cystic fibrosis (unless indications noted in proven indications above are present) Administration of monthly Synagis prophylaxis after an infant or child has experienced a breakthrough RSV hospitalization during the current season if child had met criteria for palivizumab Prophylaxis for primary asthma prevention or to reduce subsequent episodes of wheezing in infants and children Synagis prophylaxis for prevention of nosocomial disease Treatment of symptomatic RSV disease | | |
|---------|--|---|--|
| Notes | | *NOTE: Approval for up to 5 doses per single RSV "season" ** Inform ation regarding RSV season may be found at: • Centers for Disease a nd Prevention (CDC) surveillance reports (http://www.cdc.gov/surveill ance/nrevss/rsv/index.html) • http://uhc-cs-10.uhc.com/sites/cspm/CS SP/Pages/Synagis.aspx ***NOTE: Infants in a neonatal intensive care unit who qualify for prophylaxis may receive the first dose 48 to 72 ho urs before discharge to home or promptly after discharge. If the first d ose is administered in the hospital, this dose will be considered the fir st dose of the maximum 5 dose series for the season. And any subse quent doses received in the hospital setting, are also considered as p art of the maximum 5 dose series. For infants born during the RSV "se ason," fewer than 5 monthly doses may be needed. | |

| Product Name: Synagis* | | | | |
|------------------------------------|-----------------------------------|--|--|--|
| Diagnosis | is Congenital Heart Disease (CHD) | | | |
| Guideline Type Prior Authorization | | | | |

1 - ONE of the following:

1.1 ONE of the following for patients age 0 to less than 12 months:

1.1.1 Patient has hemodynamically significant congenital heart disease (CHD) including ONE of the following:

- Acyanotic heart disease and receiving medication to control congestive heart failure and will require cardiac surgical procedures
- Moderate to severe pulmonary hypertension

| Documentation that decisions regarding prophylaxis for infants with cyanotic heart defects were made in consultation with a pediatric cardiologist |
|---|
| OR |
| 1.1.2 The patient is undergoing cardiac transplantation during the RSV "season" |
| OR |
| 1.2 BOTH of the following: |
| 1.2.1 The patient is greater than or equal to 12 months to less than 24 months of age: |
| AND |
| 1.2.2 ONE of the following: |
| After cardiac bypass At the conclusion of extracorporeal membrane oxygenation The patient is undergoing cardiac transplantation during the RSV "season" |
| AND |
| 2 - Administered during RSV season** |
| AND |
| 3 - Monthly dose of Synagis does not exceed 15 milligram per kilogram per dose |
| AND |
| 4 - Monthly dose of Synagis does not exceed 5 doses per single RSV "season"*** |
| AND |

| 5 - | The | patient | does | not mee | t ONF | of the | following | situations |
|-----|------|---------|------|---------|-------|--------|-----------|------------|
| • | 1110 | pation | 0000 | not mot | | | ronowing | Situations |

| • | atrial septal defa aortic stenosis, Infants with con- surgery, unless Infants with card Routine use of p disease, CLD, a airway because gestation) is pre- Routine use of p proven indication Administration of a breakthrough for palivizumab Prophylaxis for wheezing in infa Synagis prophy | dren with hemodynamically insignificant heart disease (e.g., secundum ect, small ventricular septal defect, pulmonic stenosis, uncomplicated mild coarctation of the aorta, and patent ductus arteriosus) genital heart disease and cardiac lesions adequately corrected by they continue to require medication for congestive heart failure diomyopathy sufficiently mild that they do not require pharmacotherapy prophylaxis in children with Down syndrome [unless qualifying heart irway clearance issues (the inability to clear secretions from the upper of ineffective cough), or prematurity (less than 29 weeks, 0 days esent] prophylaxis in children with cystic fibrosis (unless indications noted in ns above are present) of monthly Synagis prophylaxis after an infant or child has experienced RSV hospitalization during the current season if child had met criteria primary asthma prevention or to reduce subsequent episodes of ants and children laxis for prevention of nosocomial disease mptomatic RSV disease |
|-------|--|---|
| Notes | | *NOTE: Approval for up to 5 doses per single RSV "season" ** Inform ation regarding RSV season may be found at: • Centers for Disease a nd Prevention (CDC) surveillance reports (http://www.cdc.gov/surveill ance/nrevss/rsv/index.html) • http://uhc-cs-10.uhc.com/sites/cspm/CS SP/Pages/Synagis.aspx ***NOTE: Infants in a neonatal intensive care unit who qualify for prophylaxis may receive the first dose 48 to 72 ho urs before discharge to home or promptly after discharge. If the first d ose is administered in the hospital, this dose will be considered the fir st dose of the maximum 5 dose series for the season. And any subse quent doses received in the hospital setting, are also considered as p art of the maximum 5 dose series. For infants born during the RSV "se ason," fewer than 5 monthly doses may be needed. |

| Product Name: Synagis* | | | | |
|------------------------------------|--|--|--|--|
| Diagnosis | iagnosis Congenital abnormalities of the airway or neuromuscular disease | | | |
| Guideline Type Prior Authorization | | | | |
| | | | | |

- **1** ALL of the following:
 - 1.1 Patient is age 0 to less than 12 months

AND 1.2 Patient has ONE of the following: Neuromuscular disease • A congenital anomaly that impairs the ability to clear secretions from the lower airway because of ineffective cough AND 2 - Administered during RSV season** AND 3 - Monthly dose of Synagis does not exceed 15 milligram per kilogram per dose AND 4 - Monthly dose of Synagis does not exceed 5 doses per single RSV "season"*** AND **5** - The patient does not meet ONE of the following situations Infants and children with hemodynamically insignificant heart disease (e.g., secundum • atrial septal defect, small ventricular septal defect, pulmonic stenosis, uncomplicated aortic stenosis, mild coarctation of the aorta, and patent ductus arteriosus) Infants with congenital heart disease and cardiac lesions adequately corrected by ٠ surgery, unless they continue to require medication for congestive heart failure Infants with cardiomyopathy sufficiently mild that they do not require pharmacotherapy • Routine use of prophylaxis in children with Down syndrome [unless gualifying heart • disease, CLD, airway clearance issues (the inability to clear secretions from the upper airway because of ineffective cough), or prematurity (less than 29 weeks, 0 days gestation) is present] Routine use of prophylaxis in children with cystic fibrosis (unless indications noted in proven indications above are present)

- Administration of monthly Synagis prophylaxis after an infant or child has experienced a breakthrough RSV hospitalization during the current season if child had met criteria for palivizumab
- Prophylaxis for primary asthma prevention or to reduce subsequent episodes of wheezing in infants and children
- Synagis prophylaxis for prevention of nosocomial disease
- Treatment of symptomatic RSV disease

| Notes | *NOTE: Approval for up to 5 doses per single RSV "season" ** Inform ation regarding RSV season may be found at: • Centers for Disease a nd Prevention (CDC) surveillance reports (http://www.cdc.gov/surveill ance/nrevss/rsv/index.html) • http://uhc-cs-10.uhc.com/sites/cspm/CS SP/Pages/Synagis.aspx ***NOTE: Infants in a neonatal intensive care unit who qualify for prophylaxis may receive the first dose 48 to 72 ho urs before discharge to home or promptly after discharge. If the first d ose is administered in the hospital, this dose will be considered the fir st dose of the maximum 5 dose series for the season. And any subse quent doses received in the hospital setting, are also considered as p art of the maximum 5 dose series. For infants born during the RSV "se ason," fewer than 5 monthly doses may be needed. |
|-------|---|

| Product Name: Synagis* | | | |
|------------------------------------|--|--|--|
| Diagnosis | iagnosis Immunocompromised children less than 24 months of age | | |
| Guideline Type Prior Authorization | | | |

- **1** BOTH of the following:
- 1.1 Patient is less than 24 months of age

AND

1.2 The patient is immunocompromised (e.g. receiving cancer chemotherapy, undergoing hematopoietic stem cell transplantation, or solid organ transplantation)

AND

2 - Administered during RSV season**

AND

3 - Monthly dose of Synagis does not exceed 15 milligram per kilogram per dose

AND

4 - Monthly dose of Synagis does not exceed 5 doses per single RSV "season"***

AND

5 - The patient does not meet ONE of the following situations

- Infants and children with hemodynamically insignificant heart disease (e.g., secundum atrial septal defect, small ventricular septal defect, pulmonic stenosis, uncomplicated aortic stenosis, mild coarctation of the aorta, and patent ductus arteriosus)
- Infants with congenital heart disease and cardiac lesions adequately corrected by surgery, unless they continue to require medication for congestive heart failure
- Infants with cardiomyopathy sufficiently mild that they do not require pharmacotherapy
- Routine use of prophylaxis in children with Down syndrome [unless qualifying heart disease, CLD, airway clearance issues (the inability to clear secretions from the upper airway because of ineffective cough), or prematurity (less than 29 weeks, 0 days gestation) is present]
- Routine use of prophylaxis in children with cystic fibrosis (unless indications noted in proven indications above are present)
- Administration of monthly Synagis prophylaxis after an infant or child has experienced a breakthrough RSV hospitalization during the current season if child had met criteria for palivizumab
- Prophylaxis for primary asthma prevention or to reduce subsequent episodes of wheezing in infants and children
- Synagis prophylaxis for prevention of nosocomial disease
- Treatment of symptomatic RSV disease

| *NOTE: Approval for up to 5 doses per single RSV "season" ** Inform ation regarding RSV season may be found at: • Centers for Disease a nd Prevention (CDC) surveillance reports (http://www.cdc.gov/surveill ance/nrevss/rsv/index.html) • http://uhc-cs-10.uhc.com/sites/cspm/CS SP/Pages/Synagis.aspx ***NOTE: Infants in a neonatal intensive care unit who qualify for prophylaxis may receive the first dose 48 to 72 ho urs before discharge to home or promptly after discharge. If the first d ose is administered in the hospital, this dose will be considered the fir st dose of the maximum 5 dose series for the season. And any subse quent doses received in the hospital setting, are also considered as p |
|--|

| art of the maximum 5 dose series. For infants born during the RSV "se ason," fewer than 5 monthly doses may be needed. |
|--|
| uson, newer than o monthly usees may be needed. |

| Product Name: Synagis* | | |
|--|--|--|
| Diagnosis | Cystic fibrosis (CF) | |
| Guideline Type | Prior Authorization | |
| Approval Criteria | | |
| 1 - ONE of the following | g: | |
| 1.1 BOTH of the follow | ving for patients age 0 to less than 12 months: | |
| 1.1.1 Patient has cys | tic fibrosis | |
| | | |
| | AND | |
| | ical evidence of at least ONE of the following: | |
| Chronic lung dis Nutritional comp Failure to thrive pediatric growth | promise defined as weight for length less than the 10th percentile on a | |
| | OR | |
| 1.2 BOTH of the follow | ving: | |
| 1.2.1 Patient is greater than or equal to 12 months to less than 24 months of age | | |
| AND | | |
| 1.2.2 Patient has mai | nifestations of severe lung disease including ONE of the following: | |
| Abnormalities o stable | Abnormalities on chest radiography or chest computed tomography that persists when | |
| Vveight for lengt | th less than the 10th percentile on a pediatric growth chart | |

AND

2 - Administered during RSV season**

AND

3 - Monthly dose of Synagis does not exceed 15 milligram per kilogram per dose

AND

4 - Monthly dose of Synagis does not exceed 5 doses per single RSV "season"***

AND

5 - The patient does not meet ONE of the following situations

- Infants and children with hemodynamically insignificant heart disease (e.g., secundum atrial septal defect, small ventricular septal defect, pulmonic stenosis, uncomplicated aortic stenosis, mild coarctation of the aorta, and patent ductus arteriosus)
- Infants with congenital heart disease and cardiac lesions adequately corrected by surgery, unless they continue to require medication for congestive heart failure
- Infants with cardiomyopathy sufficiently mild that they do not require pharmacotherapy
- Routine use of prophylaxis in children with Down syndrome [unless qualifying heart disease, CLD, airway clearance issues (the inability to clear secretions from the upper airway because of ineffective cough), or prematurity (less than 29 weeks, 0 days gestation) is present]
- Routine use of prophylaxis in children with cystic fibrosis (unless indications noted in proven indications above are present)
- Administration of monthly Synagis prophylaxis after an infant or child has experienced a breakthrough RSV hospitalization during the current season if child had met criteria for palivizumab
- Prophylaxis for primary asthma prevention or to reduce subsequent episodes of wheezing in infants and children
- Synagis prophylaxis for prevention of nosocomial disease
- Treatment of symptomatic RSV disease

| *NOTE: Approval for up to 5 doses per single RSV "season" ** Inform ation regarding RSV season may be found at: • Centers for Disease a nd Prevention (CDC) surveillance reports (http://www.cdc.gov/surveill ance/nrevss/rsv/index.html) • http://uhc-cs-10.uhc.com/sites/cspm/CS |
|---|
| SP/Pages/Synagis.aspx ***NOTE: Infants in a neonatal intensive care |

| unit who qualify for prophylaxis may receive the first dose 48 to 72 ho urs before discharge to home or promptly after discharge. If the first d ose is administered in the hospital, this dose will be considered the fir st dose of the maximum 5 dose series for the season. And any subse quent doses received in the hospital setting, are also considered as p art of the maximum 5 dose series. For infants born during the RSV "se |
|---|
| ason," fewer than 5 monthly doses may be needed. |

2. Background

Benefit/Coverage/Program Information

Additional Information

In most of North America, peak RSV activity typically occurs between November and March, usually beginning in November or December, peaking in January or February, and ending by the end of March or sometime in April. Communities in the southern United States, particularly some communities in the state of Florida, tend to experience the earliest onset of RSV. Data from the Centers for Disease Control and Prevention (CDC) have identified variations in the onset and offset of the RSV "season" in the state of Florida that could affect the timing of Synagis administration. ¹⁰

- Despite varied onsets, the RSV "season" is of the same duration (5 months) in the different regions of Florida.
- On the basis of the epidemiology of RSV in Alaska, particularly in remote regions where the burden of RSV disease is significantly greater than the general US population, the selection of Alaska Native infants eligible for prophylaxis may differ from the remainder of the United States. Clinicians may wish to use RSV surveillance data generated by the state of Alaska to assist in determining onset and end of the RSV season for qualifying infants.
- Limited information is available concerning the burden of RSV disease among Native American populations. However, special consideration may be prudent for Navajo and White Mountain Apache infants in the first year of life.

For analysis of National Respiratory and Enteric Virus Surveillance System (NREVSS) reports in the CDC Morbidity and Mortality Weekly Report, season onset is defined as the first of 2 consecutive weeks during which the mean percentage of specimens testing positive for RSV antigen is \geq 10% and RSV "season" offset is defined as the last of 2 consecutive weeks during which the mean percentage of positive specimens is \geq 10%. Use of specimens to determine the start of the RSV "season" requires that the number of specimens tested be statistically significant.

3. Revision History

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Systane, Refresh, Gonak, Genteal, Tears Naturale

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99534 | Systane, Refresh, Gonak, Genteal, Tears Naturale |
|----------|--|
|----------|--|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: brand Systane, brand Refresh, brand Gonak, brand Genteal, Tears Naturale | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - History of failure, contraindication, or intolerance to ALL of the following:

• Generic equivalents for drops, ointments and gel formulations for Systane, Refresh, Gonak, Genteal, Tears Naturale, and Generic equivalent to the requested brand product

• sodium chloride ophthalmic ointment

2. Revision History

| Date | Notes |
|-----------|-------------------------------------|
| 5/20/2021 | Arizona Medicaid 7.1 Implementation |

Takhzyro

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-104868 Takhzyro

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 3/17/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Takhzyro | |
|------------------------|-----------------------|
| Approval Length | 8 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of hereditary angioedema (HAE) as confirmed by ONE of the following:

1.1 C1 inhibitor (C1-INH) deficiency or dysfunction (Type I or II HAE) as documented by one of the following (per laboratory standard):

• C1-INH antigenic level below the lower limit of normal

• C1-INH functional level below the lower limit of normal

OR

1.2 HAE with normal C1 inhibitor levels and one of the following:

- Confirmed presence of a FXII, angiopoietin-1 or plasminogen gene mutation
- Recurring angioedema attacks that are refractory to high-dose antihistamines with confirmed family history of angioedema

AND

2 - BOTH of the following:

2.1 For prophylaxis against HAE attacks

AND

2.2 Not used in combination with other products indicated for prophylaxis against HAE attacks (e.g., Cinryze, Haegarda)

AND

3 - BOTH of the following:

3.1 Prescriber attests that patient has experienced attacks of a severity and-or frequency such that they would clinically benefit from prophylactic therapy with Takhzyro

AND

3.2 Documentation of baseline HAE attack rate is greater than or equal to one attack per 4 weeks

AND

4 - Prescribed by ONE of the following:

- Immunologist
- Allergist

AND

5 - ONE of the following:

5.1 History of failure, contraindication, or intolerance to Haegarda

OR

5.2 Patient is currently on Takhzyro therapy

| Product Name: Takhzyro | |
|------------------------|---------------------|
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response, defined as a clinically significant reduction in the rate and-or number of hereditary angioedema (HAE) attacks, while on Takhzyro therapy

AND

2 - Reduction in the utilization of on-demand therapies used for acute attacks (e.g., Berinert, Ruconest, Firazyr, Kalbitor) as determined by claims information, while on Takhzyro therapy

AND

3 - Prescribed by ONE of the following:

- Immunologist
- Allergist

AND

4 - BOTH of the following:

4.1 For prophylaxis against hereditary angioedema (HAE) attacks

AND

4.2 Not used in combination with other products indicated for prophylaxis against HAE attacks (e.g., Cinryze, Haegarda

AND

5 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting the number of acute HAE attacks in the previous 6 months while on Takhzyro therapy*

| | *Authorization: • Patient experienced no (zero) acute HAE attacks in t he previous 6 months: Takhzyro 300mg given every 4 weeks for 12 m onths (**Patients experiencing unexpected breakthrough HAE attacks once switched to every 4 week dosing will require additional review to allow for 2 weeks dosing.**) • Patient experienced one or more acute HAE attacks in the previous 6 months: Takhzyro 300mg given every 2 weeks for 6 months |
|--|---|
|--|---|

2. Revision History

| Date | Notes |
|-----------|--|
| 3/16/2022 | Added new GPI. Added submission of records to initial and reauth cri teria |

Talicia and Mycobutin

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-101395 | Talicia and Mycobutin |
|-----------|-----------------------|
|-----------|-----------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 1/4/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Mycobutin | |
|-------------------------|---|
| Diagnosis | Mycobacterium Avium Complex Prophylaxis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of Mycobacterium Avium Complex Prophylaxis

AND

2 - Prescribed by or in consultation with an HIV or infectious disease specialist

AND

3 - Member has failed azithromycin or clarithromycin or is intolerant to the medication due to significant adverse effects or both are contraindicated

AND

4 - If request is for brand Mycobutin and the member is allergic to the generic formulation, the prescriber must submit the FDA MedWatch form

AND

5 - The requested dosage does not exceed 450 mg per day

| Product Name: Mycobutin | |
|---|--|
| Mycobacterium Avium Complex Prophylaxis | |
| 12 month(s) | |
| Reauthorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Member is responding positively to therapy

| Product Name: Mycobutin | |
|-------------------------|---|
| Diagnosis | Mycobacterium Avium Complex Prophylaxis |
| Approval Length | 12 month(s) |
| Guideline Type | Quantity Limit |

1 - For doses that exceed 450mg, the use of this drug is supported by information from ONE of the following appropriate compendia of current literature:

- Food and Drug Administration (FDA) approved indications and limits
- Published practice guidelines and treatment protocols
- Comparative data evaluating the efficacy, type and frequency of side effects and potential drug interactions among alternative products as well as the risks, benefits and potential member outcomes
- Drug Facts and Comparisons
- American Hospital Formulary Service Drug Information
- • United States Pharmacopeia Drug Information
- DRUGDEX Information System
- UpToDate
- MicroMedex
- Peer-reviewed medical literature, including randomized clinical trials, outcomes, research data and pharmacoeconomic studies
- Other drug reference resources

| Product Name: Mycobutin | |
|-------------------------|---|
| Diagnosis | Helicobacter pylori Infection (off-label) |
| Approval Length | 14 Day(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of H. pylori infection

AND

2 - Prescribed in combination with amoxicillin and a proton pump inhibitor

AND

3 - If request is for brand Mycobutin, inability to use generic rifabutin (e.g., contraindications to excipients in rifabutin)

| Product Name: Talicia | | | |
|---|---|--|--|
| Diagnosis | Helicobacter pylori Infection | | |
| Approval Length | 14 Day(s) | | |
| Guideline Type | Prior Authorization | | |
| Approval Criteria | | | |
| 1 - Diagnosis of H. pylo | ori infection | | |
| | | | |
| | AND | | |
| 2 - The medication is p disease specialist | 2 - The medication is prescribed by or in consultation with a gastroenterologist or infectious disease specialist | | |
| | AND | | |
| 3 - One of the following | : | | |
| 3.1 Member has tried 3 first-line treatment regimens listed in the table in background section (One of which must be Rifabutin triple therapy) | | | |
| OR | | | |
| 3.2 Both of the followi | ng: | | |
| 3.2.1 Culture and sensitivity report indicate resistance or lack of susceptibility of H. pylori to all first-line treatment regimens except Rifabutin triple therapy | | | |
| AND | | | |
| 3.2.2 Member must h | ave tried and failed Rifabutin triple therapy | | |

| Product Name: Mycobutin | |
|-------------------------|--------------------------|
| Diagnosis | Tuberculosis (off-label) |
| Approval Length | 12 month(s) |

| Therapy Stage | Initial Authorization | | |
|--|---|--|--|
| Guideline Type | Prior Authorization | | |
| | | | |
| Approval Criteria | Approval Criteria | | |
| 1 - Diagnosis of tuberculosis infection | | | |
| | | | |
| AND | | | |
| | | | |
| 2 - Prescribed by or in | consultation with an HIV or infectious disease specialist | | |
| | | | |
| | AND | | |
| | | | |
| 3 - Current treatment with protease inhibitors or non-nucleoside reverse transcriptase | | | |
| inhibitors (NNRTIs) for the treatment of HIV infection | | | |
| AND | | | |
| | | | |
| 4 - If the request is for brand Mycobutin, inability to use generic rifabutin (e.g., contraindications to excipients in rifabutin). | | | |

| Product Name: Mycobutin | |
|-------------------------|--------------------------|
| Diagnosis | Tuberculosis (off-label) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

