



# Nebraska Medicaid Preferred Drug List with Prior Authorization Criteria

July 2026 PDL

PDL updated July 1, 2026, **Highlights** indicate change from previous posting

For the most up to date list of covered drugs consult the **Drug Lookup** on the Nebraska Medicaid website at <https://ne.primetherapeutics.com/drug-lookup>.

- **PDMP Check Requirements** – Nebraska Medicaid providers are required to check the prescription drug history in the statewide PDMP before prescribing CII controlled substances to certain Medicaid beneficiaries (exemption to this requirement are for beneficiaries receiving cancer treatment, hospice/palliative care, or in long-term care facilities). If not able to check the PDMP, then provider is required to document good faith effort, including reasons why unable to conduct the check and may be required to submit documentation to the State upon request.
  - PDMP check requirements are under Section 5042 of the SUPPORT for Patients and Communities Act, consistent with section 1944 of the Social Security Act [42 U.S.C. 1396w-3a].
- **Opioids** – The maximum opioid dose covered is 90 Morphine Milligram Equivalents (MME) per day.

## Non-Preferred Drug Coverage

Class and drug-specific therapeutic trial and failure requirements are found within this document. Examples of non-preferred exception criteria include:

- Adverse reaction to preferred drugs
- Allergy to preferred drugs
- Contraindication to preferred drugs
- Documentation of inability to swallow solid dosage forms

Specific Class Prior Authorization forms can be found within the PDL class listings and at:

<https://nebraska.fhsc.com/priorauth/paforms.asp>

- [Immunomodulators Self-Injectable PA Form](#)
- [Opioid Dependence Treatment PA Form](#)
- [Opioid Dependence Treatment Informed Consent](#)
- [Growth Hormone PA Form](#)
- [HAE Treatments PA Form](#)
- [Hepatitis C PA Form](#)

For all other class medically-necessary coverage, quantity, and high dose requests use the following:

[Documentation of Medical Necessity PA Form](#)

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**ACNE AGENTS, TOPICAL**

| Preferred Agents   | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|--|--|--|
| adapalene (generic Differin) <b>GEL (OTC/Rx), GEL PUMP</b><br>adapalene/BPO (generic Epiduo)<br>benzoyl peroxide (BPO) <b>WASH, LOTION</b><br>benzoyl peroxide <b>GEL OTC</b><br>clindamycin/BPO (generic BenzaClin) <b>GEL, PUMP</b><br>clindamycin/BPO (generic Duac)<br>clindamycin phosphate <b>PLEDGET</b><br>clindamycin phosphate <b>SOLUTION</b><br>erythromycin <b>GEL</b><br>erythromycin <b>SOLN</b><br>erythromycin-BPO (generic Benzamycin)<br>tretinoin (generic Avita, Retin-A) <sup>AL</sup> , <b>CREAM, GEL</b> | adapalene (generic Differin) <b>CREAM</b><br>adapalene/BPO (generic Epiduo Forte)<br>ALTRENO (tretinoin) <sup>AL</sup><br>AMZEEQ (minocycline)<br>ATRALIN (tretinoin)<br>AVAR (sulfacetamide sodium/sulfur)<br>AVITA (tretinoin)<br>AZELEX (azelaic acid)<br>BENZEFOAM (benzoyl peroxide)<br>benzoyl peroxide <b>CLEANSER, CLEANSING BAR OTC</b><br>benzoyl peroxide <b>FOAM</b> (generic BenzePro)<br>benzoyl peroxide <b>GEL Rx</b><br>benzoyl peroxide <b>TOWELETTE OTC</b><br>clindamycin <b>FOAM, LOTION</b><br>clindamycin <b>GEL</b><br>clindamycin phosphate (generic Clindagel) <b>GEL</b><br>clindamycin/BPO (generic Acanya) <b>GEL</b><br>clindamycin/BPO <b>PUMP</b> (generic Onexton) <sup>AL</sup><br>clindamycin/tretinoin (generic Veltin, Ziana)<br>dapsone (generic Aczone)<br>DIFFERIN (adapalene) <b>CREAM, GEL-OTC, GEL PUMP, LOTION</b><br>erythromycin <b>PLEDGET</b><br>EVOCLIN (clindamycin) <b>FOAM</b> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed THREE preferred agents within this drug class</li> <li>▪ All retinoid products have a maximum age limit of 20 without a diagnosis of acne</li> </ul> |