1 - Member is responding positively to therapy

2. Background

| Drug Name | Dosing Regimen | Dose Limit/ Maximur Dose |
|----------------------|---|-----------------------------------|
| Azithromycin | MAC: 1,200 mg PO once weekly or 600 mg PO twice weekly | 500 mg/c |
| Clarithromycin | MAC: 500 mg PO BID | 1.5 g/day |
| clarithromycin | H. pylori infection: | See |
| triple regimen | 14 days: | dosing regimen |
| | PPI (standard or double dose) BID; Clarithromycin 500 mg; | |
| | Amoxicillin 1,000 mg or metronidazole 500 mg TID (if penicillin allergy) | |
| bismuth | H. pylori infection: | See |
| quadruple regimen | 10-14 days: | dosing regimen |
| | PPI (standard dose) BID; bismuth subcitrate (120- 300 mg) or subsalicylate (300 mg) QID; tetracycline 500 mg QID; metronidazole 250 mg QID or 500 mg TID-QID | |
| concomitant regimen | <i>H. pylori</i> infection: | See |
| | 10-14 days: | dosing regimen |
| | PPI (standard dose) BID; Clarithromycin 500 mg; Amoxicillin 1,000 mg; | 5 |
| | Metronidazole or tinidazole 500 mg | |
| sequential regimen | <i>H. pylori</i> infection: | See |
| | 5-7 days of BID PPI (standard dose) + amoxicillin 1,000 mg; followed by 5-7 days of BID PPI, clarithromycin 500 mg + metronidazole/tinidazole | dosing regimen |
| hybrid regimen | H. pylori infection: | See |
| | 7 days of BID PPI (standard dose) + amoxicillin 1,000 mg; followed by 7 days of BID PPI, amoxicillin + | dosing regimen |

| | clarithromycin 500 mg + metronidazole/tinidazole | |
|------------------------------------|---|--------------------------|
| levofloxacin triple regimen | <i>H. pylori</i> infection: 10-14 days: PPI (standard dose) BID; levofloxacin 500 mg QD; amoxicillin 1,000 mg BID | See dosing regimen |
| levofloxacin sequential regimen | <i>H. pylori</i> infection: | See dosing regimen |
| | 5-7 days of BID PPI (standard dose) + amoxicillin 1,000 mg; followed by 5-7 days of BID PPI, amoxicillin + metronidazole/tinidazole + QD levofloxacin 500 mg | |
| rifabutin triple | <i>H. pylori</i> infection: 10 days of BID PPI (standard dose) + amoxicillin 1,000 mg BID + rifabutin 300 mg QD | See dosing regimen |

3. Revision History

| Date | Notes |
|----------|----------------------------|
| 1/4/2022 | Corrected Talicia criteria |

Taltz - Arizona

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99799 Taltz - Arizona

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|--------------------|-----------|
| P&T Approval Date: | |
| P&T Revision Date: | |

1. Criteria

| Product Name: Taltz | |
|---------------------|-----------------------|
| Diagnosis | Plaque Psoriasis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - One of the following:

1.1 Submission of medical records (e.g., chart notes, laboratory values) documenting ALL of the following:

1.1.1 Diagnosis of chronic moderate to severe plaque psoriasis

AND

1.1.2 Greater than or equal to 3 percent body surface area involvement, palmoplantar, facial, or genital involvement, or severe scalp psoriasis

AND

1.1.3 BOTH of the following:

1.1.3.1 History of failure to ONE of the following topical therapies, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial):*

- Corticosteroids (e.g., betamethasone, clobetasol, desonide)
- Vitamin D analogs (e.g., calcitriol, calcipotriene)
- Tazarotene
- Calcineurin inhibitors (e.g., tacrolimus, pimecrolimus)
- Anthralin
- Coal tar

AND

1.1.3.2 History of failure to a 3 month trial of methotrexate at the maximally indicated dose within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document date, and duration of trial)*

AND

1.1.4 History of failure, contraindication, or intolerance to ALL of the following preferred biologic products (document drug, date, and duration of trial):*

- Humira (adalimumab)
- Enbrel (etanercept)

• Otezla (apremilast)

AND

1.1.5 History of failure, contraindication, or intolerance to ALL of the following nonpreferred biologic products (document drug, date, and duration of trial): *

Cimzia

AND

1.1.6 Patient is not receiving Taltz in combination with ONE of the following:

- Biologic disease-modifying anti-rheumatic drugs (DMARD) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Cosentyx (secukin umab), Orencia (abatacept)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

1.1.7 Prescribed by or in consultation with a dermatologist

OR

1.2 ALL of the following:

1.2.1 Patient is currently on Taltz therapy as documented by claims history or medical records (document date, and duration of therapy)

AND

1.2.2 Diagnosis of chronic moderate to severe plaque psoriasis

AND

1.2.3 Patient is not receiving Taltz in combination with ONE of the following:

• Biologic DMARD [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Cosentyx (secukinumab), Orencia (abatacept)]

- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND 1.2.4 Prescribed by or in consultation with a dermatologist Notes *Note: Claims history may be used in conjunction as documentation of drug, date, and duration of trials

| Product Name: Taltz | |
|---------------------|---------------------|
| Diagnosis | Plaque Psoriasis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Taltz therapy

AND

- 2 Patient is not receiving Taltz in combination with ONE of the following:
 - Biologic disease-modifying anti-rheumatic drugs (DMARD) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Cosentyx (secukinumab), Orencia (abatacept)]
 - Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
 - Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with a dermatologist

| Product Name: Taltz | |
|---------------------|-----------------------|
| Diagnosis | Psoriatic Arthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

1 - One of the following:

1.1 Submission of medical records (e.g., chart notes, laboratory values) documenting ALL of the following:

1.1.1 Diagnosis of active psoriatic arthritis

AND

1.1.2 History of failure to a 3 month trial of methotrexate at the maximally indicated dose within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document date, and duration of trial)*

AND

1.1.3 History of failure, contraindication, or intolerance to THREE of the following preferred biologic products (document drug, date, and duration of trial):*

- Humira (adalimumab)
- Enbrel (etanercept)
- Otezla (apremilast)
- Xeljanz (tofacitinib)

AND

1.1.4 History of failure, contraindication, or intolerance to THREE of the following non-preferred biologic products (document drug, date, and duration of trial):*

- Orencia
- Cimzia

Simponi AND **1.1.5** Patient is not receiving Taltz in combination with ONE of the following: Biologic disease-modifying anti-rheumatic drugs (DMARD) [e.g., Humira • (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Cosentyx (secukinumab), Orencia (abatacept)] Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)] • Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)] • AND **1.1.6** Prescribed by or in consultation with ONE of the following: Rheumatologist . Dermatologist OR **1.2** ALL of the following: **1.2.1** Patient is currently on Taltz therapy as documented by claims history or medical records (document date, and duration of therapy) AND **1.2.2** Diagnosis of active psoriatic arthritis AND **1.2.3** Patient is not receiving Taltz in combination with ONE of the following: Biologic DMARD [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi • (golimumab), Cosentyx (secukinumab), Orencia (abatacept)] Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)] • Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)] •

| | AND |
|---|---|
| 1.2.4 Prescribed b Rheumatolog Dermatologis | |
| Notes | *Note: Claims history may be used in conjunction as documentation of drug, date, and duration of trials |

| Product Name: Taltz | |
|---------------------|---------------------|
| Diagnosis | Psoriatic Arthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Documentation of positive clinical response to Taltz therapy

AND

- **2** Patient is not receiving Taltz in combination with ONE of the following:
 - Biologic disease-modifying anti-rheumatic drugs (DMARD) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Cosentyx (secukinumab), Orencia (abatacept)]
 - Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
 - Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

- **3** Prescribed by or in consultation with ONE of the following:
 - Rheumatologist
 - Dermatologist

| Product Name: Taltz | |
|---------------------|------------------------|
| Diagnosis | Ankylosing Spondylitis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

1 - One of the following:

1.1 Submission of medical records (e.g., chart notes, laboratory values) documenting ALL of the following:

1.1.1 Diagnosis of active ankylosing spondylitis

AND

1.1.2 History of failure to TWO nonsteroidal anti-inflammatory drugs (NSAIDs) (e.g., ibuprofen, naproxen) at maximally indicated doses, each used for at least 4 weeks within the last 3 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trials)*

AND

1.1.3 History of failure, contraindication, or intolerance to BOTH of the following preferred biologic products (document drug, date, and duration of trial):

- Humira (adalimumab)
- Enbrel (etanercept)

AND

1.1.4 History of failure, contraindication, or intolerance to BOTH of the following non-preferred biologic products (document drug, date, and duration of trial):*

- Cimzia
- Simponi

AND

1.1.5 Patient is not receiving Taltz in combination with ONE of the following:

- Biologic disease-modifying anti-rheumatic drugs (DMARD) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Cosentyx (secukinumab), Orencia (abatacept)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

1.1.6 Prescribed by or in consultation with a rheumatologist

OR

1.2 ALL of the following:

1.2.1 Patient is currently on Taltz therapy as documented by claims history or medical records (document date, and duration of therapy)

AND

1.2.2 Diagnosis of active ankylosing spondylitis

AND

1.2.3 Patient is not receiving Taltz in combination with ONE of the following:

- Biologic DMARD [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Cosentyx (secukinumab), Orencia (abatacept)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

1.2.4 Prescribed by or in consultation with a rheumatologist

| *Note: Claims history may be used in conjunction as documentation of drug, date, and duration of trials |
|---|
| urug, date, and duration of thats |

| Product Name: Taltz | |
|---------------------|------------------------|
| Diagnosis | Ankylosing Spondylitis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Documentation of positive clinical response to Taltz therapy

AND

- **2** Patient is not receiving Taltz in combination with ONE of the following:
 - Biologic disease-modifying anti-rheumatic drugs (DMARD) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Cosentyx (secukinumab), Orencia (abatacept)]
 - Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
 - Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with a rheumatologist

| Product Name: Taltz | |
|---------------------|--|
| Diagnosis | Non-radiographic axial spondyloarthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - One of the following:

1.1 Submission of medical records (e.g., chart notes, laboratory values) documenting ALL of the following:

1.1.1 Diagnosis of active non-radiographic axial spondyloarthritis

AND

1.1.2 History of failure, contraindication, or intolerance to BOTH of the following preferred biologic products (document drug, date, and duration of trial):*

- Humira (adalimumab)
- Enbrel (etanercept)

AND

1.1.3 History of failure, contraindication, or intolerance to BOTH of the following nonpreferred biologic products (document drug, date, and duration of trial):*

- Cimzia
- Simponi

AND

1.1.4 History of failure to TWO nonsteroidal anti-inflammatory drugs (NSAIDs) (e.g., ibuprofen, naproxen) at maximally indicated doses, each used for at least 4 weeks, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trials)*

AND

1.1.5 Patient is not receiving Taltz in combination with ONE of the following:

- Biologic disease-modifying anti-rheumatic drugs (DMARD) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Cosentyx (secukinumab), Orencia (abatacept)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]

1.1.6 Prescribed by or in consultation with a rheumatologist

OR

1.2 ALL of the following:

1.2.1 Patient is currently on Taltz therapy as documented by claims history or medical records (document date, and duration of therapy)

AND

1.2.2 Diagnosis of active non-radiographic axial spondyloarthritis

AND

1.2.3 Patient is not receiving Taltz in combination with ONE of the following:

- Biologic DMARD [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Cosentyx (secukinumab), Orencia (abatacept)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]

AND

1.2.4 Prescribed by or in consultation with a rheumatologist

| Notes | *Note: Claims history may be used in conjunction as documentation of |
|-------|--|
| | drug, date, and duration of trials |

| Product Name: Taltz | |
|---------------------|--|
| Diagnosis | Non-radiographic axial spondyloarthritis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

| Approval Criteria |
|--|
| 1 - Documentation of positive clinical response to Taltz therapy |
| AND |
| 2 - Patient is not receiving Taltz in combination with ONE of the following: |
| Biologic disease-modifying anti-rheumatic drugs (DMARD) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Cosentyx (secukinumab), Orencia (abatacept)] Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)] |
| AND |
| 3 - Prescribed by or in consultation with a rheumatologist |

| Date | Notes |
|-----------|-----------------|
| 6/25/2021 | Updated Program |

Tarceva

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99779 Tarceva

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Tarceva, generic erlotinib | |
|--|-----------------------|
| Diagnosis | Pancreatic Cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of pancreatic cancer

- 2 Disease is ONE of the following:
 - Locally advanced Unresectable •
 - •
 - Metastatic •

AND

3 - Used in combination with Gemzar (gemcitabine)

| Product Name: Brand Tarceva, generic erlotinib | |
|--|---------------------|
| Diagnosis | Pancreatic Cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Tarceva therapy

| Product Name: Brand Tarceva, generic erlotinib | |
|--|------------------------------------|
| Diagnosis | Non-Small Cell Lung Cancer (NSCLC) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | - |

Approval Criteria

1 - Diagnosis of non-small cell lung cancer (NSCLC)

2 - Disease is ONE of the following:

- Metastatic
- Recurrent

AND

- **3** ONE of the following:
 - Tumors are positive for epidermal growth factor receptor (EGFR)exon 19 deletions
 - Tumors are positive for exon 21 (L858R) substitution mutations
 - Tumors are positive for a known sensitizing EGFR mutation (e.g. in-frame exon 20 insertions, exon 18 G719 mutation, exon 21 L861Q mutation)

| Product Name: Brand Tarceva, generic erlotinib | |
|--|------------------------------------|
| Diagnosis | Non-Small Cell Lung Cancer (NSCLC) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Tarceva therapy

| Product Name: Brand Tarceva, generic erlotinib | |
|--|-----------------------|
| Diagnosis | Chordoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of chordoma

| Product Name: Brand Tarceva, generic erlotinib | |
|--|---------------------|
| Diagnosis | Chordoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Tarceva therapy

| Product Name: Brand Tarceva, generic erlotinib | |
|--|-----------------------|
| Diagnosis | Kidney Cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- **1** Both of the following:
 - •
 - Diagnosis of kidney cancer Disease is stage IV or relapsed •

AND

2 - Disease is of non-clear cell histology

| Product Name: Brand Tarceva, generic erlotinib | |
|--|-----------------|
| Diagnosis | Kidney Cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |

| Guideline Type | Prior Authorization |
|----------------|---------------------|
|----------------|---------------------|

1 - Patient does not show evidence of progressive disease while on Tarceva therapy

| Product Name: Brand Tarceva, generic erlotinib | |
|--|--------------------------------------|
| Diagnosis | Central Nervous System (CNS) Cancers |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of metastatic brain cancer from Non-Small Cell Lung Cancer (NSCLC)

AND

2 - ONE of the following:

- Tumors are positive for epidermal growth factor receptor (EGFR) exon 19 deletions
- Tumors are positive for exon 21 (L858R) substitution mutations
- Tumors are positive for a known sensitizing EGFR mutation (e.g., in-frame exon 20 insertions, exon 18 G719 mutation, exon 21 L861Q mutation)

| Product Name: Brand Tarceva, generic erlotinib | |
|--|--------------------------------------|
| Diagnosis | Central Nervous System (CNS) Cancers |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Tarceva therapy

| Product Name: Brand Tarceva, generic erlotinib | |
|--|-----------------------|
| Diagnosis | Vulvar cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of vulvar cancer

| Product Name: Brand Tarceva, generic erlotinib | |
|--|---------------------|
| Diagnosis | Vulvar cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Tarceva therapy

| Product Name: Brand Tarceva, generic erlotinib | |
|--|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Tarceva will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Brand Tarceva, generic erlotinib | |
|--|---------------------------|
| Diagnosis | NCCN Recommended Regimens |

| Approval Length | 12 month(s) |
|-----------------|---------------------|
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

1 - Documentation of positive clinical response to Tarceva therapy

| Date | Notes |
|----------|-------------------------------------|
| 6/2/2021 | Arizona Medicaid 7.1 Implementation |

Targretin

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99771 Targretin

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Targretin caps, generic bexarotene caps, Targretin gel | |
|--|---------------------------|
| Diagnosis | Cutaneous T-Cell Lymphoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of cutaneous T-cell lymphoma (CTCL)

2 - History of failure, contraindication, or intolerance to at least one prior therapy (including skin-directed therapies [e.g., corticosteroids (clobetasol, diflorasone, halobetasol, augmented betamethasone dipropionate), phototherapy, or systemic therapies [e.g. Interferons])

| Product Name: Brand Targretin caps, generic bexarotene caps, Targretin gel | |
|--|---------------------------|
| Diagnosis | Cutaneous T-Cell Lymphoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Patient has not had disease progression while on therapy

| Product Name: Brand Targretin caps, generic bexarotene caps, Targretin gel | |
|--|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Targretin will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Brand Targretin caps, generic bexarotene caps, Targretin gel | |
|--|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Documentation of positive clinical response to Targretin therapy

| Date | Notes |
|----------|-------------------------------------|
| 6/2/2021 | Arizona Medicaid 7.1 Implementation |

Tarpeyo (budesonide)

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-113527 | Tarpeyo (budesonide) |
|-----------|----------------------|
|-----------|----------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 9/8/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Tarpeyo | |
|---|---------------------|
| Approval Length | 9 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of primary immunoglobulin A nephropathy (IgAN) as confirmed by a kidney biopsy | |
| | AND |

2 - Patient is at risk of rapid disease progression [e.g., generally a urine protein-to-creatinine ratio (UPCR) greater than or equal to 1.5 g/g, or by other criteria such as clinical risk scoring using the International IgAN Prediction Tool]

AND

3 - Used to reduce proteinuria

AND

4 - Estimated glomerular filtration rate (eGFR) greater than or equal to 35 mL/min/1.73 m2

AND

5 - One of the following:

5.1 Patient has been on a minimum 90-day trial of a maximally tolerated dose and will continue to receive therapy with one of the following:

- An angiotensin-converting enzyme (ACE) inhibitor (e.g., benazepril, lisinopril)
- An angiotensin II receptor blocker (ARB) (e.g., losartan, valsartan)

OR

5.2 Patient has a contraindication or intolerance to both ACE inhibitors and ARBs

AND

6 - Trial and failure, contraindication, or intolerance to another glucocorticoid (e.g., methylprednisolone, prednisone)

AND

7 - Prescribed by or in consultation with a nephrologist

| Date | Notes |
|----------|---|
| 9/8/2022 | Removed references, no clinical criteria changes. |

Tasigna

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99772 Tasigna

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Tasigna | |
|---|--------------------------|
| Diagnosis | Chronic Myeloid Leukemia |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of chronic myeloid leukemia | |

2 - ONE of the following:

2.1 Patient is not a candidate for imatinib (Gleevec) as attested by physician

OR

2.2 Patient is currently on Tasigna therapy

| Product Name: Tasigna | |
|-----------------------|--------------------------|
| Diagnosis | Chronic Myeloid Leukemia |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Tasigna therapy

| Product Name: Tasigna | |
|-----------------------|---------------------------------------|
| Diagnosis | Gastrointestinal Stromal Tumor (GIST) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of progressive gastrointestinal stromal tumor (GIST)

AND

2 - History of failure, contraindication, or intolerance to ALL of the following:

- Gleevec (imatinib)
- Sutent (sunitinib)
- Stivarga (regorafenib)

| Gastrointestinal Stromal Tumor (GIST) |
|---------------------------------------|
| 12 month(s) |
| Reauthorization |
| Prior Authorization |
| 1 F |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Tasigna therapy

| Product Name: Tasigna | |
|-----------------------|------------------------------------|
| Diagnosis | Acute Lymphoblastic Leukemia (ALL) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of Philadelphia chromosome-positive acute lymphoblastic leukemia (Ph+ALL)

| Product Name: Tasigna | |
|-----------------------|------------------------------------|
| Diagnosis | Acute Lymphoblastic Leukemia (ALL) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| | |

1 - Patient does not show evidence of progressive disease while on Tasigna therapy

| Product Name: Tasigna | |
|-----------------------|--|
| Diagnosis | Myeloid/Lymphoid Neoplasms with Eosinophilia and Tyrosine Kinase Fusion Genes |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of myeloid/lymphoid neoplasms with eosinophilia and ABL1 (gene) rearrangement

AND

2 - Neoplasm is in blast or chronic phase

| Product Name: Tasigna | |
|-----------------------|--|
| Diagnosis | Myeloid/Lymphoid Neoplasms with Eosinophilia and Tyrosine Kinase Fusion Genes |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Tasigna therapy

| Product Name: Tasigna | |
|-----------------------|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |

| Therapy Stage | Initial Authorization |
|----------------|-----------------------|
| Guideline Type | Prior Authorization |

1 - Tasigna will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Tasigna | |
|-----------------------|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | • |

Approval Criteria

1 - Documentation of positive clinical response to Tasigna therapy

| Date | Notes |
|----------|-------------------------------------|
| 6/2/2021 | Arizona Medicaid 7.1 Implementation |

Tavalisse - ARIZONA

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99785 Tavalisse - ARIZONA

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Tavalisse | | |
|--|---------------------------------------|--|
| Diagnosis | Chronic immune thrombocytopenia (ITP) | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of chronic immune thrombocytopenia (ITP) | | |

2 - ONE of the following:

2.1 BOTH of the following:

2.1.1 History of failure, contraindication, or intolerance to ONE of the following:

- Corticosteroids
- Immunoglobulins

AND

2.1.2 History of failure, contraindication, or intolerance to both of the preferred alternatives*

- Nplate (romiplostim)
- Promacta Tablet (eltrombopag)

OR

2.2 Patient is currently on Tavalisse therapy

| Product Name: Tavalisse | |
|-------------------------|---------------------------------------|
| Diagnosis | Chronic immune thrombocytopenia (ITP) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Tavalisse therapy

| Date | Notes |
|----------|--------------------|
| 6/8/2021 | 7/1 Implementation |

Tegsedi

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99652 Tegsedi

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Tegsedi | |
|---|--|
| Hereditary transthyretin-mediated (hATTR) amyloidosis with polyneuropathy | |
| 12 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |
| | |
| | |

1 - BOTH of the following:

| Diagnosis of Hereditary transthyretin-mediated (hATTR) amyloidosis with polyneuropathy | | |
|--|--|--|
| Documentation that the patient has a pathogenic transthyretin (TTR) mutation (e.g., V30M) | | |
| | | |
| AND | | |
| 2 - Prescribed by or in consultation with a neurologist | | |
| AND | | |
| 3 - Documentation of ONE of the following: | | |
| Patient has a baseline polyneuropathy disability (PND) score less than or equal to IIIb Patient has a baseline familial amyloidotic polyneuropathy (FAP) Stage 1 or 2 Patient has a baseline neuropathy impairment (NIS) score greater than or equal to 10 and less than or equal to 130 | | |
| AND | | |
| 4 - Patient has not had a liver transplant | | |
| AND | | |
| 5 - Presence of clinical signs and symptoms of the disease (e.g., peripheral sensorimotor polyneuropathy, autonomic neuropathy, motor disability, etc.) | | |
| AND | | |
| 6 - Patient is not receiving Tegsedi in combination with ONE of the following: | | |
| Oligonucleotide agents [e.g., Onpattro (patisiran)] Tafamidis (e.g., Vyndaqel, Vyndamax) | | |

Product Name: Tegsedi

| Diagnosis | Hereditary transthyretin-mediated (hATTR) amyloidosis with polyneuropathy |
|---|---|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| Approval Criteria 1 - Patient has previou | sly received treatment with Tegsedi |
| | AND |
| 2 - Prescribed by or in | consultation with a neurologist |
| | AND |
| 3 - Documentation of C | ONE of the following: |
| Patient continues to have a polyneuropathy disability (PND) score less than or equal to IIIb Patient continues to have a familial amyloidotic polyneuropathy (FAP) Stage 1 or 2 Patient continues to have a neuropathy impairment (NIS) score greater than or equal to 10 and less than or equal to 130 | |
| | AND |
| 4 - Documentation that the patient has experienced a positive clinical response to Tegsedi therapy (e.g., improved neurologic impairment, motor function, quality of life, slowing of disease progression, etc.) | |
| | AND |
| 5 - Patient is not receiv | ving Tegsedi in combination with ONE of the following: |
| | e agents [e.g., Onpattro (patisiran)] , Vyndaqel, Vyndamax) |

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Temodar (temozolomide)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-106301 | Temodar (temozolomide) | |
|-----------|------------------------|--|
|-----------|------------------------|--|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 4/19/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Temodar, generic temozolomide | |
|---|-----------------------|
| Diagnosis Central Nervous Systems (CNS) Tumor | |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- **1** Diagnosis of ONE of the following types of central nervous system tumors:
 - Intracranial and Spinal Ependymoma (excluding Subependymoma)

- Low-Grade Infiltrative Supratentorial Astrocytoma/Oligodendroglioma
- Medulloblastoma
- Anaplastic Gliomas
- Glioblastoma
- Limited or extensive brain metastases
- Primary CNS (central nervous system) lymphoma

2 - For Brand Temodar requests ONLY: Trial and failure to generic temozolomide (verified via paid pharmacy claims or submission of medical records/chart notes)

| Product Name: Brand Temodar, generic temozolomide | |
|---|-------------------------------------|
| Diagnosis | Central Nervous Systems (CNS) Tumor |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on therapy

| Product Name: Brand Temodar, generic temozolomide | |
|---|--------------------------------------|
| Diagnosis | Cutaneous Melanoma or Uveal Melanoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- **1** Diagnosis of ONE of the following types of melanoma:
 - Metastatic cutaneous melanoma
 - Metastatic uveal melanoma

2 - For Brand Temodar requests ONLY: Trial and failure to generic temozolomide (verified via paid pharmacy claims or submission of medical records/chart notes)

| Product Name: Brand Temodar, generic temozolomide | |
|---|--------------------------------------|
| Diagnosis | Cutaneous Melanoma or Uveal Melanoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on therapy

| Product Name: Brand Temodar, generic temozolomide | |
|---|-----------------------------------|
| Diagnosis | Neuroendocrine and Adrenal Tumors |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of ONE of the following types of neuroendocrine tumors:

- Bronchopulmonary/thymic disease
- Poorly controlled carcinoid syndrome in lung or thymus
- Pancreas
- Pheochromocytoma/paraganglioma
- Poorly differentiated (High Grade)/ large or small cell

AND

2 - For Brand Temodar requests ONLY: Trial and failure to generic temozolomide (verified via paid pharmacy claims or submission of medical records/chart notes)

| Product Name: Brand Temodar, generic temozolomide | |
|---|--|
| Neuroendocrine and Adrenal Tumors | |
| 12 month(s) | |
| Reauthorization | |
| Prior Authorization | |
| | |

1 - Patient does not show evidence of progressive disease while on therapy

| Product Name: Brand Temodar, generic temozolomide | |
|---|-----------------------------|
| Diagnosis | Primary Cutaneous Lymphomas |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of ONE of the following types of primary cutaneous lymphomas:

- Mycosis fungoides (MF)
- Sézary syndrome (SS)
- Primary cutaneous anaplastic large cell lymphoma

AND

2 - For Brand Temodar requests ONLY: Trial and failure to generic temozolomide (verified via paid pharmacy claims or submission of medical records/chart notes)

| Product Name: Brand Temodar, generic temozolomide | |
|---|-----------------------------|
| Diagnosis | Primary Cutaneous Lymphomas |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |

| Guideline Type Prior Authorization |
|------------------------------------|
|------------------------------------|

1 - Patient does not show evidence of progressive disease while on therapy

| Product Name: Brand Temodar, generic temozolomide | |
|---|-----------------------|
| Diagnosis | Soft Tissue Sarcoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - One of the following:

1.1 ONE of the following:

- Diagnosis of angiosarcoma
- Diagnosis of unresectable or progressive retroperitoneal/intra-abdominal soft tissue sarcoma
- Diagnosis of rhabdomyosarcoma
- Undifferentiated pleomorphic sarcoma

OR

1.2 BOTH of the following:

1.2.1 Diagnosis of soft tissue sarcoma of the extremity/superficial trunk, head/neck

AND

1.2.2 ONE of the following:

- Disease is stage IV
- Disease has disseminated metastases

OR

1.3 BOTH of the following:

- Diagnosis of solitary fibrous tumor/hemangiopericytoma Used in combination with Avastin (bevacizumab) •
- •

AND

2 - For Brand Temodar requests ONLY: Trial and failure to generic temozolomide (verified via paid pharmacy claims or submission of medical records/chart notes)

| Product Name: Brand Temodar, generic temozolomide | |
|---|---------------------|
| Diagnosis | Soft Tissue Sarcoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on therapy

| Product Name: Brand Temodar, generic temozolomide | |
|---|-----------------------|
| Diagnosis | Bone Cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- 1 Diagnosis of ONE of the following:
 - Ewing's sarcoma family of tumors •
 - Mesenchymal chondrosarcoma •

2 - ONE of the following:

- Disease has relapsed ٠
- Disease is progressive following primary treatment Used as second-line therapy for metastatic disease •
- •

AND

3 - Used in combination with Campostar (irinotecan)

AND

4 - For Brand Temodar requests ONLY: Trial and failure to generic temozolomide (verified via paid pharmacy claims or submission of medical records/chart notes)

| Product Name: Brand Temodar, generic temozolomide | |
|---|---------------------|
| Diagnosis | Bone Cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on therapy

| Product Name: Brand Temodar, generic temozolomide | |
|---|-----------------------|
| Diagnosis | Uterine Sarcoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| | |

1 - Diagnosis of recurrent or metastatic uterine sarcoma

AND

2 - For Brand Temodar requests ONLY: Trial and failure to generic temozolomide (verified via paid pharmacy claims or submission of medical records/chart notes)

| 5 | ine Sarcoma |
|----------------------|-----------------|
| Approval Length 12 m | |
| | nonth(s) |
| Therapy Stage Reau | uthorization |
| Guideline Type Prior | r Authorization |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on therapy

| Product Name: Brand Temodar, generic temozolomide | |
|---|-------------------------------|
| Diagnosis | Small Cell Lung Cancer (SCLC) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of small cell lung cancer (SCLC)

AND

2 - ONE of the following:

2.1 Relapse within 6 months following complete or partial response or stable disease with initial treatment

OR

2.2 Primary progressive disease

AND

3 - For Brand Temodar requests ONLY: Trial and failure to generic temozolomide (verified via paid pharmacy claims or submission of medical records/chart notes)

| Product Name: Brand Temodar, generic temozolomide | |
|---|-------------------------------|
| Diagnosis | Small Cell Lung Cancer (SCLC) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on therapy

| Product Name: Brand Temodar, generic temozolomide | |
|---|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Temodar will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

AND

2 - For Brand Temodar requests ONLY: Trial and failure to generic temozolomide (verified via paid pharmacy claims or submission of medical records/chart notes)

| Product Name: Brand Temodar, generic temozolomide | |
|---|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to therapy

2. Revision History

| Date | Notes |
|-----------|--|
| 4/19/2022 | Added step through generic for Brand Temodar only. Removed refer ence to drug name in reauth sections. |

Test Strips

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99549 Test Strips

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Non-preferred Test Strip Products | |
|---|--------------|
| Approval Length | 12 month(s) |
| Guideline Type | Step Therapy |
| | |

Approval Criteria

1 - History of failure, contraindication, or intolerance to BOTH of the following:

- True Metrix
- Accu-Chek

| | OR | |
|--|----|--|
| 2 - Patient is on an insulin pump | | |
| | OR | |
| 3 - Patient is visually impaired | | |

| Product Name: Preferred or non-preferred test strip products | |
|--|----------------|
| Approval Length | 12 month(s) |
| Guideline Type | Quantity Limit |

1 - ONE of the following:

1.1 For Insulin Dependent or Pregnant patients, the physician must confirm the patient requires a greater quantity because of more frequent blood glucose testing (e.g., patients on intravenous insulin infusions)

OR

1.2 For Non-Insulin Dependent Patients, ONE the following:

1.2.1 The patient is experiencing or is prone to hypoglycemia or hyperglycemia and requires additional testing to achieve glycemic control

OR

1.2.2 The patient's physician is adjusting medications and the patient requires additional blood glucose testing during this time

1.2.3 The patient's physician is adjusting MNT (medical nutrition therapy) and the patient requires additional blood glucose testing during this time

OR

1.2.4 The patient requires additional testing due to fluctuations in blood glucose due to physical activity or exercise

OR

1.2.5 Other circumstances where prescribing physician confirms that the patient requires a greater quantity because of more frequent blood glucose testing (clinical review required by OptumRx reviewing pharmacist and/or medical director)

2. Revision History

| Date | Notes |
|----------|-------------------------------------|
| 7/1/2021 | Arizona Medicaid 7.1 Implementation |

Testosterone - AZM

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-107461 Testosterone - AZM

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Initial Authorization Prior Authorization

Formulary Note

Guideline Note:

| Effective Date: | 6/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

Product Name: Brand Androgel pump, generic testosterone 1.62% pump (generic Androgel
pump), Brand Androgel gel, generic testosterone gel (generic Androgel), testosterone
enanthate, Androderm, testosterone topical 30mg/act solution, testosterone cypionate, Brand
Testim, generic testosterone 50mg/5gm TD gel (generic Testim), Brand Vogelxo, generic
testosterone TD gel (generic Vogelxo), Jatenzo, TlandoDiagnosisHypogonadismApproval Length12 month(s)

Approval Criteria

Therapy Stage

Guideline Type

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting ONE of the following:

1.1 TWO pre-treatment serum total testosterone levels less than 300 ng/dL (less than 10.4 nmol/L) or less than the reference range for the lab, taken at separate times (Document lab value and date for both levels)

OR

1.2 BOTH of the following:

1.2.1 Patient has a condition that may cause altered sex-hormone binding globulin (SHBG) (e.g., thyroid disorder, HIV disease, liver disorder, diabetes, obesity)

AND

1.2.2 ONE pre-treatment calculated free or bioavailable testosterone level less than 50 pg/mL (less than 5 ng/dL or less than 0.17 nmol/L) or less than the reference range for the lab (This may require treatment to be temporarily held. Document lab value and date)

OR

1.3 Patient has a history of ONE of the following:

- Bilateral orchiectomy
- Panhypopituitarism
- A genetic disorder known to cause hypogonadism (e.g., congenital anorchia, Klinefelter's syndrome)

AND

2 - Patient is NOT taking ONE of the following growth hormones, unless diagnosed with panhypopituitarism:

- Genotropin
- Humatrope
- Norditropin FlexPro
- Nutropin AQ
- Omnitrope

| Saizen | |
|--|--|
| | |
| AND | |
| 3 - Patient is NOT taking with an Aromatase inhibitor (eg, Arimidex [anastrozole], Femara [letrozole], Aromasin [exemestane]) | |
| AND | |
| 4 - Patient was male at birth | |
| AND | |
| 5 - Diagnosis of hypogonadism | |
| AND | |
| 6 - ONE of the following: | |
| Significant reduction in weight (less than 90 percent ideal body weight) (e.g., AIDS wasting syndrome) Osteopenia Osteoporosis Decreased bone density | |
| Decreased bone density Decreased libido Organic cause of testosterone deficiency (eg, injury, tumor, infection, or genetic defects) | |
| AND | |
| 7 - If the request is for generic Androgel, patient must have tried and failed Brand Androgel (verified via paid pharmacy claims or submission of medical records) | |
| AND | |
| 8 - If the request is for JATENZO or TLANDO, patient must have tried and failed Brand | |

Androgel or Androderm (Applies to Jatenzo and Tlando only) (verified via paid pharmacy claims or submission of medical records)

Product Name: Brand Androgel pump, generic testosterone 1.62% pump (generic Androgel pump), Brand Androgel gel, generic testosterone gel (generic Androgel), testosterone enanthate, Androderm, testosterone topical 30mg/act solution, testosterone cypionate, Brand Testim, generic testosterone 50mg/5gm TD gel (generic Testim), Brand Vogelxo, generic testosterone TD gel (generic Vogelxo), Jatenzo, Tlando

| Diagnosis | Gender Dysphoria |
|-----------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient is using hormones to change physical characteristics

AND

2 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of gender dysphoria, as defined by the current version of the Diagnostic and Statistical Manual of Mental Disorders (DSM)

AND

3 - Patient is NOT taking ONE of the following growth hormones, unless diagnosed with panhypopituitarism:

- Genotropin
- Humatrope
- Norditropin FlexPro
- Nutropin AQ
- Omnitrope
- Saizen

AND

4 - Patient is NOT taking with an Aromatase inhibitor (eg, Arimidex [anastrozole], Femara [letrozole], Aromasin [exemestane])

AND

5 - If the request is for generic Androgel, patient must have tried and failed Brand Androgel (verified via paid pharmacy claims or submission of medical records)

AND

6 - If the request is for JATENZO or TLANDO, patient must have tried and failed Brand Androgel or Androderm (Applies to Jatenzo and Tlando only) (verified via paid pharmacy claims or submission of medical records)

Product Name: Brand Androgel pump, generic testosterone 1.62% pump (generic Androgel pump), Brand Androgel gel, generic testosterone gel (generic Androgel), testosterone enanthate, Androderm, testosterone topical 30mg/act solution, testosterone cypionate, Brand Testim, generic testosterone 50mg/5gm TD gel (generic Testim), Brand Vogelxo, generic testosterone TD gel (generic Vogelxo), Jatenzo, Tlando

| Diagnosis | Gender Dysphoria, hypogonadism |
|-----------------|--------------------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting ONE of the following:

1.1 Follow-up total serum testosterone level drawn within the past 12 months is within or below the normal male limits of the reporting lab (document value and date)

OR

1.2 Follow-up total serum testosterone level drawn within the past 12 months is outside of upper male limits of normal for the reporting lab and the dose is adjusted (document value and date)

OR

1.3 BOTH of the following:

1.3.1 Patient has a condition that may cause altered sex-hormone binding globulin (SHBG) (e.g., thyroid disorder, HIV disease, liver disorder, diabetes, obesity)

AND

1.3.2 ONE of the following:

1.3.2.1 Follow-up calculated free or bioavailable testosterone level drawn within the past 12 months is within or below the normal male limits of the reporting lab (document lab value and date)

OR

1.3.2.2 Follow-up calculated free or bioavailable testosterone level drawn within the past 12 months is outside of upper male limits of normal for the reporting lab and the dose is adjusted (document value and date)

AND

2 - Patient is NOT taking ONE of the following growth hormones, unless diagnosed with panhypopituitarism:

- Genotropin
- Humatrope
- Norditropin FlexPro
- Nutropin AQ
- Omnitrope
- Saizen

AND

3 - Patient is NOT taking with an Aromatase inhibitor (eg, Arimidex [anastrozole], Femara [letrozole], Aromasin [exemestane])

2. Revision History

| Date | Notes |
|-----------|--|
| 5/24/2022 | Added Jatenzo and Tlando as NP targets. Added submission of records to criteria. |

Tezspire (tezepelumab-ekko)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-104976 | Tezspire (tezepelumab-ekko) |
|-----------|-----------------------------|
|-----------|-----------------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 4/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Tezspire | |
|------------------------|--|
| 6 Month(s) [A] | |
| Initial Authorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of severe asthma

AND

2 - Patient is 12 years of age or older

AND

- **3** One of the following: [2,3]
 - Patient has had two or more asthma exacerbations requiring systemic corticosteroids (e.g., prednisone) within the past 12 months
 - Prior asthma-related hospitalization within the past 12 months

AND

4 - Patient is currently being treated with one of the following unless there is a contraindication or intolerance to these medications:

4.1 Both of the following: [2,3]

- High-dose inhaled corticosteroid (ICS) (i.e., greater than 500 mcg fluticasone propionate equivalent/day)
- Additional asthma controller medication (e.g., leukotriene receptor antagonist [e.g., montelukast], long-acting beta-2 agonist [LABA] [e.g., salmeterol], tiotropium)

OR

4.2 One maximally-dosed combination ICS/LABA product (e.g., Advair [fluticasone propionate/salmeterol], Symbicort [budesonide/formoterol], Breo Ellipta [fluticasone/vilanterol]) [B]

AND

5 - Prescribed by or in consultation with one of the following:

- Pulmonologist
- Allergist/Immunologist

| Product Name: Tezspire | |
|------------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting positive clinical response to therapy as evidenced by one of the following:

- A reduction in asthma exacerbations
- Improvement in forced expiratory volume in 1 second (FEV1) from baseline

AND

2 - Patient continues to be treated with an inhaled corticosteroid (ICS) (e.g., fluticasone, budesonide) with or without additional asthma controller medication (e.g., leukotriene receptor antagonist [e.g., montelukast], long-acting beta-2 agonist [LABA] [e.g., salmeterol], tiotropium) unless there is a contraindication or intolerance to these medications [4]

AND

3 - Prescribed by or in consultation with one of the following:

- Pulmonologist
- Allergist/Immunologist

2. Endnotes

- A. The Global Initiative for Asthma (GINA) Global Strategy for Asthma Management and Prevention update recommends that patients with asthma should be reviewed regularly to monitor their symptom control, risk factors and occurrence of exacerbations, as well as to document the response to any treatment changes. Ideally, after initiation of treatment, patients should be re-evaluated in 3 to 6 months. [4]
- B. The Global Initiative for Asthma (GINA) Global Strategy for Asthma Management and Prevention guideline recommend patients with severe asthma should be treated with maximal optimized high dose ICS-LABA therapy. [4]

3. Revision History

| Date | Notes |
|-----------|---|
| 3/22/2022 | New Program mirrors ORx with Submission of Records added to initi al and reauth |

Thalomid

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99780 Thalomid

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Thalomid | |
|-----------------------------------|-----------------------|
| Diagnosis | Multiple Myeloma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of multiple myeloma | |

| Product Name: Thalomid | |
|------------------------|---------------------|
| Diagnosis | Multiple Myeloma |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | * |

1 - Patient does not show evidence of progressive disease while on Thalomid therapy

| Product Name: Thalomid | |
|------------------------|---------------------------------|
| Diagnosis | Erythema Nodosum Leprosum (ENL) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of moderate to severe erythema nodosum leprosum (ENL)

AND

2 - ONE of the following:

2.1 Used for acute treatment

OR

2.2 Used as maintenance therapy for prevention & suppression of cutaneous manifestations of ENL recurrence

| Product Name: Thalomid | |
|------------------------|---------------------------------|
| Diagnosis | Erythema Nodosum Leprosum (ENL) |
| Approval Length | 12 month(s) |

| Therapy Stage | Reauthorization |
|----------------|---------------------|
| Guideline Type | Prior Authorization |
| | |

1 - Documentation of positive clinical response to Thalomid therapy

| Product Name: Thalomid | |
|------------------------|------------------------------|
| Diagnosis | Aphthous Stomatitis or Ulcer |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | • |

Approval Criteria

1 - Diagnosis of severe, recurrent aphthous stomatitis or ulcer

| Product Name: Thalomid | |
|------------------------|------------------------------|
| Diagnosis | Aphthous Stomatitis or Ulcer |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Thalomid therapy

| Product Name: Thalomid | |
|------------------------|-----------------------|
| Diagnosis | Pyoderma Gangrenosum |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

1 - Diagnosis of pyoderma gangrenosum

AND

2 - Used as third line treatment

| Product Name: Thalomid | |
|------------------------|----------------------|
| Diagnosis | Pyoderma Gangrenosum |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Thalomid therapy

| Product Name: Thalomid | |
|---|--|
| Cutaneous Manifestations Systemic Lupus Erythematosus (SLE) | |
| 12 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Diagnosis of cutaneous manifestations of systemic lupus erythematosus (SLE)

| Product Name: Thalomid | |
|------------------------|---|
| Diagnosis | Cutaneous Manifestations Systemic Lupus Erythematosus (SLE) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |

| Guideline Type | Prior Authorization |
|-------------------|---------------------|
| | |
| Approval Criteria | |
| | |

1 - Documentation of positive clinical response to Thalomid therapy

| Product Name: Thalomid | |
|------------------------|-----------------------|
| Diagnosis | B-Cell Lymphomas |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of Castleman's Disease (CD)

AND

2 - NOT used as first line therapy

| Product Name: Thalomid | |
|------------------------|---------------------|
| Diagnosis | B-Cell Lymphomas |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Thalomid therapy

| Product Name: Thalomid | |
|------------------------|---------------------------------|
| Diagnosis | Myelofibrosis-Associated Anemia |
| Approval Length | 12 month(s) |

| Therapy Stage | Initial Authorization | |
|---|-----------------------|--|
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of primary | y myelofibrosis | |
| | | |
| | AND | |
| 2 - One of the following: | | |
| 2.1 Both of the following: | | |
| 2.1.1 Serum erythropoietin levels less than 500 mU/mL | | |
| | | |
| | AND | |
| 2.1.2 History of failure, contraindication, or intolerance to erythropoietins [e.g., Procrit (epoetin alfa)] | | |
| | OR | |

2.2 Serum erythropoietin levels greater than or equal to 500 mU/mL

| Product Name: Thalomid | |
|------------------------|---------------------------------|
| Diagnosis | Myelofibrosis-Associated Anemia |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation that member has evidence of symptom improvement or reduction in spleen-liver volume while on Thalomid

| Product Name: Thalomid | | |
|--|--|--|
| Diagnosis | Acquired Immunodeficiency Syndrome (AIDS)- Related Kaposi Sarcoma | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of Acquired Immunodeficiency Syndrome (AIDS)- Related Kaposi Sarcoma | | |
| AND | | |
| 2 - Patient is currently being treated with antiretroviral therapy (ART) | | |
| AND | | |
| 3 - Not used as first line therapy | | |

| AIDS- Related Kaposi Sarcoma |
|------------------------------|
| - |
| 12 month(s) |
| Reauthorization |
| Prior Authorization |
| Re |

1 - Patient does not show evidence of progressive disease while on Thalomid therapy

| Product Name: Thalomid | |
|------------------------|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |

| Guideline Type | Prior Authorization |
|----------------|---------------------|
|----------------|---------------------|

1 - Thalomid will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Thalomid | |
|------------------------|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Thalomid therapy

2. Revision History

| Date | Notes |
|----------|-------------------------------------|
| 6/3/2021 | Arizona Medicaid 7.1 Implementation |

Tobramycin Inhalation - ARIZONA

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99653 Tobramycin Inhalation - ARIZONA

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Bethkis, Kitabis | |
|---------------------------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of cystic fibrosis (CF) | |