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**ACNE AGENTS, TOPICAL (Continued)**

| Preferred Agents | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|------------------|--|--|
|                  | FABIOR (tazarotene) <b>FOAM</b><br>NEUAC (clindamycin/BPO)<br>OVACE PLUS (sulfacetamide sodium)<br>RETIN-A MICRO (tretinoin)<br>sulfacetamide<br>sulfacetamide/sulfur<br>sulfacetamide sodium/ sulfur<br><b>CLEANSER</b><br>SUMADAN (sulfacetamide/sulfur)<br>tazarotene (generic Tazorac) <b>CREAM</b><br>tazarotene <b>FOAM</b> (generic Fabior)<br>tazarotene <b>GEL</b> (generic Tazorac)<br>TRETIN-X (tretinoin)<br>tretinoin (generic Atralin) <sup>AL</sup> <b>GEL</b><br>tretinoin microspheres (generic Retin-A Micro) <sup>AL</sup> <b>GEL, GEL PUMP</b><br>WINLEVI (clascoterone) <sup>AL</sup> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed THREE preferred agents within this drug class</li> <li>▪ All retinoid products have a maximum age limit of 20 without a diagnosis of acne</li> </ul> |

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**ALZHEIMER’S AGENTS**

| Preferred Agents   | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|--|--|--|
| <b>CHOLINESTERASE INHIBITORS</b>   |  | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a 120-day trial of ONE preferred agent within the last 6 months;</li> <li><b>OR</b></li> <li>▪ Current, stabilized therapy of the non-preferred agent within the previous 45 days</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Donepezil 23:</b> Requires donepezil 10mg/day for at least 3 months AND clinical reason as to why 5mg or 10mg tablets can't be used (to deliver 20mg or 25mg)</li> </ul> |
| donepezil (generic Aricept)<br>donepezil ODT (generic Aricept ODT)<br>rivastigmine <b>PATCH</b> (generic Exelon Patch) | ADLARITY (donepezil) <b>PATCH</b><br>ARICEPT (donepezil)<br>donepezil 23 (generic Aricept 23) <sup>CL</sup><br>EXELON (rivastigmine) <b>PATCH</b><br>galantamine (generic Razadyne)<br><b>SOLN, TAB</b><br>galantamine ER (generic Razadyne ER)<br>rivastigmine <b>CAPS</b> (generic Exelon)<br>ZUNVEYL DR (benzgalantamine) |  |
| <b>NMDA RECEPTOR ANTAGONIST</b>  |  |  |
| memantine (generic Namenda)  | memantine ER (generic Namenda XR)<br>memantine <b>SOLN</b> (generic Namenda)<br>memantine/donepezil (generic Namzaric)<br>NAMENDA (memantine)<br>NAMZARIC (memantine/donepezil)  |  |

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**ANALGESICS, OPIOID LONG-ACTING**

| Preferred Agents   | Non-Preferred Agents   | Prior Authorization/Class Criteria  |
|--|--|---|
| BUTRANS (buprenorphine) <sup>QL</sup> <b>PATCH</b><br>fentanyl 25, 50, 75, 100 mcg <b>PATCH</b> <sup>QL</sup><br>morphine ER <b>TABLET</b> (generic MS Contin, Oramorph SR)<br>OXYCONTIN <sup>CL</sup> (oxycodone ER)<br>tramadol ER (generic Ultram ER) | BELBUCA (buprenorphine) <sup>AL,QL</sup><br><b>BUCCAL</b><br>buprenorphine PATCH (generic Butrans) <sup>QL</sup><br>fentanyl 37.5/62.5/87.5 mcg <b>PATCH</b> <sup>QL</sup><br>hydrocodone ER (generic Hysingla ER) <sup>QL</sup><br>hydrocodone bitartrate ER (generic Zohydro ER)<br>hydromorphone ER (generic Exalgo)<br>HYSINGLA ER (hydrocodone ER)<br>methadone <b>TABLET</b> <sup>CL</sup><br>methadone <b>ORAL SYR</b> <sup>CL</sup><br>methadone <b>SOL TABLET</b> <sup>CL</sup><br>morphine ER (generic Avinza, Kadian) <b>CAPS</b><br>NUCYNTA ER (tapentadol)<br>oxycodone ER (generic Oxycontin) <sup>CL</sup><br>oxymorphone ER (generic Opana ER)<br>tapentadol ER (Generic Nucynta ER) <sup>NR</sup><br>tramadol ER (generic ConZip) | The Center for Disease Control (CDC) does not recommend long-acting opioids when beginning opioid treatment. <ul style="list-style-type: none"> <li>▪ Preferred agents require previous use of a long-acting opioid or documentation of a trial on a short acting agent within 90 days</li> <li>▪ Non-preferred agents will be approved with failure on, or intolerance to TWO preferred agents within this drug class</li> </ul> Drug-specific criteria: <ul style="list-style-type: none"> <li>▪ <b>Methadone (all formulations):</b> Will only be approved for use in pain control or end of life care. Trial of preferred agent not required for end-of-life care</li> <li>▪ <b>Oxycontin/ oxycodone ER:</b> Pain contract required for maximum quantity authorization</li> </ul> |

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**ANALGESICS, OPIOID SHORT-ACTING<sup>QL</sup>**

| Preferred Agents   | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|--|--|--|
| <b>ORAL</b>  |  |  |
| acetaminophen/codeine <b>ELIXIR, TAB</b><br>codeine <b>TAB</b><br>hydrocodone/APAP <b>SOLN, TAB</b><br>hydrocodone/ibuprofen<br>hydromorphone <b>TAB</b><br>morphine <b>CONC SOLN, SOLN, TAB</b><br>oxycodone <b>TAB, SOLN</b><br>oxycodone/APAP<br>tramadol 50 <b>TAB<sup>AL</sup></b> (generic Ultram) | butalbital/caffeine/APAP/codeine<br>butalbital compound w/codeine<br>(butalbital/ASA/caffeine/codeine)<br>carisoprodol compound-codeine<br>(carisoprodol/ASA/codeine)<br>dihydrocodeine/APAP/caffeine<br>hydrocodone/APAP SOLN (generic<br>Zolvit) <sup>NR</sup><br>hydromorphone <b>LIQUID,</b><br><b>SUPPOSITORY</b> (generic Dilaudid)<br>levorphanol (generic Xyvona)<br>meperidine (generic Demerol)<br>morphine <b>SUPPOSITORIES</b><br>NALOCET (oxycodone/APAP)<br>NUCYNTA (tapentadol) <sup>CL</sup><br>oxycodone <b>CAPS</b><br>oxycodone/APAP <b>SOLN</b><br>oxycodone <b>CONCENTRATE</b><br>oxymorphone IR (generic Opana)<br>pentazocine/naloxone<br>ROXICODONE (oxycodone)<br>ROXYBOND (oxycodone)<br>SEGLENTIS (celecoxib/tramadol) <sup>AL</sup><br>tapentadol (generic Nucynta) <sup>CL, NR</sup><br>tramadol 25mg<br>tramadol 75mg<br>tramadol 100mg (generic Ultram) <sup>AL</sup><br>tramadol (generic Qdolo) <sup>AL</sup> <b>SOLN</b><br>tramadol/APAP (generic Ultracet) | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed THREE preferred agents within this drug class within the last 12 months.</li> <li>▪ Note: for short acting opiate tablets and capsules there is a maximum quantity limit of #150 per 30 days.</li> <li>▪ Opiate limits for opiate naïve patients will consist of:               <ul style="list-style-type: none"> <li>-prescriptions limited to a 7 day supply, AND</li> <li>-initial opiate prescription fill limited to maximum of 50 Morphine Milligram Equivalents (MME) per day</li> </ul>               These limits may only be exceeded with patient specific documentation of medical necessity, with examples such as, cancer diagnosis, end-of-life care, palliative care, Sickle Cell Anemia, or prescriber attestation that patient is not recently opiate naïve             </li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Nucynta/ tapentadol:</b> Approved only for diagnosis of acute pain, for 30 days or less</li> </ul> |

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**ANALGESICS, OPIOID SHORT-ACTING<sup>QL</sup> (Continued)**

| Preferred Agents                        | Non-Preferred Agents   | Prior Authorization/Class Criteria  |
|---|--|---|
| <b>NASAL</b>                            |  | Drug-specific criteria:<br><ul style="list-style-type: none"> <li>▪ <b>Actiq®/Fentora®/ fentanyl transmucosal/Onsolis:</b> Approved only for diagnosis of cancer AND current use of long-acting opiate</li> </ul> |
|   | butorphanol <b>SPRAY<sup>QL</sup></b>  |   |
| <b>BUCCAL/TRANSMUCOSAL<sup>CL</sup></b> |  |   |
|   | fentanyl <b>TRANSMUCOSAL</b> (generic Actiq) <sup>CL</sup><br>FENTORA (fentanyl) <sup>CL</sup> |   |