Product Name: Brand TOBI Nebulizer Solution, generic tobramycin solution for inhalation, TOBI Podhaler

| Approval Length | 12 month(s) | | |
|--|---------------------------------------|--|--|
| Therapy Stage | Initial Authorization | | |
| Guideline Type | Prior Authorization | | |
| | | | |
| Approval Criteria | | | |
| 1 - Diagnosis of cystic | 1 - Diagnosis of cystic fibrosis (CF) | | |
| | | | |
| | AND | | |
| 2 - Lung infection with positive culture demonstrating Pseudomonas aeruginosa infection | | | |
| AND | | | |
| 3 - History of failure, intolerance, or contraindication to BOTH of the following | | | |
| Brand BethkisKitabis | | | |

Product Name: Brand TOBI Nebulizer Solution, generic tobramycin solution for inhalation,
TOBI PodhalerApproval Length12 month(s)Therapy StageReauthorizationGuideline TypePrior Authorization

Approval Criteria

1 - Documentation of positive clinical response to therapy

2. Revision History

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Topical NSAIDs

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99574 Topical NSAIDs

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Flector Patch, generic diclofenac epolamine 1.3% patch | | | |
|--|---------------------|--|--|
| Approval Length | 2 Week(s) | | |
| Guideline Type | Prior Authorization | | |
| | | | |
| Approval Criteria | Approval Criteria | | |
| 1 - Diagnosis of acute pain due to minor strains, sprains, or contusions | | | |
| | | | |
| AND | | | |
| | | | |

2 - ONE of the following:

2.1 The patient did not receive adequate pain relief when treated with at least three preferred non-steroidal anti-inflammatory drugs (NSAIDs) (An inadequate response to treatment is defined as pain and/or inflammatory symptoms not resolved after 14 days of therapy)

- Diclofenac DR (Generic Voltaren)
- Diclofenac ER (Generic Voltaren ER)
- Etodolac (Generic Lodine)
- Etodolac ER (Generic Lodine ER)
- Fenoprofen (Generic Nalfon)
- Flurbiprofen (Generic Ansaid)
- Ibuprofen
- Indomethacin (Generic Indocin)
- Ketorolac (Generic Toradol)
- Mefenamic (Generic Ponstel)
- Meloxicam (Generic Mobic)
- Nabumetone (Generic Relafen)
- Nabumetone DS (Generic Relafen DS)
- Naproxen (Generic Anaprox)
- Naproxen DR (Generic Anaprox DR)
- Naproxen EC (Generic Anaprox EC)
- Oxaprozin (Generic Daypro)
- Piroxicam (Generic Feldene)
- Sulindac (Generic Clinoril)

OR

2.2 The patient has one of the following risk factors for NSAID-induced adverse GI (gastrointestinal) events:

- Patient is greater than or equal to 65 years of age
- Prior history of peptic, gastric, or duodenal ulcer
- History of NSAID-related ulcer
- History of clinically significant GI (gastrointestinal) bleeding
- Untreated or active H. Pylori gastritis
- Concurrent use of oral corticosteroids (e.g. prednisone, prednisolone, dexamethasone)
- Concurrent use of anticoagulants (e.g. warfarin, heparin)
- Concurrent use of antiplatelets (e.g. aspirin including low-dose, clopidogrel)

| Product Name: Pennsaid 2%, diclofenac sodium soln 1.5% | |
|--|-------------|
| Approval Length | 12 month(s) |

| Guideline Type | Prior Authorization | |
|---|---|--|
| Approval Criteria 1 - Patient has a diagnosis of pain due to osteoarthritis of the knee(s) | | |
| | AND | |
| 2 - ONE of the following | g: | |
| non-steroidal anti-inflar | t receive adequate pain relief when treated with at least three preferred nmatory drugs (NSAIDs) (An inadequate response to treatment is inflammatory symptoms not resolved after 14 days of therapy) | |
| Diclofenac ER (Etodolac (Gene Etodolac ER (G Fenoprofen (Ge Flurbiprofen (Ge Ibuprofen Indomethacin (Ge Ketorolac (Gene Mefenamic (Ge Meloxicam (Gene Nabumetone (Gene Nabumetone DS Naproxen (Gene Naproxen DR (Gene | eneric Lodine ER) eneric Nalfon) eneric Ansaid) Generic Indocin) eric Toradol) neric Ponstel) neric Mobic) Generic Relafen) S (Generic Relafen DS) eric Anaprox) Generic Anaprox DR) Generic Anaprox EC) neric Daypro) eric Feldene) | |
| OR | | |
| 2.2 The patient has one of the following risk factors for NSAID-induced adverse GI (gastrointestinal) events: | | |
| Prior history of p History of NSAI History of clinical | er than or equal to 65 years of age peptic, gastric, or duodenal ulcer D-related ulcer ally significant GI bleeding tive H. Pylori gastritis | |

- Concurrent use of oral corticosteroids (e.g. prednisone, prednisolone, dexamethasone)
- Concurrent use of anticoagulants (e.g. warfarin, heparin)
- Concurrent use of antiplatelets (e.g. aspirin including low-dose, clopidogrel)

AND

3 - Patient has a history of failure, intolerance, or contraindication to diclofenac topical gel 1% (Rx formulation), or Voltaren OTC (over the counter)

| Product Name: generic diclofenac topical gel 1% (Rx formulation), Voltaren OTC | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - The patient has a diagnosis of pain due to osteoarthritis of joints amenable to topical treatment, including but not limited to the hands, knees, ankles, elbows, feet, and wrists

AND

2 - ONE of the following:

2.1 The patient did not receive adequate pain relief when treated with at least three preferred non-steroidal anti-inflammatory drugs (NSAIDs) (An inadequate response to treatment is defined as pain and/or inflammatory symptoms not resolved after 14 days of therapy)

- Diclofenac DR (Generic Voltaren)
- Diclofenac ER (Generic Voltaren ER)
- Etodolac (Generic Lodine)
- Etodolac ER (Generic Lodine ER)
- Fenoprofen (Generic Nalfon)
- Flurbiprofen (Generic Ansaid)
- Ibuprofen
- Indomethacin (Generic Indocin)
- Ketorolac (Generic Toradol)
- Mefenamic (Generic Ponstel)
- Meloxicam (Generic Mobic)
- Nabumetone (Generic Relafen)
- Nabumetone DS (Generic Relaten DS)
- Naproxen (Generic Anaprox)

- Naproxen DR (Generic Anaprox DR)
- Naproxen EC (Generic Anaprox EC)
- Oxaprozin (Generic Daypro)
- Piroxicam (Generic Feldene)
- Sulindac (Generic Clinoril)

OR

2.2 The patient has one of the following risk factors for NSAID-induced adverse GI (gastrointestinal) events:

- Patient is greater than or equal to 65 years of age
- Prior history of peptic, gastric, or duodenal ulcer
- History of NSAID-related ulcer
- History of clinically significant GI bleeding
- Untreated or active H. Pylori gastritis
- Concurrent use of oral corticosteroids (e.g. prednisone, prednisolone, dexamethasone)
- Concurrent use of anticoagulants (e.g. warfarin, heparin)
- Concurrent use of antiplatelets (e.g. aspirin including low-dose, clopidogrel)

Toujeo Solostar, Toujeo Max Solostar, Semglee, Basaglar

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99570 | Toujeo Solostar, | Toujeo Max Solostar, | Semglee, Basaglar |
|----------|------------------|----------------------|-------------------|
|----------|------------------|----------------------|-------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Toujeo Solostar, Toujeo Max Solostar, Semglee, Basaglar | |
|---|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Requests for Toujeo Solostar, Toujeo Max Solostar, Semglee, Basaglar should be denied. The plan's preferred products are.

- Lantus
- Lantus Solostar
- Levemir

- Levemir FlexPen
- Humulin R U 500 (PA Required)

Trelegy Ellipta - ARIZONA

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99500 Trelegy Ellipta - ARIZONA

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Trelegy Ellipta | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| Approval Criteria | |
| Diagnosis of chronic obstructive pulmonary disease (COPD), including chronic bronchitis and-or emphysema | |
| AND | |

2 - History of failure, contraindication, or intolerance to treatment with at least a 30 day trial of both of the following used in combination:

- Stiolto Respimat (tiotropium-olodaterol) Flovent HFA (fluticasone propionate) •
- •

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona Standard to Medicaid Arizona Standard for 7 /1 go live |

Tremfya - AZ Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99730 Tremfya - AZ

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: 12/9/2021 |
|---------------------------|
|---------------------------|

1. Criteria

| Product Name: Tremfya | |
|-----------------------|-----------------------|
| Diagnosis | Plaque Psoriasis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, laboratory values) documenting ALL of the following:

1.1 Diagnosis of chronic moderate to severe plaque psoriasis

AND

1.2 Greater than or equal to 3% body surface area involvement, palmoplantar, facial, or genital involvement, or severe scalp psoriasis

AND

1.3 BOTH of the following:

1.3.1 History of failure to ONE of the following topical therapies, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*:

- Corticosteroids (e.g., betamethasone, clobetasol, desonide)
- Vitamin D analogs (e.g., calcitriol, calcipotriene)
- Tazarotene
- Calcineurin inhibitors (e.g., tacrolimus, pimecrolimus)
- Anthralin
- Coal tar

AND

1.3.2 History of failure to a 3 month trial of methotrexate at the maximally indicated dose within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document date and duration of trial)*

AND

1.4 History of failure, contraindication, or intolerance to ALL of the following preferred biologic products (document drug, date, and duration of trial)*:

- Humira (adalimumab)
- Enbrel (etanercept)
- Otezla (apremilast)

AND

1.5 Patient is not receiving Tremfya in combination with one of the following: Biologic disease modifying antirheumatic drug (DMARD) [e.g., Humira (adalimumab), • Cimzia (certolizumab), Simponi (golimumab), Cosentyx (secukinumab), Orencia (abatacept)] Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)] • Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)] • AND 1.6 Prescribed by or in consultation with a dermatologist OR **2** - All of the following: **2.1** Patient is currently on Tremfya therapy as documented by claims history or medical records (document date and duration of therapy) AND 2.2 Diagnosis of chronic moderate to severe plaque psoriasis AND **2.3** Patient is not receiving Tremfya in combination with one of the following: Biologic disease modifying antirheumatic drug (DMARD) [e.g., Humira (adalimumab), • Cimzia (certolizumab), Simponi (golimumab), Cosentyx (secukinumab), Orencia (abatacept)] Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)] Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)] AND 2.4 Prescribed by or in consultation with a dermatologist

| Notes | *Claims history may be used in conjunction as documentation of drug, date, and duration of trial |
|-------|--|
| | |

| Product Name: Tremfya | |
|-----------------------|---------------------|
| Diagnosis | Plaque Psoriasis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Tremfya therapy

AND

- **2** Patient is not receiving Tremfya in combination with one of the following:
 - Biologic disease modifying antirheumatic drug (DMARD) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Cosentyx (secukinumab), Orencia (abatacept)]
 - Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
 - Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with a dermatologist

| Product Name: Tremfya | | |
|-----------------------|---------------------------|--|
| Diagnosis | Psoriatic Arthritis (PsA) | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |

1 - Submission of medical records (e.g., chart notes, laboratory values) documenting ALL of the following: **1.1** Diagnosis of active psoriatic arthritis AND **1.2** History of failure to a 3 month trial of methotrexate at the maximally indicated dose within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document date and duration of trial) AND **1.3** History of failure, contraindication, or intolerance to THREE of the following preferred biologic products (document drug, date, and duration of trial): Humira (adalimumab) • Enbrel (etanercept) • Otezla (apremilast) • Xeljanz (tofacitinib) • AND **1.4** Patient is not receiving Tremfya in combination with ONE of the following: Biologic disease modifying antirheumatic drug (DMARD) [e.g., Humira (adalimumab), • Cimzia (certolizumab), Simponi (golimumab), Cosentyx (secukinumab), Orencia (abatacept)] Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)] • Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)] • AND **1.5** Prescribed by, or in consultation with, ONE of the following: Rheumatologist Dermatologist •

OR

2 - ALL of the following:

2.1 Patient is currently on Tremfya therapy as documented by claims history or medical records (document date and duration of therapy)

AND

2.2 Diagnosis of active psoriatic arthritis

AND

2.3 Patient is not receiving Tremfya in combination with ONE of the following:

- Biologic DMARD [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Cosentyx (secukinumab), Orencia (abatacept)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

2.4 Prescribed by, or in consultation with, ONE of the following:

- Rheumatologist
- Dermatologist

| Product Name: Tremfya | | |
|-----------------------|---------------------------|--|
| Diagnosis | Psoriatic Arthritis (PsA) | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Reauthorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |

| 1 | - Documentation of | nositive | clinical re | esnonse to | Tremfva | therany |
|---|--------------------|----------|-------------|------------|---------|---------|
| | - Documentation of | positive | Chinical It | esponse io | псппуа | петару |

AND

2 - Patient is not receiving Tremfya in combination with ONE of the following:

- Biologic disease modifying antirheumatic drug (DMARD) [e.g., Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Cosentyx (secukinumab), Orencia (abatacept)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g., Otezla (apremilast)]

AND

3 - Prescribed by or in consultation with ONE of the following:

- Rheumatologist
- Dermatologist

| Date | Notes |
|----------|--------------------|
| 6/3/2021 | 7/1 Implementation |

Tretinoin Capsules - ARIZONA

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99513 Tretinoin Capsules - ARIZONA

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|--------------------|-----------|
| P&T Approval Date: | |
| P&T Revision Date: | |

1. Criteria

| Product Name: Tretinoin capsules | | |
|----------------------------------|------------------------------------|--|
| Diagnosis | Acute Promyelocytic Leukemia (APL) | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |

1 - Diagnosis of acute promyelocytic leukemia

| Product Name: Tretinoin capsules | | |
|------------------------------------|--|--|
| Acute Promyelocytic Leukemia (APL) | | |
| 12 month(s) | | |
| Reauthorization | | |
| Prior Authorization | | |
| | | |

Approval Criteria

1 - Documentation of positive clinical response to tretinoin capsules

| Product Name: Tretinoin capsules | |
|----------------------------------|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Tretinoin capsules will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Tretinoin capsules | |
|----------------------------------|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to tretinoin capsules

| Date | Notes |
|----------|--------------------|
| 4/8/2021 | 7/1 Implementation |

Tretinoin Topical

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99591 Tretinoin Topical

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Retin-A cream and gel* | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - One of the following: | |
| 1.1 Patient is 26 years of age or less | |

| | OR |
|---|--|
| 1.2 Both of the followi | ng: |
| Patient is greateDiagnosis of ac | er than 26 years of age ne vulgaris |
| | AND |
| 2 - The patient must hat ALL of the following: | we a history of therapeutic failure, contraindication, or intolerance to |
| benzoyl peroxide topical clindamycin topical erythromycin | |
| Notes | *Only Brand Covered |

| Date | Notes |
|------------|-----------------------------------|
| 10/29/2021 | Changed effective date to 12/1/21 |

Trikafta

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-103832 Trikafta

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 2/18/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Trikafta | |
|---------------------------------------|-----------------------|
| Diagnosis | Cystic Fibrosis |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of cystic fibrosis (CF) | |

AND

2 - Submission of laboratory results documenting that the patient has at least one F508del mutation in the cystic fibrosis transmembrane conductance regulator (CFTR) gene or a mutation in the CFTR gene that is responsive to Trikafta based on in vitro data

AND

3 - The patient is 6 years of age or older

AND

4 - Prescribed by, or in consultation with, a specialist affiliated with a CF care center

| Product Name: Trikafta | |
|------------------------|---------------------|
| Diagnosis | Cystic Fibrosis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Provider attests that the patient has achieved a clinically meaningful response while on Trikafta therapy to ONE of the following:

- Lung function as demonstrated by percent predicted expiratory volume in 1 second (ppFEV1)
- Body mass index (BMI)
- Pulmonary exacerbations
- Quality of life as demonstrated by Cystic Fibrosis Questionnaire-Revised (CFQ-R) respiratory domain score

AND

2 - Prescribed by, or in consultation with, a specialist affiliated with a cystic fibrosis (CF) care center

| Date | Notes |
|-----------|--|
| 2/17/2022 | Updated age and mutation requirements to reflect labeling update |

Triptans - AZ

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99544 Triptans - AZ

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

Product Name: Brand Amerge, generic sumatriptan nasal spray, brand Imitrex tablets, Brand Imitrex injection, generic sumatriptan 6mg PFS, generic almotriptan, brand Maxalt, brand Maxalt MLT, Onzetra Xsail, brand Relpax, generic eletriptan, brand Treximet, generic sumatriptan naproxen, Zembrace, brand Zomig, brand Zomig ZMT, brand Frova, generic frovatriptan, Tosymra

| Diagnosis | Non-preferred products |
|-----------------|------------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of migraine headaches with or without aura

AND

2 - Patient has a history of failure, contraindication, or intolerance to a trial of at least three preferred products (document drugs, duration, and date of trials)*

- brand Imitex Nasal Spray
- naratriptan (generic Amerge)
- rizatriptan (generic Maxalt)
- sumatriptan (Generic Imitrex)
- zolmitriptan (Generic Zomig)

| cartridge, auto-injec | | | | |
|---|--|--|--|--|
| Diagnosis Migraine Headaches with or without Aura | | | | |
| Approval Length | 12 month(s) | | | |
| Guideline Type | Quantity Limits | | | |
| 1 - Diagnosis of mig | raine headaches with or without aura | | | |
| | AND | | | |
| 2 - Prescribed by or | in consultation with one of the following: | | | |
| | | | | |

- Neurologist
- Pain management specialist

AND

3 - Patient is currently receiving prophylactic therapy with at least ONE of the following:

3.1 Amitriptyline (Elavil)

| OR |
|--|
| 3.2 One of the following beta-blockers: |
| atenolol metoprolol nadolol** propranolol timolol** |
| OR |
| 3.3 Divalproex sodium (Depakote/Depakote ER) |
| OR |
| 3.4 OnabotulinumtoxinA (Botox) *** |
| OR |
| 3.5 Topiramate (Topamax) |
| OR |
| 3.6 Venlafaxine (Effexor/Effexor XR) |
| OR |
| 3.7 Calcitonin gene-related peptide (CGRP) receptor antagonists [e.g., Aimovig (erenumab), Emgality (galcanezumab)] |
| AND |

4 - One of the following:

4.1 Higher dose or quantity is supported in the dosage and administration section of the manufacturer's prescribing information

OR

4.2 Higher dose or quantity is supported by one of the following compendia:

- American Hospital Formulary Service Drug Information
- Thomson Micromedex DrugDex
- Clinical pharmacology
- United States Pharmacopoeia-National Formulary (USP-NF)

OR

4.3 Physician provides evidence from published biomedical literature to support safety and additional efficacy at doses/quantities greater than those approved by the Food and Drug Administration (FDA) for the diagnosis indicated

AND

5 - Physician acknowledges that the potential benefit outweighs the risk associated with the higher dose or quantity

| * See "Quantity Limits" table in background section for quantity limits * * Nadolol and timolol are non-preferred and should not be included in denial to provider *** OpphetulinumtexinA (Retex) is a medical bopofit |
|--|
| denial to provider *** OnabotulinumtoxinA (Botox) is a medical benefit, should not be included in denial to provider |

| Product Name: Brand I cartridge, auto-injector | mitrex (inj, cartridge, auto-injector and PFS), generic sumatriptan (inj, and PFS)* | | |
|--|---|--|--|
| Diagnosis Cluster Headaches | | | |
| Approval Length | 12 month(s) | | |
| Guideline Type | Quantity Limit | | |
| | | | |

Approval Criteria

1 - Diagnosis of cluster headaches

AND

2 - Prescribed by or in consultation with one of the following:

- Neurologist
- Pain management specialist

AND

3 - Patient has experienced at least 2 cluster periods lasting from 7 days to 365 days, separated by pain-free periods lasting at least three months.

AND

4 - One of the following:

4.1 Higher dose or quantity is supported in the dosage and administration section of the manufacturer's prescribing information

OR

4.2 Higher dose or quantity is supported by one of the following compendia:

- American Hospital Formulary Service Drug Information
- Thomson Micromedex DrugDex
- Clinical pharmacology
- United States Pharmacopoeia-National Formulary (USP-NF)

OR

4.3 Physician provides evidence from published biomedical literature to support safety and additional efficacy at doses/quantities greater than those approved by the Food and Drug Administration (FDA) for the diagnosis indicated

AND

5 - Physician acknowledges that the potential benefit outweighs the risk associated with the higher dose or quantity

Notes

* See "Quantity Limits" table in background section for quantity limits

Product Name: Brand Amerge, generic naratriptan, Brand Frova, generic frovatriptan, Brand Imitrex tablets and nasal spray, generic sumatriptan tablets and nasal spray, generic almotriptan, Brand Maxalt and Maxalt MLT, generic rizatriptan and rizatriptan MLT, Onzetra Xsail, Brand Relpax, generic eletriptan, Brand Treximet, generic sumatriptan-naproxen, Zembrace Sym Touch, Brand Zomig and Zomig ZMT, generic zolmitriptan and zolmitriptan ZMT, brand Zomig nasal, generic zolmitriptan nasal spray, Tosymra *

| Approval Length | 12 month(s) |
|-----------------|----------------|
| Guideline Type | Quantity Limit |

Approval Criteria

1 - Diagnosis of migraine headaches with or without aura

AND

- **2** Prescribed by or in consultation with one of the following:
 - Neurologist
 - Pain management specialist

AND

3 - Patient is currently receiving prophylactic therapy with at least ONE of the following:

3.1 Amitriptyline (Elavil)

OR

3.2 One of the following beta-blockers:

| atenolol metoprolol nadolol** propranolol timolol** | |
|---|---|
| | OR |
| 3.3 Divalproex sodium (Depakote/Depakote | ER) |
| | OR |
| 3.4 OnabotulinumtoxinA (Botox) *** | |
| | OR |
| 3.5 Topiramate (Topamax) | |
| | OR |
| 3.6 Venlafaxine (Effexor/Effexor XR) | |
| | OR |
| 3.7 Calcitonin gene-related peptide (CGRP) Emgality (galcanezumab)] | receptor antagonists [e.g., Aimovig (erenumab), |
| | AND |
| 4 - One of the following: | |
| 4.1 Higher dose or quantity is supported in the manufacturer's prescribing information | he dosage and administration section of the |

4.2 Higher dose or quantity is supported by one of the following compendia:

- American Hospital Formulary Service Drug Information
- Thomson Micromedex DrugDex
- Clinical pharmacology
- United States Pharmacopoeia-National Formulary (USP-NF)

OR

OR

4.3 Physician provides evidence from published biomedical literature to support safety and additional efficacy at doses/quantities greater than those approved by the FDA (Food and Drug Administration) for the diagnosis indicated

AND

5 - Physician acknowledges that the potential benefit outweighs the risk associated with the higher dose or quantity

| Notes | * See "Quantity Limits" table in background section for quantity limits * |
|-------|---|
| | * Nadolol and timolol are non-preferred and should not be included in |
| | denial to provider *** OnabotulinumtoxinA (Botox) is a medical benefit, |
| | should not be included in denial to provider |

| Product Name: Brand Zomig nasal spray, generic zolmitriptan nasal spray | | |
|---|--------------|--|
| Approval Length 12 month(s) | | |
| Guideline Type | Step Therapy | |

Approval Criteria

1 - Patient has a history of failure, contraindication, or intolerance to a trial of Imitrex Nasal Spray

2 - If the request is for generic zolmitriptan nasal spray, patient must have tried and failed Brand Zomig Spray

2. Background

| Benefit/Coverage/Program Inform | ation | |
|---------------------------------|----------------|----------------|
| Quantity Limits | | |
| Quantity Limits | | |
| Drug Name | Strength | Quantity Limit |
| Brand Amerge | 1mg, 2.5mg | 9 tabs/month |
| generic naratriptan | | |
| Brand Frova | 2.5mg | 9 tabs/month |
| Generic frovatriptan | | |
| Brand Imitrex tablets | 25mg, 50mg, | 9 tabs/month |
| generic sumatriptan tablets | 100mg | |
| Brand Maxalt | 5mg, 10mg | 9 tabs/month |
| Generic rizatriptan | | |
| Brand Maxalt MLT | 5mg, 10mg | 9 tabs/month |
| Generic rizatriptan ODT | | |
| Generic almotriptan | 6.25mg, 12.5mg | 6 tabs/month |
| Relpax | 20mg, 40mg | 6 tabs/month |
| Generic eletriptan | | |
| Brand Zomig | 2.5mg, 5mg | 6 tabs/month |
| Generic zolmitriptan | | |

| Brand Zomig ZMT Generic zolmitriptan ODT | 2.5mg, 5mg | 6 tabs/month | |
|--|---------------------------|--|--|
| Brand Imitrex Nasal Spray Generic sumatriptan nasal spray | 5mg, 20mg | 6 spray devices/month | |
| Zomig Nasal Spray | 2.5mg, 5mg | 6 spray devices/month | |
| Treximet Generic sumatriptan/naproxen | 85mg/500 mg, 10mg/60mg | 9 tabs/month | |
| Onzetra Xsail | 11mg | 1 box (8 units)/month | |
| Zembrace SymTouch | 3mg | 1 box (4 units)/month | |
| Brand Imitrex Generic Sumatriptan Autoinjector/Cartridge Refills | 4mg/0.5mL 6mg/0.5mL | 8 autoinjectors or cartridge refills/month (4 boxes/month) | |
| Brand Imitrex Generic Sumatriptan Vials | 6mg/0.5mL | 10 vials/month (2 boxes/month) | |
| Generic Sumatriptan Pre-filled Syringe | 6mg/0.5mL | 8 prefilled syringes (4 boxes/month) | |
| Tosymra nasal spray | 10mg | 6 units per month | |

Twyneo (tretinoin-benzoyl peroxide 0.1-3% cream)

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 6/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Twyneo | |
|----------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting both of the following:

- **1.1** Both of the following:
 - Patient is 9 years of age or older

• Diagnosis of acne vulgaris

AND

1.2 The patient must have a history of therapeutic failure, contraindication, or intolerance to ALL of the following (verified via paid pharmacy claims or submission of medical records):

- benzoyl peroxide
- topical clindamycin
- topical erythromycin
- topical tretinoin (Brand Retin-A)

| Date | Notes |
|-----------|-------------|
| 5/24/2022 | New program |

Tykerb

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99775 Tykerb

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Tykerb, generic lapatinib | |
|---|--|
| Breast Cancer | |
| 12 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - One of the following:

1.1 BOTH of the following:

1.1.1 Diagnosis of recurrent or stage IV hormone receptor positive, human epidermal growth factor receptor 2-positive (HER2+) breast cancer

AND

1.1.2 Used in combination with an aromatase inhibitor [e.g., Aromasin (exemestane), Femara (letrozole), Arimidex (anastrozole)]

OR

1.2 BOTH of the following:

1.2.1 Diagnosis of advanced or stage IV human epidermal growth factor receptor 2-positive (HER2+) breast cancer

AND

1.2.2 Used in combination with ONE of the following:

• Herceptin (trastuzumab)

• Xeloda (capecitabine)

| Product Name: Brand Tykerb, generic lapatinib | |
|---|--------------------------------------|
| Diagnosis | Central Nervous System (CNS) Cancers |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - One of the following:

1.1 ALL of the following:

1.1.1 Diagnosis of recurrent, central nervous system (CNS) cancer with metastatic lesions

AND

1.1.2 Tykerb is active against primary (breast) tumor

AND

1.1.3 Used in combination with Xeloda (capecitabine)

OR

1.2 ALL of the following:

1.2.1 Diagnosis of recurrent intracranial or spinal ependymoma (excluding subependymoma)

AND

1.2.2 Patient has received previous radiation therapy

AND

- **1.2.3** Patient has received ONE of the following:
 - Gross total or subtotal resection
 - Localized recurrence
 - Evidence of metastasis (brain, spine, or cerebral spinal fluid)

AND

1.2.4 Used in combination with Temodar (temozolomide)

| Product Name: Brand Tykerb, generic lapatinib | |
|---|----------|
| Diagnosis | Chordoma |

| Approval Length | 12 month(s) |
|-----------------|-----------------------|
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of epidermal growth factor receptor (EGFR) -positive, recurrent chordoma

| Product Name: Brand Tykerb, generic lapatinib | |
|---|-----------------------|
| Diagnosis | Colon Cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of unresectable, advanced or metastatic colon cancer (Human epidermal growth factor receptor 2 (HER2)-amplified and RAS wild type)

AND

2 - Patient has not previously been treated with a Human epidermal growth factor receptor 2 (HER2) inhibitor [e.g., Kanjinti (trastuzumab), Perjeta (pertuzumab), Nerlynx (neratinib)]

AND

3 - Patient has previously been treated with ONE of the following regimens:

- Oxaliplatin-based therapy without irinotecan
- Irinotecan-based therapy without oxaliplatin
- FOLFOXIRI (fluorouracil, leucovorin, oxaliplatin, and irinotecan) regimen
- A fluoropyrimidine without irinotecan or oxaliplatin

4 - Used in combination with trastuzumab

| Product Name: Brand Tykerb, generic lapatinib | |
|---|-----------------------|
| Diagnosis | Rectal Cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of unresectable, advanced or metastatic rectal cancer (Human epidermal growth factor receptor 2 (HER2)-amplified and RAS wild type)

AND

2 - Patient has not previously been treated with a Human epidermal growth factor receptor 2 (HER2) inhibitor [e.g., Kanjinti (trastuzumab), Perjeta (pertuzumab), Nerlynx (neratinib)]

AND

3 - Patient has previously been treated with ONE of the following regimens:

- Oxaliplatin-based therapy without irinotecan
- Irinotecan-based therapy without oxaliplatin
- FOLFOXIRI (fluorouracil, leucovorin, oxaliplatin, and irinotecan) regimen
- A fluoropyrimidine without irinotecan or oxaliplatin

AND

4 - Used in combination with trastuzumab

| Product Name: Brand Tykerb, generic lapatinib | |
|---|--|
| Diagnosis | Breast Cancer, Central Nervous System (CNS) Cancers, Chordoma, Colon Cancer, Rectal Cancer |
| Approval Length | 12 month(s) |

| Therapy Stage | Reauthorization |
|----------------|---------------------|
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Tykerb therapy

| Product Name: Brand Tykerb, generic lapatinib | | |
|---|--|--|
| NCCN Recommended Regimens | | |
| 12 month(s) | | |
| Initial Authorization | | |
| Prior Authorization | | |
| 1 | | |

Approval Criteria

1 - Tykerb will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Brand Tykerb, generic lapatinib | | |
|---|--|--|
| NCCN Recommended Regimens | | |
| 12 month(s) | | |
| Reauthorization | | |
| Prior Authorization | | |
| | | |

Approval Criteria

1 - Documentation of positive clinical response to Tykerb therapy

| Date | Notes |
|----------|-------------------------------------|
| 6/2/2021 | Arizona Medicaid 7.1 Implementation |

Tymlos - Arizona

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99789 | Tymlos - Arizona |
|----------|------------------|
|----------|------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Tymlos | | |
|----------------------|---|--|
| Diagnosis | Postmenopausal patients with osteoporosis at high risk for fracture | |
| Approval Length | 24 Months** | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |

1 - Diagnosis of postmenopausal osteoporosis

AND

2 - ONE of the following:

2.1 Bone Mineral Density (BMD) T-score less than or equal to -3.5 based on BMD measurements from lumbar spine (at least two vertebral bodies), hip (femoral neck, total hip), or radius (one-third radius site) [NOTE: Provider must submit patient specific BMD T-score]

OR

2.2 BOTH of the following:

2.2.1 BMD T-score between -2.5 and -3.5 (BMD T-score greater than -3.5 and less than or equal to -2.5) based on BMD measurements from lumbar spine (at least two vertebral bodies), hip (femoral neck, total hip), or radius (one-third radius site) [NOTE: Provider must submit patient specific BMD T-score]

AND

2.2.2 ONE of the following:

2.2.2.1 History of ONE of the following resulting from minimal trauma:

- Vertebral compression fracture
- Fracture of the hip
- Fracture of the distal radius
- Fracture of the pelvis
- Fracture of the proximal humerus

OR

2.2.2.2 History of failure, contraindication, or intolerance to ALL of the following (Document drug, date, and duration of trial)

- bisphosphonate (e.g. ALENDRONATE, IBANDRONATE)
- selective estrogen receptor modulator (SERM) (e.g RALOXIFENE)
- Prolia (DENOSUMAB)
- Forteo.(TERIPARATIDE)

OR

2.3 ALL of the following:

2.3.1 BMD T-score between -1 and -2.5 (BMD T-score greater than -2.5 and less than or equal to -1) based on BMD measurements from lumbar spine (at least two vertebral bodies), hip (femoral neck, total hip), or radius (one-third radius site) [NOTE: Provider must submit patient specific BMD T-score]

AND

2.3.2 ONE of the following:

2.3.2.1 History of ONE of the following resulting from minimal trauma:

- Vertebral compression fracture
- Fracture of the hip
- Fracture of the distal radius
- Fracture of the pelvis
- Fracture of the proximal humerus

OR

2.3.2.2 ONE of the following Fracture Risk Assessment Tool (FRAX) 10-year fracture probabilities:

- Major osteoporotic fracture at 20 percent or more
- Hip fracture at 3 percent or more

AND

2.3.3 History of failure, contraindication, or intolerance to ALL of the following (Document drug, date, and duration of trial)

- bisphosphonate (e.g. ALENDRONATE, IBANDRONATE)
- selective estrogen receptor modulator (SERM) (e.g RALOXIFENE)
- Prolia (DENOSUMAB)
- Forteo.(TERIPARATIDE)

| | AND |
|--|--|
| | has not exceeded a total of 24 months* of cumulative use of nalogs (e.g., Teriparatide Injection, Forteo, Tymlos) during the patient's |
| | AND |
| 4 - Prescriber attests to the following: The information provided is true and accurate to the best of their knowledge and they understand that OptumRx may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided | |
| Notes | *Claims history may be used in conjunction as documentation of drug, date, and duration of trial **Duration of coverage will be limited to 24 months of cumulative parathyroid hormone analog therapy (e.g., Forte o, Tymlos) in the patient's lifetime |

| Date | Notes |
|----------|------------------|
| 7/1/2021 | Update Guideline |

Uloric

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99501 Uloric

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Uloric, generic febuxostat | |
|---|--------------|
| Approval Length | 12 month(s) |
| Guideline Type | Step Therapy |
| | |
| Approval Criteria | |
| 1 - History of failure, contraindication or intolerance to allopurinol (generic Zyloprim) | |

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona Standard to Medicaid Arizona Standard for 7 /1 go live |

Ultomiris (ravulizumab-cwvz)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-114466 | Ultomiris | (ravulizumab-cwvz) |
|-----------|-----------|--------------------|
|-----------|-----------|--------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Ultomiris | |
|-------------------------|---|
| Diagnosis | Paroxysmal Nocturnal Hemoglobinuria (PNH) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of paroxysmal nocturnal hemoglobinuria (PNH)

2 - Patient is one month of age and older

AND

3 - Prescribed by or in consultation with a hematologist/oncologist

| Product Name: Ultomiris | |
|-------------------------|---|
| Diagnosis | Paroxysmal Nocturnal Hemoglobinuria (PNH) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response (e.g., hemoglobin stabilization, decrease in the number of red blood cell transfusions) to therapy

| Product Name: Ultomiris | |
|-------------------------|---|
| Diagnosis | Atypical Hemolytic Uremic Syndrome (aHUS) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of atypical hemolytic uremic syndrome (aHUS)

AND

2 - Patient is one month of age and older

3 - Prescribed by or in consultation with one of the following:

- Hematologist
- Nephrologist

| Product Name: Ultomiris | |
|-------------------------|---|
| Diagnosis | Atypical Hemolytic Uremic Syndrome (aHUS) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response (e.g., hemoglobin stabilization, decrease in the number of red blood cell transfusions) to therapy

| Product Name: Ultomiris | |
|-------------------------|-------------------------------------|
| Diagnosis | Generalized Myasthenia Gravis (gMG) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of generalized myasthenia gravis (gMG)

AND

2 - Patient is anti-acetylcholine receptor (AChR) antibody positive

3 - One of the following:

3.1 Trial and failure, contraindication, or intolerance to two preferred immunosuppressive therapies (e.g., glucocorticoids, azathioprine, cyclosporine, mycophenolate mofetil, methotrexate, tacrolimus)

OR

3.2 Both of the following:

3.2.1 Trial and failure, contraindication, or intolerance to one preferred immunosuppressive therapy (e.g., glucocorticoids, azathioprine, cyclosporine, mycophenolate mofetil, methotrexate, tacrolimus)

AND

3.2.2 Trial and failure, contraindication, or intolerance to one of the following:

- Chronic plasmapheresis or plasma exchange (PE)
- Intravenous immunoglobulin (IVIG)

AND

4 - Prescribed by or in consultation with a neurologist

| Product Name: Ultomiris | |
|-------------------------|-------------------------------------|
| Diagnosis | Generalized Myasthenia Gravis (gMG) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Documentation of positive clinical response to therapy

| Date | Notes |
|-----------|-------------|
| 9/26/2022 | New Program |

Valchlor

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99693 Valchlor

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|--------------------|-----------|
| P&T Approval Date: | |
| P&T Revision Date: | |

1. Criteria

| Product Name: Valchlor | |
|------------------------|-----------------------------|
| Diagnosis | Primary Cutaneous Lymphomas |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

- 1 Diagnosis of ONE of the following:
 - Chronic or smoldering T-cell leukemia-lymphoma •
 - Primary cutaneous marginal zone or follicle center B-cell lymphoma •
 - Lymphomatoid papulosis (LyP) with extensive lesions Mycosis fungoides (MF)-Sezary syndrome (SS) •
 - •

| Product Name: Valchlor | |
|-----------------------------|--|
| Primary Cutaneous Lymphomas | |
| 12 month(s) | |
| Reauthorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Valchlor

| Product Name: Valchlor | |
|------------------------|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Valchlor will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Valchlor | |
|------------------------|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Valchlor therapy

| Date | Notes |
|----------|--------------------|
| 4/8/2021 | 7/1 Implementation |

Valsartan oral solution

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-114467 | Valsartan oral solution |
|-----------|-------------------------|
|-----------|-------------------------|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Valsartan oral solution | | |
|--|----------------------------------|--|
| Diagnosis | Patients 7 years of age or older | |
| Approval Length | 12 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Patient is 7 years of age or older | | |

| AND | |
|--|--|
| 2 - Patient cannot take solid dosage form due to swallowing issues | |

2. Revision History

| Date | Notes |
|-----------|-------------|
| 9/26/2022 | New program |

Vancomycin - AZ

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99527 Vancomycin - AZ

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Firvanq oral solution, Brand Vancocin, generic vancomycin capsules, vancomycin oral solution | |
|--|---|
| Diagnosis | Clostridioides difficile-associated diarrhea (CDAD) [previously known as Clostridium difficile-associated diarrhea] |
| Approval Length | 10 Day(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of Clostridioides difficile-associated diarrhea (CDAD) [previously known as Clostridium difficile-associated diarrhea]

AND

2 - If the request is for vancomycin oral solution, the prescriber provides a reason or special circumstance the patient cannot use Firvanq and vancomycin capsules*

| Notes | NOTE: *Vancomycin oral solution is non-preferred. Firvanq and vanco |
|-------|---|
| | mycin capsules are preferred. |

| Product Name: Brand Firvanq oral solution, Brand Vancocin, generic vancomycin capsules, vancomycin oral solution | |
|--|---|
| Diagnosis | Clostridioides difficile-associated diarrhea (CDAD) [previously known as Clostridium difficile-associated diarrhea] |
| Approval Length | 12 Week(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Recurrence of Clostridioides difficile infection [previously known as Clostridium difficileassociated diarrhea] after prior treatment with oral vancomycin

| Notes | NOTE: *Vancomycin oral solution is non-preferred. Firvanq and vanco |
|-------|---|
| | mycin capsules are preferred. |

| Product Name: Brand Firvanq oral solution, Brand Vancocin, generic vancomycin capsules, vancomycin oral solution | |
|--|-----------------------|
| Diagnosis | Staphylococcus aureus |
| Approval Length | 10 Day(s) |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of Enterocolitis due to Staphylococcus aureus

| | AND |
|-------|--|
| | or vancomycin oral solution, the prescriber provides a reason or special tient cannot use Firvanq and vancomycin capsules* |
| Notes | NOTE: *Vancomycin oral solution is non-preferred. Firvanq and vanco mycin capsules are preferred. |

2. Revision History

Г

| Date | Notes |
|-----------|--------------------|
| 5/18/2021 | 7/1 Implementation |

1

Vecamyl

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99655 Vecamyl

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Vecamyl | |
|-----------------------|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of moderately severe to severe essential hypertension

OR

2 - Diagnosis of uncomplicated malignant hypertension

| Product Name: Vecamyl | |
|------------------------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| Guideline Type Prior Authorization | |

Approval Criteria

1 - Documentation of a positive clinical response to Vecamyl therapy

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Velphoro (sucroferric oxyhydroxide), Auryxia (ferric citrate)

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99596 Velphoro (sucroferric oxyhydroxide), Auryxia (ferric citrate)

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Velphoro, Auryxia | | | |
|--|---------------------|--|--|
| Approval Length | 12 month(s) | | |
| Guideline Type | Prior Authorization | | |
| | | | |
| Approval Criteria | | | |
| 1 - One of the following | | | |
| Diagnosis of hyperphosphatemia Diagnosis of End Stage Renal Disease | | | |

2 - Adherence to and trial and failure to Sevelamer and Fosrenol at maxiumum dosages (MUST be verified via paid pharmacy claims or submission of medical records)

- Sevelamer Carbonate at the maximum dosage 800mg/15 per day •
- Sevelamer Powder Packets at maximum dosage 2.4gm packet 4 per day
 Fosrenol Chewable Tablets and/or Powder Packets at the maximum dosage 4500 mg per day in divided doses

| Notes | 1. Approval will not be granted for requests based on potential side eff ects, i.e., constipation 2. Approval will not be granted for submitted pri |
|-------|---|
| | or authorizations based on pill burden. Velphoro and Sevelamer are b oth taken 3 times a day. |

| Date | Notes |
|------------|--|
| 11/23/2021 | Added 'verified via paid pharmacy claims or submission of medical re cords' to t/f req |

Veltassa

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-114517 Veltassa

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Veltassa | |
|------------------------|-----------------------------------|
| Diagnosis | Non-Life Threatening Hyperkalemia |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| | |

1 - Diagnosis of non-life threatening hyperkalemia

2 - Where clinically appropriate, medications known to cause hyperkalemia (e.g. angiotensinconverting enzyme inhibitor, angiotensin II receptor blocker, aldosterone antagonist, nonsteroidal anti-inflammatory drugs [NSAIDs]) have been discontinued or reduced to the lowest effective dose

AND

3 - Where clinically appropriate, loop or thiazide diuretic therapy for potassium removal has failed

AND

4 - Patient follows a low potassium diet (less than or equal to 3 grams per day)

AND

5 - History of failure, intolerance, or contraindication to Lokelma

| Product Name: Veltassa | |
|------------------------|-----------------------------------|
| Diagnosis | Non-Life Threatening Hyperkalemia |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient has a positive clinical response to Veltassa therapy

AND

2 - Patient continues to require treatment for hyperkalemia

3 - Where clinically appropriate, medications known to cause hyperkalemia (e.g. angiotensinconverting enzyme inhibitor, angiotensin II receptor blocker, aldosterone antagonist, nonsteroidal anti-inflammatory drugs [NSAIDs])) have been discontinued or reduced to the lowest effective dose

| Date | Notes |
|-----------|--------------------------------------|
| 9/26/2022 | Added step through preferred Lokelma |

Vemlidy

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99731 Vemlidy

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Vemlidy | |
|---|---|
| Diagnosis | Treatment-Naïve Chronic Hepatitis B Infection |
| Approval Length | 12 month(s) |
| Guideline Type | Step Therapy |
| | |
| Approval Criteria | |
| 1 - Patient has a contraindication to entecavir therapy | |

Product Name: Vemlidy

| Diagnosis | Treatment-Experienced Chronic Hepatitis B Infection | |
|--|---|--|
| Approval Length | 12 month(s) | |
| Guideline Type | Step Therapy | |
| | | |
| Approval Criteria | | |
| 1 - One of the following: | | |
| 1.1 BOTH of the following: | | |
| 1.1.1 Patient is currently on Viread therapy | | |
| | AND | |
| 1.1.2 ONE of the follo | owing: | |
| Patient has a creatinine clearance less than 60 mL per minute Patient has a diagnosis of osteoporosis | | |
| | OR | |
| 1.2 Patient is currently | y on Vemlidy therapy | |

| Date | Notes |
|-----------|--------------------|
| 5/18/2021 | 7/1 Implementation |

Ventolin, Proventil, generic albuterol

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99557 Ventolin, Proventil, generic albuterol

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Ventolin, Proventil, generic albuterol | |
|--|---|
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Requests for Ventol preferred product is Pro | in, Proventil, and generic albuterol should be denied. The plan's pair. |
| Notes | ProAir is the only preferred albuterol. Patients on other albuterol formu lations are to be transitioned to ProAir. |

| Date | Notes |
|-----------|-------------------------------------|
| 6/29/2021 | Arizona Medicaid 7.1 Implementation |

Verkazia (cyclosporine ophthalmic emulsion 0.1%)

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-107454 | Verkazia (cyclosporine ophthalmic emulsion 0.1%) |
|-----------|--|
|-----------|--|

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 6/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Verkazia | |
|------------------------|-----------------------|
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting all of the following:

1.1 Diagnosis of moderate to severe vernal keratoconjunctivitis confirmed by the presence of

clinical signs and symptoms (e.g., itching, photophobia, giant papillae at the upper tarsal conjunctiva or at the limbus, thick mucus discharge, conjunctival hyperaemia)

AND

1.2 Trial and failure, contraindication, or intolerance to one of the following (verified via pharmacy paid claims or submission of medical records):

- Topical ophthalmic "dual-acting" mast cell stabilizer and antihistamine (e.g., olopatadine, azelastine)
- Topical ophthalmic mast cell stabilizers (e.g., cromolyn)

AND

1.3 Trial and failure, contraindication, or intolerance, for short term use (up to 2 to 3 weeks), of topical ophthalmic corticosteroids (e.g., dexamethasone, prednisolone, fluorometholone) ((verified via pharmacy paid claims or submission of medical records)

AND

2 - Prescribed by or in consultation with ONE of the following:

- Ophthalmologist
- Optometrist

| Product Name: Verkazia | |
|------------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting positive clinical response to therapy as evidenced by an improvement in clinical signs and symptoms (e.g., itching, photophobia, papillary hypertrophy, mucus discharge, conjunctival hyperaemia)

| Date | Notes |
|-----------|-------------|
| 5/24/2022 | New program |

Vijoice (alpelisib)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-108523 Vijoice (alpelisib)

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 7/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Vijoice | |
|--|-----------------------|
| Approval Length | 6 Months |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of PIK3CA-Related Overgrowth Spectrum (PROS) | |

2 - Submission of documentation of mutation in the PIK3CA gene

AND

3 - Patient is 2 years of age or older

AND

4 - Submission of documentation of severe clinical manifestations (e.g., Congenital Lipomatous Overgrowth, Vascular malformations, Epidermal nevi, Scoliosis/skeletal and spinal [CLOVES], Facial Infiltrating Lipomatosis [FIL], Klippel-Trenaunay Syndrome [KTS], Megalencephaly-Capillary Malformation Polymicrogyria [MCAP])

AND

5 - Prescribed by or in consultation with a physician who specializes in the treatment of PROS

| Product Name: Vijoice | |
|-----------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of documentation of positive clinical response to therapy (e.g., radiological response defined as $a \ge 20\%$ reduction from baseline in the sum of target lesion volume)

AND

2 - Prescribed by or in consultation with a physician who specializes in the treatment of PROS

| Date | Notes |
|-----------|-------------|
| 6/22/2022 | New program |

Vitamin B-12

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99535 Vitamin B-12

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Vitamin B-12 | | | |
|--|---------------------|--|--|
| Approval Length | 12 month(s) | | |
| Guideline Type | Prior Authorization | | |
| | | | |
| Approval Criteria | | | |
| 1 - Provider has submitted lab work documenting a Vitamin B-12 deficiency. | | | |

| Date | Notes |
|-----------|-------------------------------------|
| 5/20/2021 | Arizona Medicaid 7.1 Implementation |

Vitamin C

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99532 Vitamin C

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Vitamin C | | |
|--|--|--|
| Approval Length 12 month(s) | | |
| Guideline Type Prior Authorization | | |
| | | |
| Approval Criteria | | |
| 1 - Provider has submitted lab work documenting a Vitamin C deficiency | | |

| Date | Notes |
|-----------|--------------------|
| 5/19/2021 | 7/1 Implementation |

Vitamin D

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99533 Vitamin D

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Vitamin D | | |
|--|--|--|
| Approval Length 12 month(s) | | |
| Guideline Type Prior Authorization | | |
| | | |
| Approval Criteria | | |
| 1 - Provider has submitted lab work documenting a Vitamin D deficiency | | |

| Date | Notes |
|-----------|--------------------|
| 5/19/2021 | 7/1 Implementation |

Vivjoa (oteseconazole)

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-114156 | Vivjoa (oteseconazole) | |
|-----------|------------------------|--|
| | | |

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Vivjoa | | |
|--|---------------------|--|
| Approval Length | 4 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of recurrent vulvovaginal candidiasis (RVVC) | | |
| AND | | |

| 2 - Patient is NOT of reproductive potential | |
|---|--|
| AND | |
| 3 - Diagnosis of RVVC confirmed by one of the following: | |
| Positive potassium hydroxide (KOH) preparation Vaginal fungal culture | |
| AND | |
| 4 - Patient has experienced 3 or more symptomatic episodes of vulvovaginal candidiasis (VVC) within the past 12 months | |
| AND | |
| 5 - Trial and failure, contraindication, or intolerance to both of the following: | |
| One intravaginal product (e.g., clotrimazole, miconazole, tioconazole, terconazole, boric acid) Oral fluconazole | |

| Date | Notes |
|-----------|-------------|
| 9/26/2022 | New Program |

Vonjo (pacritinib)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-107466 | Vonjo (pacritinib) |
|-----------|--------------------|
|-----------|--------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 6/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Vonjo | |
|---------------------|-----------------------|
| Diagnosis | Myelofibrosis |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting ALL of the following:

| 1.1 Diagnosis of ONE of the following: |
|---|
| Primary myelofibrosis Post-polycythemia vera myelofibrosis Post-essential thrombocythemia myelofibrosis |
| AND |
| 1.2 Disease is intermediate or high risk |
| AND |
| 1.3 Pre-treatment platelet count below 50 x 10^9 L |
| AND |
| 2 - Prescribed by or in consultation with ONE of the following: |
| HematologistOncologist |

| Product Name: Vonjo | |
|---------------------|---------------------|
| Diagnosis | Myelofibrosis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting positive clinical response to therapy (e.g., symptom improvement, spleen volume reduction)

| Product Name: Vonjo | |
|---------------------|---------------------------|
| Diagnosis | NCCN Recommended Regimens |

| Approval Length | 12 month(s) |
|-----------------|---------------------|
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - This drug will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B

| Date | Notes |
|-----------|-------------|
| 5/24/2022 | New Program |

Voquezna Triple Pak (vonoprazan, amoxicillin, clarithromycin), Voquezna Dual Pak (vonoprazan, amoxicillin)

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-114518 Voquezna Triple Pak (vonoprazan, amoxicillin, clarithromycin), Voquezna Dual Pak (vonoprazan, amoxicillin)

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Voquezna Dual Pak, Voquezna Triple Pak | | |
|--|---------------------|--|
| Approval Length | 1 month [A] | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of Helicobacter pylori infection | | |

2 - Trial and failure, contraindication, or intolerance to BOTH of the following first line treatment regimens

- Clarithromycin based therapy (e.g., clarithromycin based triple therapy, clarithromycin based concomitant therapy) [D]
- Bismuth quadruple therapy (e.g., bismuth and metronidazole and tetracycline and proton pump inhibitor [PPI])

| Date | Notes |
|-----------|-------------|
| 9/26/2022 | New Program |

Votrient

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99701 Votrient

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Votrient | |
|---|--|
| Diagnosis | Renal Cell Carcinoma (RCC)/Kidney Cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of renal cell carcinoma (RCC) | |

2 - ONE of the following:

- Disease is relapsed Stage IV disease •
- •

| Product Name: Votrient | | |
|--|---------------------------|--|
| Diagnosis | Soft Tissue Sarcoma (STS) | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| Approval Criteria 1 - One of the following: 1.1 Diagnosis of ONE of the following: Angiosarcoma Alveolar soft part sarcoma Pleomorphic rhabdomyosarcoma Retroperitoneal/Intra-abdominal disease that is unresectable or progressive Soft tissue sarcoma of the extremity/superficial trunk or head/neck with disease that is stage IV or recurrent and has disseminated metastases Solitary fibrous tumor/hemangiopericytoma | | |
| OR | | |
| 1.2 BOTH of the follow | ving: | |
| 1.2.1 Diagnosis of progressive gastrointestinal stromal tumors (GIST) | | |
| AND | | |

1.2.2 History of failure, contraindication, or intolerance to ALL of the following:

- Gleevec (imatinib) •
- Sutent (sunitinib)
- Stivarga (regorafenib) •

| Product Name: Votrient | |
|------------------------|-----------------------|
| Diagnosis | Thyroid Carcinoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- 1 One of the following:
- 1.1 ALL of the following:
- **1.1.1** Diagnosis of ONE of the following:
 - Follicular carcinoma •
 - Hürthle cell carcinoma
 - Papillary carcinoma

AND

- **1.1.2** ONE of the following:
 - Unresectable locoregional recurrent disease •
 - Persistent disease •
 - Metastatic disease •

AND

1.1.3 ONE of the following:

- Patient has symptomatic disease Patient has progressive disease •
- •

AND **1.1.4** ONE of the following: Disease is refractory to radioactive iodine treatment • Distant metastatic disease not amenable to radioactive iodine treatment • OR 1.2 ALL of the following: 1.2.1 Diagnosis of medullary carcinoma AND 1.2.2 ONE of the following: Disease is progressive • Disease is symptomatic with distant metastases • AND **1.2.3** History of failure, contraindication, or intolerance to ONE of the following:

- Caprelsa (vandetanib)
- Cometriq (cabozantinib)

| Product Name: Votrient | | |
|------------------------|-----------------------|--|
| Diagnosis | Uterine Sarcoma | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | Approval Criteria | |

1 - Diagnosis of uterine sarcoma

AND

2 - One of the following:

- Disease is recurrent
- Disease is metastatic

AND

3 - Disease has progressed following previous cytotoxic chemotherapy (e.g., doxorubicin, docetaxel/gemcitabine, etc.)

| Product Name: Votrient | |
|------------------------|-----------------------|
| Diagnosis | Ovarian Cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| | |

Approval Criteria

- **1** Diagnosis of ONE of the following:
 - Epithelial ovarian cancer
 - Fallopian tube cancer
 - Primary peritoneal cancer

AND

2 - ONE of the following:

- Disease is persistent
- Disease is recurrent

| Product Name: Votrient | |
|------------------------|---|
| Diagnosis | Renal Cell Carcinoma (RCC)/Kidney Cancer, Soft Tissue Sarcoma (STS), Thyroid Carcinoma, Uterine Sarcoma, Ovarian Cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Votrient therapy

| Product Name: Votrient | |
|------------------------|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Votrient will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Votrient | |
|------------------------|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Votrient therapy

| Date | Notes |
|-----------|--------------------|
| 4/13/2021 | 7/1 Implementation |

Voxzogo (vosoritide)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-107444 | Voxzogo (vosoritide) |
|-----------|----------------------|
|-----------|----------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 6/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| Product Name: Voxzogo | |
|--|-----------------------|
| Diagnosis | Achondroplasia |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Patient is 5 years of age or older | |

2 - Patient has open epiphyses

AND

3 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of achondroplasia as confirmed by one of the following: [2, 3]

3.1 Both of the following:

3.1.1 Patient has clinical manifestations characteristic of achondroplasia (e.g., macrocephaly, frontal bossing, midface retrusion, disproportionate short stature with rhizomelic shortening of the arms and the legs, brachydactyly, trident configuration of the hands, thoracolumbar kyphosis, and accentuated lumbar lordosis)

AND

3.1.2 Patient has radiographic findings characteristic of achondroplasia (e.g., large calvaria and narrowing of the foramen magnum region, undertubulated, shortened long bones with metaphyseal abnormalities, narrowing of the interpedicular distance of the caudal spine, square ilia and horizontal acetabula, small sacrosciatic notches, proximal scooping of the femoral metaphyses, and short and narrow chest)

OR

3.2 Molecular genetic testing confirmed c.1138G>A or c.1138G>C variant (i.e., p.Gly380Arg mutation) in the fibroblast growth factor receptor-3 (FGFR3) gene

AND

4 - Patient did not have limb-lengthening surgery in the previous 18 months and does not plan on having limb-lengthening surgery while on Voxzogo therapy

AND

5 - Prescribed by or in consultation with one of the following:

- Clinical geneticist
- Endocrinologist
- A physician who has specialized expertise in the management of achondroplasia

| Product Name: Voxzogo | |
|-----------------------|---------------------|
| Diagnosis | Achondroplasia |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient continues to have open epiphyses

AND

2 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting positive clinical response to therapy as evidenced by one of the following:

- Improvement in annualized growth velocity (AGV) compared to baseline
- Improvement in height Z-score compared to baseline

AND

3 - Prescribed by or in consultation with one of the following:

- Clinical geneticist
- Endocrinologist
- A physician who has specialized expertise in the management of achondroplasia

| Product Name: Voxzogo | |
|-----------------------|--------------------------------|
| Diagnosis | Idiopathic Short Stature (ISS) |

| Approval Length | N/A - Requests for non-approvable diagnoses should not be approved |
|-----------------|--|
| Guideline Type | Prior Authorization |
| | |
| | |

Approval Criteria

1 - Requests for coverage for diagnosis of Idiopathic Short Stature (ISS) are not authorized and will not be approved

| Notes | Approval Length: N/A - Requests for Idiopathic Short Stature (ISS) sh |
|-------|---|
| | ould not be approved. Deny as a benefit exclusion. |

| Date | Notes |
|-----------|-------------|
| 5/24/2022 | New Program |

Vtama (tapinarof)

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-112050 Vtama (tapinarof)

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 9/1/2022 |
|-----------------|----------|
|-----------------|----------|

1. Criteria

| 6 month(s) |
|----------------------|
| nitial Authorization |
| Prior Authorization |
| n |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging, paid claims history) documenting a diagnosis of plaque psoriasis

2 - Submission of medical records (e.g., chart notes, lab work, imaging, paid claims history) documenting a minimum duration of a 4 week trial and failure, contraindication, or intolerance to TWO of the following topical therapies:

- Corticosteroids (e.g., betamethasone, clobetasol)
- Vitamin D analogs (e.g., calcitriol, calcipotriene)
- Tazarotene
- Calcineurin inhibitors (e.g., tacrolimus, pimecrolimus)
- Anthralin
- Coal tar

AND

3 - Prescribed by or in consultation with a dermatologist

| Product Name: Vtama | |
|---------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging, paid claims history) documenting positive clinical response to therapy as evidenced by one of the following:

- Reduction in the body surface area (BSA) involvement from baseline
- Improvement in symptoms (e.g., pruritus, inflammation) from baseline

| Date | Notes |
|-----------|-------------|
| 8/19/2022 | New Program |

Vyndaqel and Vyndamax

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99867 | Vyndaqel and Vyndamax |
|--------------|---|
| Formulary | Medicaid - Arizona SP (AZM, AZMREF, AZMDDD) |
| Formulary No | ote |

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Vyndaqel, Vyndamax | |
|----------------------------------|---|
| Diagnosis | Transthyretin (ATTR)-mediated amyloidosis with cardiomyopathy (ATTR-CM) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Diagnosis of transthyretin (ATTR)-mediated amyloidosis with cardiomyopathy (ATTR-CM)

2 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting ONE of the following:

2.1 Documentation that the patient has a pathogenic transthyretin (TTR) mutation (e.g., V30M)

OR

2.2 Cardiac or noncardiac tissue biopsy demonstrating histologic confirmation of ATTR amyloid deposits

OR

2.3 Submission of medical records (e.g., chart notes, lab work, imaging) documenting ALL of the following

2.3.1 Echocardiogram or cardiac magnetic resonance imaging suggestive of amyloidosis

AND

2.3.2 Radionuclide imaging (99mTc-DPD, 99mTc-PYP, or 99m Tc-HMDP) showing grade 2 or 3 cardiac uptake*

AND

2.3.3 Absence of monoclonal protein identified in serum, urine immunofixation (IFE), serum free light chain (sFLC) assay

AND

3 - Prescribed by, or in consultation, with a cardiologist

4 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting presence of clinical signs and symptoms of cardiomyopathy (e.g., heart failure, dyspnea, edema, hepatomegaly, ascites, angina, etc.)

AND

5 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting BOTH of the following:

5.1 ONE of the following:

5.1.1 Patient has New York Heart Association (NYHA) Functional Class I or II heart failure

OR

5.1.2 BOTH of the following:

5.1.2.1 Patient has New York Heart Association (NYHA) Functional Class III heart failure

AND

5.1.2.2 Patient's cardiopulmonary functional status allows patient to ambulate 100 meters or greater in six minutes or less

AND

5.2 Patient has an N-terminal pro-B-type naturetic peptide (NT-proBNP) level greater than or equal to 600 picograms/milliliter

AND

6 - One of the following:

6.1 Paid claims or submission of medical records (e.g., chart notes) verifying patient is not receiving Vyndaqel or Vyndamax in combination with either of the following:

- Onpattro (patisiran)
- Tegsedi (inotersen)

OR

6.2 If the patient is receiving Vyndaqel/Vyndamax in combination with Onpattro (patisiran) or Tegsedi (inotersen), the physician attests that he/she will coordinate care with other specialist(s) involved in the patient's amyloidosis treatment plan to determine optimal long term monotherapy** treatment regimen

| Notes | NOTE: *May require prior authorization and notification ** Referring to |
|-------|---|
| | monotherapy with Vyndaqel/Vyndamax, Onpattro, or Tegsedi |

| Product Name: Vyndaqel, Vyndamax | |
|----------------------------------|---|
| Diagnosis | Transthyretin (ATTR)-mediated amyloidosis with cardiomyopathy (ATTR-CM) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes) documenting that the patient has experienced a positive clinical response to Vyndaqel or Vyndamax (e.g., improved symptoms, quality of life, slowing of disease progression, decreased hospitalizations, etc.)

AND

2 - Prescribed by or in consultation with a cardiologist

AND

3 - Submission of medical records (e.g., chart notes) documenting that patient continues to have New York Heart Association (NYHA) Functional Class I, II, or III heart failure

4 - Paid claims or submission of medical records (e.g., chart notes) verifying patient is not receiving Vyndaqel or Vyndamax in combination with either of the following:

- Onpattro (patisiran)Tegsedi (inotersen)

| Date | Notes |
|-----------|---|
| 12/9/2021 | Added submission of records/paid claims where applicable. |

Vyvgart (efgartigimod alfa-fcab)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-104873 | Vyvgart (efgartigimod alfa-fcab) |
|-----------|----------------------------------|
|-----------|----------------------------------|

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 3/17/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Vyvgart | |
|-----------------------|-----------------------|
| Approval Length | 6 Months [A] |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting diagnosis of generalized myasthenia gravis (gMG)

2 - Patient is anti-acetylcholine receptor (AChR) antibody positive

AND

3 - Prior to administration, patient must be on a stable dose of at least ONE of the following therapies for the treatment of gMG:

- acetylcholinesterase (AChE) inhibitors (e.g., pyridostigmine)
- steroids (e.g., prednisone)
- non-steroidal immunosuppressive therapies (NSISTs) [e.g., azathioprine, cyclosporine, cyclophosphamide)

AND

4 - One of the following:

4.1 Prescribed medication will be administered at 10mg/kg as an intravenous infusion over one hour once weekly for 4 weeks

OR

4.2 In patients weighing 120 kg or more, prescribed medication will be administered at 1200mg per infusion over one hour once weekly for 4 weeks

AND

5 - Prescribed by or in consultation with a neurologist

| Product Name: Vyvgart | |
|-----------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g., chart notes, lab work, imaging) documenting positive clinical response to therapy

AND

2 - One of the following:

2.1 Prescribed medication will be administered at 10mg/kg as an intravenous infusion over one hour once weekly for 4 weeks

OR

2.2 In patients weighing 120 kg or more, prescribed medication will be administered at 1200mg per infusion over one hour once weekly for 4 weeks

2. Endnotes

A. In the ADAPT study all patients received cycle 1, then the time between each treatment cycle was individualized based on the duration of the patient's clinically meaningful response (with a maximum of 3 treatment cycles allowed in 26 week).

| Date | Notes |
|-----------|--|
| 3/31/2022 | New guideline, mirrors ORx with addition of submission of MR req for both initial and reauth |

Wakix

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99732 Wakix

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Wakix | |
|---------------------|-----------------------|
| Diagnosis | Narcolepsy |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g. chart notes, lab values) documenting a diagnosis of narcolepsy with BOTH of the following:

1.1 The patient has daily periods of irrepressible need to sleep or daytime lapses into sleep occurring for at least 3 months

AND

1.2 A mean sleep latency of less than or equal to 8 minutes and two or more sleep onset rapid eye movement (REM) periods (SOREMPs) are found on a MSLT (Multiple Sleep Latency Test) performed according to standard techniques following a normal overnight polysomnogram. A SOREMP (within 15 minutes of sleep onset) on the preceding nocturnal polysomnogram may replace one of the SOREMPs on the MSLT

AND

2 - Physician attestation to the following: Other causes of sleepiness have been ruled out or treated (including but not limited to obstructive sleep apnea, insufficient sleep syndrome, shift work, the effects of substances or medications or their withdrawal, sleep phase disorder, or other sleep disorders)

AND

3 - One of the following:

3.1 Patient has a history of failure, contraindication, or intolerance to all of the following:

3.1.1 One of the following:

- An amphetamine-based stimulant (e.g., amphetamine, dextroamphetamine)
- A methylphenidate-based stimulant

AND

3.1.2 Armodafinil (Nuvigil)

AND

3.1.3 Sunosi (solriamfetol)

OR

3.2 Patient has a history of or potential for a substance abuse disorder

AND

- **4** Prescribed by one of the following:
 - •
 - Neurologist Psychiatrist •
 - Sleep Medicine Specialist •

| Product Name: Wakix | |
|---------------------|---------------------|
| Diagnosis | Narcolepsy |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient has a reduction in symptoms of excessive daytime sleepiness associated with Wakix therapy

| Date | Notes |
|----------|--------------------|
| 6/3/2021 | 7/1 Implementation |

Xalkori

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99695 Xalkori

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|--------------------|-----------|
| P&T Approval Date: | |
| P&T Revision Date: | |

1. Criteria

| Product Name: Xalkori | |
|-----------------------|--|
| Diagnosis | Inflammatory Myofibroblastic Tumor (IMT) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of inflammatory myofibroblastic tumor (IMT) with anaplastic lymphoma kinase (ALK) translocation

| Product Name: Xalkori | |
|-----------------------|------------------------------------|
| Diagnosis | Non-Small Cell Lung Cancer (NSCLC) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of non-small cell lung cancer (NSCLC)

AND

- **2** Disease is ONE of the following:
 - Metastatic
 - Recurrent
 - Advanced

AND

- **3** ONE of the following:
 - Tumor is anaplastic lymphoma kinase (ALK)-positive
 - Tumor is ROS1-positive
 - Tumor is positive for mesenchymal-epithelial transition (MET) amplification
 - Tumor is positive for MET exon 14 skipping mutation

| Product Name: Xalkori | |
|-----------------------|--------------------------------------|
| Diagnosis | Central Nervous System (CNS) Cancers |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |

| Guideline Type | Prior Authorization |
|--|---------------------|
| Approval Criteria | |
| 1 - Diagnosis of metastatic brain cancer from non-small cell lung cancer (NSCLC) | |
| AND | |
| 2 - ONE of the following: | |
| Tumor is anaplastic lymphoma kinase (ALK)-positive Tumor is ROS1-positive | |

| Product Name: Xalkori | Product Name: Xalkori | |
|--|--------------------------------|--|
| Diagnosis | Anaplastic Large Cell Lymphoma | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of anapla | stic large cell lymphoma | |
| | | |
| AND | | |
| 2 - Tumor is anaplastic lymphoma kinase (ALK)-positive | | |
| AND | | |
| 3 - Disease is relapsed or refractory | | |
| | | |

Product Name: Xalkori

| Diagnosis | Inflammatory Myofibroblastic Tumor (IMT), Non-Small Cell Lung Cancer (NSCLC), Central Nervous System (CNS) Cancers, Anaplastic Large Cell Lymphoma |
|-----------------|--|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Patient does not show evidence of progressive disease while on Xalkori therapy

| N Recommended Regimens |
|------------------------|
| |
| onth(s) |
| Authorization |
| Authorization |
| |

Approval Criteria

1 - Xalkori will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Xalkori | |
|-----------------------|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to Xalkori therapy

| Date | Notes |
|----------|--------------------|
| 4/8/2021 | 7/1 Implementation |

Xeljanz, Xeljanz XR (tofacitinib)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-109905 Xeljanz, Xeljanz XR (tofacitinib)

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Xeljanz tablets or Xeljanz XR tablets | |
|---|--|
| Rheumatoid Arthritis (RA) | |
| 12 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - All of the following:

1.1 Diagnosis of moderately to severely active rheumatoid arthritis (RA)

AND

1.2 History of failure to a 3 month trial of methotrexate at the maximally indicated dose within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document date and duration of trial)*

AND

1.3 If the request is for Xeljanz XR, the patient has a history of failure, contraindication, or intolerance to all of the following:

- Humira (adalimumab)
- Enbrel (etanercept)
- Xeljanz (tofacitinib) immediate-release tablets
- Orencia (abatacept)

AND

1.4 Prescribed by or in consultation with a rheumatologist

OR

2 - All of the following:

2.1 Patient is currently on the requested therapy as documented by claims history or medical records (document drug, date, and duration of therapy)*

AND

2.2 Diagnosis of moderately to severely active RA

AND

2.3 Prescribed by or in consultation with a rheumatologist

| Notes | *Claims history may be used in conjunction as documentation of drug, date, and duration of trial |
|-------|--|
| | |

| Product Name: Xeljanz tablets or Xeljanz XR tablets | |
|---|---------------------------|
| Diagnosis | Rheumatoid Arthritis (RA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Documentation of positive clinical response to therapy

AND

2 - Prescribed by or in consultation with a rheumatologist

| Product Name: Xeljanz tablets or Xeljanz XR tablets | |
|---|---------------------------|
| Diagnosis | Psoriatic Arthritis (PsA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - All of the following:

1.1 Diagnosis of active psoriatic arthritis

AND

1.2 History of failure to a 3 month trial of methotrexate at the maximally indicated dose within the last 6 months, unless contraindicated or clinically significant adverse effects are experienced (document date and duration of trial)*

AND

1.3 If the request is for Xeljanz XR, the patient has a history of failure, contraindication, or intolerance to ALL of the following:

- Humira (adalimumab)
- Enbrel (etanercept)
- Otezla (apremilast)
- Xeljanz (tofacitinib) immediate-release
- Orencia (abatacept)

AND

1.4 Prescribed by or in consultation with one of the following:

- Rheumatologist
- Dermatologist

OR

2 - All of the following:

2.1 Patient is currently on the requested therapy as documented by claims history or medical records (document drug, date, and duration of therapy)

AND

2.2 Diagnosis of active psoriatic arthritis

AND

2.3 Prescribed by or in consultation with one of the following:

- Rheumatologist
- Dermatologist

| Notes | *Claims history may be used in conjunction as documentation of drug, date, and duration of trial |
|-------|--|
| | |

| Product Name: Xeljanz tablets or Xeljanz XR tablets | |
|---|---------------------------|
| Diagnosis | Psoriatic Arthritis (PsA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Documentation of positive clinical response to therapy

AND

- **2** Prescribed by or in consultation with one of the following:
 - Rheumatologist Dermatologist •
 - •

| Product Name: Xeljanz tablets or Xeljanz XR tablets | |
|---|-------------------------|
| Diagnosis | Ulcerative Colitis (UC) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- 1 All of the following:
- **1.1** Diagnosis of moderately to severely active ulcerative colitis (UC)

1.2 History of failure to one of the following conventional therapies at maximally indicated doses within the last 3 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*:

- Corticosteroids (e.g., prednisone, methylprednisone, budesonide)
- 6-mercaptopurine (Purinethol)
- Azathioprine (Imuran)
- Aminosalicylates (e.g., mesalamine, sulfasalazine)

AND

1.3 If the request is for Xeljanz XR, the patient has a history of failure, contraindication, or intolerance to Xeljanz (tofacitinib) immediate release tablets

AND

1.4 Prescribed by or in consultation with a gastroenterologist

OR

2 - All of the following:

2.1 Patient is currently on the requested therapy as documented by claims history or medical records (document drug, date, and duration of therapy)

AND

2.2 Diagnosis of moderately to severely active UC

AND

2.3 Prescribed by or in consultation with a gastroenterologist

| Notes | *Claims history may be used in conjunction as documentation of drug, |
|-------|--|
| | date, and duration of trial |

Product Name: Xeljanz tablets or Xeljanz XR tablets

| Diagnosis | Ulcerative Colitis (UC) |
|-----------------|-------------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Documentation of positive clinical response to therapy

AND

2 - Prescribed by or in consultation with a gastroenterologist

| Product Name: Xeljanz tablets or Xeljanz XR tablets | |
|---|-----------------------------|
| Diagnosis | Ankylosing Spondylitis (AS) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of active ankylosing spondylitis

AND

2 - Prescribed by or in consultation with a rheumatologist

AND

3 - One of the following:

3.1 Both of the following:

3.1.1 Trial and failure, contraindication, or intolerance to TWO nonsteroidal antiinflammatory drugs (NSAIDs) (e.g., ibuprofen, naproxen)

AND

3.1.2 If the request is for Xeljanz XR, the patient has a history of failure, contraindication, or intolerance to all of the following:

- Humira (adalimumab)
- Enbrel (etanercept)
- Xeljanz (tofacitinib) immediate-release tablets

OR

3.2 Patient is currently on the requested therapy as documented by claims history or medical records (document drug, date, and duration of therapy)*

| Notes | *Claims history may be used in conjunction as documentation of drug, |
|-------|--|
| | date, and duration of trial |

| Product Name: Xeljanz tablets or Xeljanz XR tablets | |
|---|-----------------------------|
| Diagnosis | Ankylosing Spondylitis (AS) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Documentation of positive clinical response to therapy

AND

2 - Prescribed by or in consultation with a rheumatologist

| Product Name: Xeljanz tablets and oral solution | |
|---|--|
| Diagnosis | Polyarticular Juvenile Idiopathic Arthritis (PJIA) |
| Approval Length | 12 month(s) |

| Therapy Stage | Initial Authorization |
|--|--|
| Guideline Type | Prior Authorization |
| Approval Criteria 1 - Diagnosis of active ankylosing spondylitis | |
| | AND |
| 2 - Prescribed by or in | consultation with a rheumatologist |
| | AND |
| 3 - One of the following | : |
| 3.1 Both of the followi | ng: |
| 3.1.1 Trial and failure DMARDs: | , contraindication, or intolerance to one of the following nonbiologic |
| leflunomidemethotrexate | |
| | AND |
| 3.1.2 History of failure solution ONLY): | e, contraindication, or intolerance to all of the following (applies to oral |
| Humira (adalimi Enbrel (etanerc Xeljanz (tofaciti Orencia (abatac | ept) nib) immediate-release tablets |
| | OR |

3.2 Patient is currently on the requested therapy as documented by claims history or medical records (document drug, date, and duration of therapy)*

| Notes | *Claims history may be used in conjunction as documentation of drug, date, and duration of trial |
|-------|--|
| | |

| Product Name: Xeljanz tablets and oral solution | |
|---|--|
| Diagnosis | Polyarticular Juvenile Idiopathic Arthritis (PJIA) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Documentation of positive clinical response to therapy

AND

2 - Prescribed by or in consultation with a rheumatologist

| Date | Notes |
|-----------|---|
| 9/26/2022 | Added criteria for PJIA. Updated product lists. |

Xenazine

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99657 Xenazine

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Xenazine, generic tetrabenazine | |
|---|---|
| Diagnosis | Chorea associated with Huntington's Disease |
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of chorea in patients with Huntington's disease | |

Product Name: Brand Xenazine, generic tetrabenazine

| Diagnosis | Tardive Dyskinesia (Off Label) |
|---|--|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of tardive | dyskinesia |
| | |
| | AND |
| | |
| 2 - One of the following | i: |
| | tent symptoms of tardive dyskinesia despite a trial of dose reduction, ation of the offending medication |
| | OR |
| 2.2 Patient is not a ca offending medication | ndidate for a trial of dose reduction, tapering, or discontinuation of the |
| | AND |
| 3 - Prescribed by or in | consultation with one of the following: |
| NeurologistPsychiatrist | |

| Product Name: Brand Xenazine, generic tetrabenazine | |
|---|--------------------------------|
| Diagnosis | Tardive Dyskinesia (Off Label) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| | |

1 - Documentation of positive clinical response to therapy

| Product Name: Brand Xenazine, generic tetrabenazine | |
|---|---------------------------------|
| Diagnosis | Tourette's syndrome (off-label) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient has tics associated with Tourette's syndrome

AND

2 - History of failure, contraindication, or intolerance to Haldol (haloperidol)

AND

3 - Prescribed by or in consultation with one of the following:

- •
- Neurologist Psychiatrist •

| Product Name: Brand Xenazine, generic tetrabenazine | |
|---|---------------------------------|
| Diagnosis | Tourette's syndrome (off-label) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

Approval Criteria

1 - Documentation of positive clinical response to therapy

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Xenleta

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99529 Xenleta

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Xenleta | |
|---|--|
| Diagnosis | Community-acquired bacterial pneumonia |
| Approval Length | 7 Day(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - One of the following: | |
| 1.1 For continuation of therapy upon hospital discharge | |

OR

1.2 As continuation of therapy when transitioning from intravenous antibiotics that are shown to be sensitive to the cultured organism for the requested indication

OR

1.3 All of the following:

1.3.1 Diagnosis of community-acquired bacterial pneumonia (CABP)

AND

1.3.2 Infection caused by an organism that is confirmed to be or likely to be susceptible to treatment with Xenleta

AND

1.3.3 History of failure, contraindication, or intolerance to three of the following antibiotics:

- Amoxicillin
- A macrolide
- Doxycycline
- A fluoroquinolone
- Combination therapy with amoxicillin/clavulanate or cephalosporin AND a macrolide or doxycycline

| Product Name: Xenleta* | | | |
|---------------------------|---------------------|--|--|
| Diagnosis | Off-Label Uses | | |
| Guideline Type | Prior Authorization | | |
| | | | |
| Approval Criteria | | | |
| 1 - One of the following: | | | |

| 1.1 For continuation of therapy upon hospital discharge | |
|--|--|
| OR | |
| 1.2 As continuation of therapy when transitioning from intravenous antibiotics that are shown to be sensitive to the cultured organism for the requested indication | |
| OR | |
| 1.3 The medication is being prescribed by or in consultation with an infectious disease specialist | |
| Notes | *Approval Duration: Based on provider recommended treatment durati ons, not to exceed 6 months |

| Date | Notes |
|-----------|--------------------|
| 5/18/2021 | 7/1 Implementation |

Xermelo

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99658 Xermelo

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Xermelo | |
|-----------------------|-----------------------------|
| Diagnosis | Carcinoid Syndrome Diarrhea |
| Approval Length | 6 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of carcinoid syndrome diarrhea

AND

2 - Diarrhea is inadequately controlled with somatostatin analog therapy (e.g., octreotide, Sandostatin LAR, Somatuline Depot)

AND

3 - Used in combination with somatostatin analog therapy (e.g., octreotide, Sandostatin LAR, Somatuline Depot)

| Product Name: Xermelo | |
|-----------------------|-----------------------------|
| Diagnosis | Carcinoid Syndrome Diarrhea |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Documentation of positive clinical response to Xermelo

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Xofluza (baloxavir)

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-114475 Xofluza (baloxavir)

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Xofluza | | |
|--|---------------------|--|
| Approval Length | 1 month(s) | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Patient is 5 years of age or older | | |
| | | |
| AND | | |
| | | |

| 2 - One of the following: |
|---|
| Patient has acute, uncomplicated influenza Used for post-exposure prophylaxis of influenza |
| AND |
| 3 - Patient has not been symptomatic for more than 48 hours |
| AND |
| 4 - Patient does NOT meet all of the following: |
| On concurrent neuraminidase inhibitors (e.g., Tamiflu, Relenza) Pregnant Hospitalized |
| AND |
| 5 - Documentation of reason why preferred generic oseltamivir is not clinically appropriate for the patient (i.e., the convenience of the patient, prescriber, or other health care provider should not be accepted) |

2. Revision History

Г

| Date | Notes |
|-----------|-------------|
| 9/26/2022 | New program |

Xolair

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99659 Xolair

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Xolair | |
|--|-----------------------|
| Diagnosis | Asthma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of moderate or severe asthma | |

AND

2 - Classification of asthma as uncontrolled or inadequately controlled as defined by ONE of the following:

2.1 Poor symptom control (e.g., Asthma Control Questionnaire [ACQ] score consistently greater than 1.5 or Asthma Control Test [ACT] score consistently less than 20)

OR

2.2 Two or more bursts of systemic corticosteroids for at least 3 days each in the previous 12 months

OR

2.3 Asthma-related emergency treatment (e.g., emergency room visit, hospital admission, or unscheduled physician's office visit for nebulizer or other urgent treatment)

OR

2.4 Airflow limitation (e.g., after appropriate bronchodilator withhold forced expiratory volume in 1 second [FEV1] less than 80 percent predicted [in the face of reduced FEV1-forced vital capacity [FVC] defined as less than the lower limit of normal])

OR

2.5 Patient is currently dependent on oral corticosteroids for the treatment of asthma

AND

3 - ONE of the following:

3.1 Baseline (pre-omalizumab treatment) serum total Immunoglobulin E (IgE) level greater than or equal to 30 IU/mL (international units per milliliter) and less than or equal to 1500 IU/mL

OR

3.2 Patient is currently dependent on oral corticosteroids for the treatment of asthma

AND

4 - Positive skin test or in vitro reactivity to a perennial aeroallergen

AND

5 - Used in combination with ONE of the following:

5.1 One maximally-dosed (appropriately adjusted for age) combination inhaled corticosteroid (ICS)-long-acting beta2-agonist (LABA) product [e.g., fluticasone propionate-salmeterol (AirDuo, Advair), budesonide-formoterol (Symbicort)]

OR

5.2 Combination therapy including BOTH of the following:

5.2.1 One high-dose (appropriately adjusted for age) inhaled corticosteroid (ICS) product [e.g., ciclesonide (Alvesco), mometasone furoate (Asmanex), beclomethasone dipropionate (QVAR)]

AND

5.2.2 One additional asthma controller medication [e.g., LABA - olodaterol (Striverdi) or indacaterol (Arcapta); leukotriene receptor antagonist – montelukast (Singulair); theophylline]

AND

6 - Patient is not receiving Xolair in combination with ONE of the following:

• Anti-interleukin 4 therapy [e.g. Dupixent (dupilumab)]

• Anti-interleukin 5 therapy [e.g. Nucala (mepolizumab), Cinqair (reslizumab), Fasenra (benralizumab)]

AND

7 - Xolair dosing for moderate to severe persistent asthma is in accordance with the United States Food and Drug Administration approved labeling

AND

8 - Prescribed by or in consultation with an allergist-immunologist or pulmonologist

| Product Name: Xolair | |
|----------------------|---------------------|
| Diagnosis | Asthma |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- 1 Documentation of positive clinical response as demonstrated by ONE of the following:
 - reduction in frequency of exacerbations
 - decreased utilization of rescue medications
 - increase in percent predicted forced expiratory volume in 1 second (FEV1) from pretreatment baseline
 - reduction in severity or frequency of asthma-related symptoms (e.g., wheezing, shortness of breath, coughing)

AND

2 - Used in combination with an inhaled corticosteroid (ICS)-containing controller medication

AND

3 - Patient is not receiving Xolair in combination with ONE of the following:

- Anti-interleukin 4 therapy [e.g. Dupixent (dupilumab)]
- Anti-interleukin 5 therapy [e.g. Nucala (mepolizumab), Cinqair (reslizumab), Fasenra (benralizumab)]

AND

4 - Xolair dosing for moderate to severe persistent asthma is in accordance with the United States Food and Drug Administration approved labeling

AND

5 - Prescribed by or in consultation with allergist-immunologist or pulmonologist

| Product Name: Xolair | |
|----------------------|------------------------------|
| Diagnosis | Chronic Idiopathic Urticaria |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of chronic idiopathic urticaria

AND

2 - ONE of the following:

2.1 Patient remains symptomatic despite at least a 2-week trial of, or history of contraindication or intolerance to, two H1-antihistamines [e.g., Allegra (fexofenadine), Benadryl (diphenhydramine), Claritin (loratadine)]*

OR

2.2 Patient remains symptomatic despite at least a 2-week trial of, or history of contraindication or intolerance to BOTH of the following taken in combination:

2.2.1 Second generation H1-antihistamine [e.g., Allegra (fexofenadine), Claritin (loratadine), Zyrtec (cetirizine)] AND 2.2.2 ONE of the following: • Different second generation H1-antihistamine [e.g., Allegra (fexofenadine), Claritin (loratadine), Zyrtec (cetirizine)] First generation H1-antihistamine [e.g., Benadryl (diphenhydramine), Chlor-Trimeton • (chlorpheniramine), Vistaril (hydroxyzine)]* H2-antihistamine [e.g., Pepcid (famotidine), Tagamet HB (cimetidine), Zantac • (ranitidine)] Leukotriene modifier [e.g., Singulair (montelukast)] • AND **3** - Xolair dosing for chronic idiopathic urticaria is in accordance with the United States Food and Drug Administration approved labeling AND 4 - Prescribed by or in consultation with an allergist-immunologist or dermatologist *Patients 65 years of age and older in whom first generation H1-antihi Notes stamines are considered high risk medications to be avoided (e.g., Be ers criteria, HEDIS) should be directed to try alternatives that are not c

| Product Name: Xolai | r |
|---------------------|------------------------------|
| Diagnosis | Chronic Idiopathic Urticaria |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

onsidered high risk.

1 - Documentation of positive clinical response (e.g., reduction in exacerbations, itch severity, hives)

AND

2 - Xolair dosing for chronic idiopathic urticaria is in accordance with the United States Food and Drug Administration approved labeling

AND

3 - Prescribed by or in consultation with allergist-immunologist or dermatologist

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Xopenex Respules

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99502 Xopenex Respules

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Brand Xopenex inhalation soln, generic levalbuterol inhalation soln | |
|---|--------------|
| Approval Length | 12 month(s) |
| Guideline Type | Step Therapy |
| | |

Approval Criteria

1 - The patient has a history of failure, contraindication, or intolerance to treatment with albuterol inhalation solution

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona Standard to Medicaid Arizona Standard for 7 /1 go live |

Xuriden

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99660 Xuriden

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Xuriden | |
|---|-----------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of a hereditary orotic aciduria | |

Product Name: Xuriden

| Approval Length | 12 month(s) |
|-----------------|---------------------|
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

1 - Documentation of positive clinical response to Xuriden therapy

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Medicaid SP to Medicaid Arizona SP for 7/1 |

Xyrem, Xywav

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99814 Xyrem, Xywav

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Xyrem, Xywav | |
|----------------------------|---|
| Diagnosis | Narcolepsy with Cataplexy (i.e., Narcolepsy Type 1) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g. chart notes, laboratory values) documenting a diagnosis of narcolepsy with cataplexy (i.e., Narcolepsy Type 1) with BOTH of the following:

1.1 The patient has daily periods of irrepressible need to sleep or daytime lapses into sleep occurring for at least three months

AND

1.2 A mean sleep latency of less than or equal to 8 minutes and two or more sleep onset rapid eye movement (REM) periods (SOREMPs) on a Multiple Sleep Latency Test (MSLT) performed according to standard techniques following a normal overnight polysomnogram. A SOREMP (within 15 minutes of sleep onset) on the preceding nocturnal polysomnogram may replace one of the SOREMPs on the MSLT

AND

2 - Physician attestation to BOTH of the following:

2.1 Patient has experienced cataplexy defined as more than one episode of sudden loss of muscle tone with retained consciousness

AND

2.2 Other causes of sleepiness have been ruled out or treated (including but not limited to obstructive sleep apnea, insufficient sleep syndrome, shift work, the effects of substances or medications, or other sleep disorders)

AND

3 - Prescribed by ONE of the following:

- Neurologist
- Psychiatrist
- Sleep Medicine Specialist

| Product Name: Xyrem, Xywav | |
|----------------------------|---|
| Diagnosis | Narcolepsy with Cataplexy (i.e., Narcolepsy Type 1) |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |

| Guideline Type | Prior Authorization |
|----------------|---------------------|
|----------------|---------------------|

1 - Documentation demonstrating a reduction in frequency of cataplexy attacks associated with therapy

OR

2 - Documentation demonstrating reduction in symptoms of excessive daytime sleepiness associated with therapy

| Product Name: Xyrem, Xywav | |
|----------------------------|--|
| Diagnosis | Narcolepsy without Cataplexy (i.e., Narcolepsy Type 2) |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Submission of medical records (e.g. chart notes, lab values) documenting a diagnosis of narcolepsy without cataplexy (i.e., Narcolepsy Type 2) with BOTH of the following:

1.1 The patient has daily periods of irrepressible need to sleep or daytime lapses into sleep occurring for at least three months

AND

1.2 A mean sleep latency of less than or equal to 8 minutes and two or more sleep onset rapid eye movement (REM) periods (SOREMPs) are found on a Multiple Sleep Latency Test (MSLT) performed according to standard techniques following a normal overnight polysomnogram. A SOREMP (within 15 minutes of sleep onset) on the preceding nocturnal polysomnogram may replace one of the SOREMPs on the MSLT

AND

2 - Physician attestation to BOTH of the following:

2.1 Cataplexy is absent

AND

2.2 Other causes of sleepiness have been ruled out or treated (including but not limited to obstructive sleep apnea, insufficient sleep syndrome, shift work, the effects of substances or medications or their withdrawal, sleep phase disorder, or other sleep disorders)

AND

3 - History of failure, contraindication, or intolerance of ALL of the following (MUST be verified via paid pharmacy claims or submission of medical records):

3.1 ONE of the following:

- Amphetamine based stimulant (e.g., amphetamine, dextroamphetamine)
- Methylphenidate based stimulant

AND

3.2 Armodafanil (Nuvigil)

AND

3.3 Sunosi (solriamfetol)

AND

- 4 Prescribed by ONE of the following:
 - Neurologist
 - Psychiatrist
 - Sleep Medicine Specialist

Product Name: Xyrem, Xywav

| Narcolepsy without Cataplexy (i.e., Narcolepsy Type 2) |
|--|
| 12 month(s) |
| Reauthorization |
| Prior Authorization |
| |

1 - Documentation demonstrating reduction in symptoms of excessive daytime sleepiness associated with therapy

| Date | Notes |
|------------|---|
| 11/23/2021 | Added 'verified via paid pharmacy claims of submission of medical re cords' to t/f requirements |

Yonsa

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99734 Yonsa

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Yonsa | |
|----------------------------------|-----------------------|
| Diagnosis | Prostate Cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of prostate cancer | |

AND

2 - ONE of the following:

2.1 Disease is metastatic

OR

2.2 Disease is regional node positive (e.g., N1)

AND

3 - Used in combination with methylprednisolone

AND

4 - ONE of the following:

4.1 Used in combination with a gonadotropin-releasing hormone (GnRH) analog [e.g., Lupron (leuprolide), Zoladex (goserelin), Trelstar (triptorelin), Vantas (histrelin), Firmagon (degarelix)]

OR

4.2 Patient has had bilateral orchiectomy

AND

5 - ONE of the following:

5.1 Prescriber provides a reason or special circumstance the patient cannot take Zytiga

5.2 Patient is currently on Yonsa therapy

| Product Name: Yonsa | |
|---------------------|---------------------|
| Diagnosis | Prostate Cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Yonsa therapy

| Product Name: Yonsa | |
|---------------------|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Yonsa will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B

| Product Name: Yonsa | |
|---------------------------|--|
| NCCN Recommended Regimens | |
| 12 month(s) | |
| Reauthorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Yonsa therapy

| Date | Notes |
|-----------|--------------------|
| 5/18/2021 | 7/1 Implementation |

Zelboraf

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99698 Zelboraf

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Zelboraf | |
|------------------------|-----------------------|
| Diagnosis | Melanoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - One of the following diagnoses:

• Unresectable melanoma

• Metastatic melanoma

AND

2 - Patient is positive for BRAF V600 mutation

| Product Name: Zelboraf | |
|------------------------|--|
| Melanoma | |
| 12 month(s) | |
| Reauthorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Zelboraf therapy

| Product Name: Zelboraf | |
|------------------------|--------------------------------------|
| Diagnosis | Central Nervous System (CNS) Cancers |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient has metastatic brain lesions

AND

2 - Zelboraf is active against primary tumor (melanoma)

AND

3 - Used in combination with Cotellic (cobimetinib)

| Product Name: Zelboraf | |
|------------------------|--------------------------------------|
| Diagnosis | Central Nervous System (CNS) Cancers |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Patient does not show evidence of progressive disease while on Zelboraf therapy

| Product Name: Zelboraf | |
|------------------------|-----------------------|
| Diagnosis | Hairy Cell Leukemia |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of hairy cell leukemia

| Product Name: Zelboraf | |
|------------------------|---------------------|
| Diagnosis | Hairy Cell Leukemia |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Zelboraf therapy

| Product Name: Zelboraf | |
|------------------------|----------------------------|
| Diagnosis | Non-Small Cell Lung Cancer |

| Approval Length | 12 month(s) | |
|---|-----------------------|--|
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | Approval Criteria | |
| 1 - Diagnosis of non-small cell lung cancer (NSCLC) | | |
| | AND | |
| 2 - Disease is ONE of the following: | | |
| MetastaticAdvancedRecurrent | | |
| | AND | |
| 3 - Cancer is positive for BRAF V600E mutation | | |

| Product Name: Zelboraf | |
|----------------------------|--|
| Non-Small Cell Lung Cancer | |
| 12 month(s) | |
| Reauthorization | |
| Prior Authorization | |
| | |

1 - Patient does not show evidence of progressive disease while on Zelboraf therapy

| Product Name: Zelboraf | |
|------------------------|-------------------------|
| Diagnosis | Erdheim-Chester Disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |

| Guideline Type | Prior Authorization |
|--|--------------------------|
| | |
| Approval Criteria | |
| 1 - Diagnosis of Erdheim-Chester Disease | |
| | AND |
| 2 - Cancer is positiv | e for BRAF V600 mutation |

| Product Name: Zelboraf | |
|------------------------|-------------------------|
| Diagnosis | Erdheim-Chester Disease |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

1 - Patient does not show evidence of progressive disease while on Zelboraf therapy

| Product Name: Zelboraf | |
|------------------------|-----------------------|
| Diagnosis | Colon Cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of colon cancer

AND

2 - Cancer is positive for BRAF V600E mutation

AND

3 - ONE of the following:

- Unresectable or advanced disease •
- Metastatic disease •

AND

4 - BOTH of the following:

4.1 Used in combination with irinotecan

AND

4.2 Used in combination with ONE of the following:

- •
- Erbitux (cetuximab) Vectibix (panitumumab) •

| Product Name: Zelboraf | |
|------------------------|---------------------|
| Diagnosis | Colon Cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Zelboraf therapy

| Product Name: Zelboraf | |
|------------------------|---------------|
| Diagnosis | Rectal Cancer |
| Approval Length | 12 month(s) |

| Therapy Stage | Initial Authorization | |
|---|--------------------------------|--|
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of rectal cancer | | |
| | AND | |
| 2 - Cancer is positive for | or BRAF V600E mutation | |
| | AND | |
| 3 - ONE of the following | g: | |
| Unresectable or advanced diseaseMetastatic disease | | |
| | AND | |
| 4 - BOTH of the followi | ng: | |
| 4.1 Used in combination with irinotecan | | |
| | AND | |
| 4.2 Used in combinati | ion with ONE of the following: | |
| Erbitux (cetuximab)Vectibix (panitumumab) | | |

| Product Name: Zelboraf | |
|------------------------|-----------------|
| Diagnosis | Rectal Cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |

| Guideline Type | Prior Authorization |
|----------------|---------------------|
| | |

1 - Patient does not show evidence of progressive disease while on Zelboraf therapy

| Product Name: Zelboraf | |
|------------------------|-----------------------|
| Diagnosis | Thyroid Cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

- 1 Diagnosis of ONE of the following:
 - Follicular carcinoma •
 - Hurthle cell carcinoma
 - Papillary carcinoma

AND

2 - ONE of the following:

- Unresectable locoregional recurrent disease Metastatic disease ٠
- •
- Persistent disease •

AND

3 - ONE of the following:

- Patient has symptomatic disease ٠
- Patient has progressive disease •

AND

4 - Disease is refractory to radioactive iodine

AND

5 - Cancer is positive for BRAF V600 mutation

| Product Name: Zelboraf | |
|------------------------|---------------------|
| Diagnosis | Thyroid Cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Zelboraf therapy

| Product Name: Zelboraf | |
|---------------------------|--|
| NCCN Recommended Regimens | |
| 12 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Zelboraf will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Zelboraf | |
|------------------------|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Documentation of positive clinical response to Zelboraf therapy

| Date | Notes |
|----------|--------------------|
| 4/8/2021 | 7/1 Implementation |

Zeposia (ozanimod)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: |
|-----------------|
|-----------------|

1. Criteria

| Product Name: Zeposia | |
|--|-----------------------|
| Diagnosis | Multiple Sclerosis |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Diagnosis of multiple sclerosis (MS) | |

AND

2 - Patient has a history of failure, contraindication, or intolerance to a trial of at least TWO of the preferred alternatives * (May require PA) (Verified via pharmacy paid claims or submission of medical records)

- Interferon Beta-1B (Extavia)
- Fingolimod (Gilenya)
- Brand Copaxone 20mg
- Brand Glatopa 40mg
- Interferon Beta-1A (Refib, Avonex)

| Notes *Note: Preferred alternatives may require PA | |
|--|--|
|--|--|

| Product Name: Zeposia | |
|--|---------------------|
| Diagnosis | Multiple Sclerosis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Documentation of positive clinical response to therapy | |

| Product Name: Zeposia | |
|-----------------------|-----------------------|
| Diagnosis | Ulcerative Colitis |
| Approval Length | 12 Weeks [B, 3, 4] |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Diagnosis of moderately to severely active ulcerative colitis

AND

2 - History of failure to one of the following conventional therapies at maximally indicated doses within the last 3 months, unless contraindicated or clinically significant adverse effects are experienced (document drug, date, and duration of trial)*:

- 6-mercaptopurine (Purinethol)
- Aminosalicylates (e.g., mesalamine [Asacol, Pentasa, Rowasa], osalazine [Dipentum], Sulfasalazine [Azulfidine, Sulfazine])
- Azathioprine (Imuran)
- Corticosteroids (e.g., prednisone, methylprednisolone)

AND

3 - One of the following:

3.1 Trial and failure, contraindication, or intolerance to TWO of the following, or attestation demonstrating a trial may be inappropriate*:

- Humira (adalimumab)
- Simponi (golimumab)
- Stelara (ustekinumab)

OR

3.2 For continuation of prior therapy

AND

4 - Prescribed by or in consultation with a gastroenterologist

AND

5 - Patient is NOT receiving Zeposia in combination with ANY of the following:

- Biologic disease modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]

| Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)] | |
|---|--|
| Notes | *Note: Claims history may be used in conjunction as documentation of drug, date, and duration of trial |

| Product Name: Zeposia | |
|-----------------------|---------------------|
| Diagnosis | Ulcerative Colitis |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Documentation of positive clinical response to therapy

AND

2 - Patient is NOT receiving Zeposia in combination with ANY of the following:

- Biologic disease modifying anti-rheumatic drug (DMARD) [e.g., Enbrel (etanercept), Humira (adalimumab), Cimzia (certolizumab)]
- Janus kinase inhibitor [e.g., Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [e.g. Otezla (apremilast)]

| Date | Notes |
|----------|--|
| 2/3/2022 | New drug-specific guideline for Zeposia, with new criteria added for UC indication |

Zimhi (naloxone)

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-114472 Zimhi (naloxone)

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Zimhi | |
|---------------------|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| | |

1 - History of failure, or intolerance to preferred naloxone products (e.g., Brand Narcan nasal spray, Kloxxado, preferred naloxone injections)

| Date | Notes |
|-----------|-------------|
| 9/26/2022 | New program |

Zolgensma (onasemnogene abeparvovec-xioi)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

| GL-99786 | Zolgensma | (onasemnogene | abeparvovec-xioi) |
|----------|-----------|---------------|-------------------|
| | | | |

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Zolgensma | |
|-------------------------------|--|
| ime Authorization in Lifetime | |
| or Authorization | |
| | |

Approval Criteria

1 - The mutation or deletion of genes in chromosome 5q resulting in one of the following: [1-8, A]

1.1 Homozygous gene deletion or mutation of SMN1 gene (e.g., homozygous deletion of exon 7 at locus 5q13)

OR

1.2 Compound heterozygous mutation of SMN1 gene (e.g., deletion of Survival of Motor Neuron 1 [SMN1] exon 7 [allele 1] and mutation of SMN1 [allele 2])

AND

2 - One of the following:

2.1 Both of the following: [1-5]

2.1.1 Diagnosis of diagnosis of SMA Type 0, I or Type II spinal muscular atrophy (SMA) confirmed by a neurologist with expertise in the treatment of SMA

AND

2.1.2 Patient is less than or equal to 2 years of age

OR

2.2 Both of the following:

2.2.1 Diagnosis of SMA based on the results of SMA newborn screening

AND

2.2.2 Patient has 3 copies or less of Survival of Motor Neuron 2 (SMN 2)

AND

3 - Patient is not dependent on either of the following:

- Invasive ventilation or tracheostomy
- Use of invasive ventilation beyond use of naps and nighttime sleep

AND

4 - Submission of medical records (e.g., chart notes, laboratory values) documenting patient's anti-AAV9 antibody titers are less than or equal to 1:50 [1]

AND

5 - Patient is not to receive concomitant SMN modifying therapy (e.g. Spinraza)

AND

6 - Prescribed by a neurologist with expertise in the diagnosis of SMA

AND

7 - Patient has never received Zolgensma treatment in their lifetime

2. Endnotes

- A. This is the definition that the clinical trials used. Also consistent with clinical guidelines. [2-8]
- B. There were 3 key clinical trials for Zolgensma (START, STR1VE, SPR1NT). START and STR1VE only enrolled patients with SMA Type 1 and SPR1NT enrolled pre-symptomatic SMA patients. [2-5]
- C. Exclusion criteria found in clinical trials. [2-5]
- D. A recent European ad-hoc consensus statement on SMA stated that there currently is no published evidence that the combination of two disease modifying therapies (e.g., Spinraza and Zolgensma) is superior to any single treatment alone. RESPOND is a phase 4 trial that will assess the efficacy and safety of Spinraza in patients with suboptimal clinical response to Zolgensma. It is planned to begin enrollment in 2021. [9-10]

3. References

1. Zolgensma Prescribing Information. AveXis Inc. Bannockburn, IL. May 2019.

- 2. Mendell J.R., Al-Zaidy S, Shell R, etc. Single-Dose Gene Replacement Therapy for Spinal Muscular Atrophy. New Eng J of Med. 2017; 377:1713-22.
- Al-Zaidy S, Pickard AS, Kotha K, et al. Health outcomes in spinal muscular atrophy type 1 following AVXS-101 gene replacement therapy. Pediatr Pulmonol. 2019;54(2):179-185.
- Day JW, Chiriboga CA, Crawford TO, et al. AVXS-101 gene-replacement therapy for spinal muscular atrophy type 1: phase 3 study (STR1VE) update. Poster presented at: The 71st Annual American Academy of Neurology Meeting, Philadelphia PA, May 4-10, 2019.
- Strauss KA, Swoboda KJ, Farrar MA, et al. AVXS-101 gene-replacement therapy in presymptomatic spinal muscular atrophy: SPR1NT study update. Poster presented at the 71st Annual American Academy of Neurology Meeting; May 4-10; 2019; Philadelphia, PA.
- 6. Markowitz JA, Sing P, Darras BT. Spinal muscular atrophy: a clinical and research update. Pediatr Neurol. 2012;46(1):1-12.
- 7. Wang CH, Finkel RS, Bertini ES, et al. Consensus statement for standard of care in spinal muscular atrophy. J Child Neurol. 2007;22(8):1027-1049.
- 8. Mercuri E, Finkel RS, Muntoni F, et al. Diagnosis and management of spinal muscular atrophy: Part 1: Recommendations for diagnosis, rehabilitation, orthopedic and nutritional care. J Neuromuscul Dis. 2018;28(2):103-115.
- Kirschner J, Butoianu N, Goemans N, et al. European ad-hoc consensus statement on gene replacement therapy for spinal muscular atrophy. Eur J Paediatr Neurol. 2020. https://doi.org/10.1016/j.ejpn.2020.07.001.
- Biogen. Biogen plans to initiate phase 4 study evaluating benefit of Spinraza® (nusinersen) in patients treated with Zolgensma® (onasemnogene abeparvovec). https://investors.biogen.com/news-releases/news-release-details/biogen-plans-initiatephase-4-study-evaluating-benefit-spinrazar. July 21, 2020. Accessed October 6, 2020.

| Date | Notes |
|-----------|--------------------|
| 6/28/2021 | 7/1 Implementation |

Zolinza

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99699 Zolinza

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Zolir | Product Name: Zolinza | |
|-----------------------|------------------------|--|
| Diagnosis | T-Cell Lymphoma | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| | | |
| Approval Criteria | | |
| 1 - Diagnosis of cuta | aneous T-cell lymphoma | |

| Product Name: Zolinza | |
|-----------------------|---------------------|
| Diagnosis | T-Cell Lymphoma |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |

1 - Patient does not show evidence of progressive disease while on Zolinza therapy

| Product Name: Zolinza | duct Name: Zolinza | |
|-----------------------|---------------------------|--|
| Diagnosis | NCCN Recommended Regimens | |
| Approval Length | 12 month(s) | |
| Therapy Stage | Initial Authorization | |
| Guideline Type | Prior Authorization | |
| Guideline Type | Phor Authonization | |

Approval Criteria

1 - Use is supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Zolinza | |
|---------------------------|--|
| NCCN Recommended Regimens | |
| 12 month(s) | |
| Reauthorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Documentation of positive clinical response to Zolinza therapy

| Date | Notes |
|----------|--------------------|
| 4/8/2021 | 7/1 Implementation |

Zontivity

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99503 Zontivity

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: |
|-----------------|
|-----------------|

1. Criteria

| Product Name: Zontivity | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - ONE of the following: | |
| History of myocardial infarction (MI) Peripheral arterial disease (PAD) | |

AND

2 - Patient does not have a history of ONE of the following:

- Stroke •
- Transient ischemic attack (TIA) Intracranial hemorrhage (ICH) •
- •

AND

3 - Patient does not have active pathological bleeding

| Date | Notes |
|-----------|---|
| 3/11/2021 | Bulk Copy C&S Arizona Standard to Medicaid Arizona Standard for 7 /1 go live |

Zortress - ARIZONA

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99504 Zortress - ARIZONA

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Zortress | |
|--|---------------------|
| Approval Length | 12 month(s) |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - Kidney transplant rejection prophylaxis in patients at low-moderate immunologic risk | |
| | |
| OR | |
| | |

2 - Liver transplant rejection prophylaxis

| Date | Notes |
|-----------|--|
| 3/11/2021 | Bulk Copy C&S Arizona Standard to Medicaid Arizona Standard for 7 /1 go live |

Ztalmy (ganaxolone)

Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-114155 Ztalmy (ganaxolone)

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 10/1/2022 |
|-----------------|-----------|
|-----------------|-----------|

1. Criteria

| Product Name: Ztalmy | |
|-----------------------|--|
| 6 month(s) | |
| Initial Authorization | |
| Prior Authorization | |
| | |

Approval Criteria

1 - Submission of documentation (e.g., chart notes) confirming diagnosis of cyclin-dependent kinase-like 5 (CDKL5) deficiency disorder (CDD)

| AND |
|---|
| 2 - Patient has a mutation in the CDKL5 gene |
| AND |
| 3 - Patient is 2 years of age or older |
| AND |
| 4 - Patient is experiencing motor seizures (e.g., bilateral tonic, generalized tonic-clonic, bilateral clonic, atonic, focal, or bilateral tonic-clonic) |
| AND |
| 5 - One of the following: |
| 5.1 Trial and failure, contraindication, or intolerance to two preferred anticonvulsants (e.g., valproic acid, levetiracetam, lamotrigine) |
| OR |
| 5.2 For continuation of prior therapy |
| AND |
| 6 - Prescribed by or in consultation with a neurologist |

| Product Name: Ztalmy | |
|----------------------|---------------------|
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

1 - Documentation of positive clinical response to therapy as evidenced by a reduction in the frequency of seizures from baseline

| Date | Notes |
|-----------|-------------|
| 9/20/2022 | New Program |

Zytiga Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99800 Zytiga

Formulary Medicaid - Arizona SP (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: |
|-----------------|
|-----------------|

1. Criteria

| Product Name: Brand Zytiga, generic abiraterone | |
|---|-----------------------|
| Diagnosis | Prostate Cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Diagnosis of prostate cancer

AND

2 - ONE of the following:

2.1 Disease is metastatic

OR

2.2 Disease is regional node positive (e.g., N1)

AND

3 - Used in combination with prednisone

AND

4 - ONE of the following:

4.1 Used in combination with a gonadotropin-releasing hormone (GnRH) analog [e.g., Lupron (leuprolide), Zoladex (goserelin), Trelstar (triptorelin), Vantas (histrelin), Firmagon (degarelix)]

OR

4.2 Patient has had bilateral orchiectomy

AND

5 - If the request is for brand Zytiga, must meet both of the following:

5.1 Patient has tried and failed generic Zytiga

AND

5.2 The prescriber provides reason or special circumstance why the patient cannot take generic Zytiga

| Product Name: Brand Zytiga, generic abiraterone | |
|---|---------------------|
| Diagnosis | Prostate Cancer |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Patient does not show evidence of progressive disease while on Zytiga therapy

| Product Name: Brand Zytiga, generic abiraterone | |
|---|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Initial Authorization |
| Guideline Type | Prior Authorization |

Approval Criteria

1 - Zytiga will be approved for uses supported by The National Comprehensive Cancer Network (NCCN) Drugs and Biologics Compendium with a Category of Evidence and Consensus of 1, 2A, or 2B.

| Product Name: Brand Zytiga, generic abiraterone | |
|---|---------------------------|
| Diagnosis | NCCN Recommended Regimens |
| Approval Length | 12 month(s) |
| Therapy Stage | Reauthorization |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

1 - Documentation of positive clinical response to Zytiga therapy

| Date | Notes |
|-----------|---|
| 6/28/2021 | Updated criteria added ABIRATERONE ACETATE TAB 500 MG |

Zyvox

Medicaid - Arizona (AZM, AZMREF, AZMDDD)



Prior Authorization Guideline

GL-99578 Zyvox

Formulary Medicaid - Arizona (AZM, AZMREF, AZMDDD)

Formulary Note

Guideline Note:

| Effective Date: | 12/9/2021 |
|--------------------|-----------|
| P&T Approval Date: | |
| P&T Revision Date: | |

1. Criteria

| Product Name: Brand Zyvox*, generic linezolid* | |
|--|---------------------|
| Diagnosis | Labeled Uses |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |
| 1 - One of the following: | |
| 1.1 For continuation of therapy upon hospital discharge | |

OR

1.2 As continuation of therapy when transitioning from intravenous antibiotics that are shown to be sensitive to the cultured organism for the requested indication

OR

1.3 BOTH of the following:

1.3.1 ONE of the following diagnoses:

- Nosocomial pneumonia
- Community-acquired pneumonia
- Skin and skin structure infections (complicated and uncomplicated)

AND

1.3.2 Infection caused by an organism that is confirmed to be or likely to be susceptible to treatment with Zyvox

OR

1.4 Invasive infection caused by or likely to be caused by vancomycin-resistant Enterococcus faecium (VRE)

| Notes | *Approval Duration: For vancomycin-resistant Enterococcus faecium, |
|-------|---|
| | authorization will be issued for 28 days. For osteomyelitis, authorizatio |
| | n will be issued for the requested duration, not to exceed 6 weeks. All |
| | other approvals will be issued for 14 days. |

| Product Name: Brand Zyvox*, generic linezolid* | |
|--|---------------------|
| Diagnosis | Off label Uses |
| Guideline Type | Prior Authorization |
| | |
| Approval Criteria | |

| 1 - For continuation of therapy upon hospital discharge | |
|--|---|
| | OR |
| 2 - As continuation of therapy when transitioning from intravenous antibiotics that are shown to be sensitive to the cultured organism for the requested indication | |
| OR | |
| 3 - The medication is being prescribed by or in consultation with an Infectious Disease specialist | |
| Notes | *Approval Duration: Based on provider recommended treatment durati ons, not to exceed 6 months. |

| Date | Notes |
|-----------|------------------|
| 7/21/2021 | Update guideline |