**ANDROGENIC AGENTS (TOPICAL)<sup>CL</sup>**

| Preferred Agents  | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|---|---|--|
| ANDROGEL (testosterone) <sup>CL</sup> <b>PUMP</b><br>testosterone <b>PUMP</b> (generic Androgel) <sup>CL</sup><br>TESTIM (testosterone)<br><b>TRANSDERMAL</b> | NATESTO (testosterone) <sup>CL</sup><br>testosterone <b>PACKET</b> (generic Androgel) <sup>CL</sup><br>testosterone <b>GEL, PACKET, PUMP</b> (generic Vogelxo)<br>testosterone (generic Axiron)<br>testosterone (generic Fortesta)<br>testosterone (generic Testim) | <ul style="list-style-type: none"> <li>▪ Preferred agents approved for diagnosis of Primary hypogonadism (congenital or acquired) or Hypogonadotropic hypogonadism. Off label use for the following will be considered with documentation of necessity: female to male transsexual – gender dysphoria, weight gain, male osteoporosis, delayed puberty in males, corticosteroid-induced hypogonadism and osteoporosis, inoperable carcinoma of the breast, postpartum breast pain and engorgement, and menopause</li> <li>▪ In addition, non-preferred agents will be approved for patients who have failed ONE preferred agent within this drug class within the last 6 months</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Androgel®:</b> Approved for Males only</li> <li>▪ <b>Natesto®:</b> Approved for Males only with diagnosis of:<br/>Primary hypogonadism (congenital or acquired) OR<br/>Hypogonadotropic hypogonadism (congenital or acquired)</li> </ul> |

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**ANGIOTENSIN MODULATORS**

| Preferred Agents  | Non-Preferred Agents   | Prior Authorization/Class Criteria  |
|---|--|---|
| <b>ACE INHIBITORS</b>   |  | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed TWO preferred agents within the last 12 months</li> <li>▪ Non-preferred combination products may be covered as individual prescriptions without prior authorization</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Epaned/enalapril oral solution/Qbrelis oral solution:</b> Clinical reason why oral tablet is not appropriate</li> </ul> |
| benazepril (generic Lotensin)<br>enalapril (generic Vasotec)<br>lisinopril (generic Prinivil, Zestril)<br>ramipril (generic Altace) | captopril (generic Capoten)<br>EPANED (enalapril) <sup>CL</sup> <b>ORAL SOLN</b><br>enalapril (generic Epaned) <sup>CL</sup> <b>ORAL SOLN</b><br>fosinopril (generic Monopril)<br>moexepiril (generic Univasc)<br>perindopril (generic Aceon)<br>QBRELIS (lisinopril) <sup>CL</sup> <b>ORAL SOLN</b><br>quinapril (generic Accupril)<br>trandolapril (generic Mavik) |   |
| <b>ACE INHIBITOR/DIURETIC COMBINATIONS</b>  |  |   |
| enalapril/HCTZ (generic Vaseretic)<br>lisinopril/HCTZ (generic Prinzide, Zestoretic)  | benazepril/HCTZ (generic Lotensin HCT)<br>captopril/HCTZ (generic Capozide)<br>fosinopril/HCTZ (generic Monopril HCT)<br>moexipril/HCTZ (generic Uniretic)<br>quinapril/HCTZ (generic Accuretic)   |   |
| <b>ANGIOTENSIN RECEPTOR BLOCKERS</b>  |  |   |
| irbesartan (generic Avapro)<br>losartan (generic Cozaar)<br>olmesartan (generic Benicar)<br>valsartan (generic Diovan)              | azilsartan medoxomil (generic Edarbi) <sup>NR</sup><br>candesartan (generic Atacand)<br>EDARBI (azilsartan)<br>eprosartan (generic Teveten)<br>telmisartan (generic Micardis)  |   |

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**ANGIOTENSIN MODULATORS (Continued)**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|--|---|--|
| <b>ANGIOTENSIN RECEPTOR BLOCKER/DIURETIC COMBINATIONS</b>              |   | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed TWO preferred agents within the last 12 months</li> <li>▪ Non-preferred combination products may be covered as individual prescriptions without prior authorization</li> </ul> |
| irbesartan/HCTZ (generic Avalide)                                      | candesartan/HCTZ (generic Atacand-HCT)  |  |
| losartan/HCTZ (generic Hyzaar)   | EDARBYCLOR (azilsartan/chlorthalidone)  |  |
| olmesartan/HCTZ (generic Benicar-HCT)                                  | telmisartan/HCTZ (generic Micardis-HCT)   |  |
| valsartan/HCTZ (generic Diovan-HCT)                                    |   |  |
| <b>ANGIOTENSIN MODULATOR/<br/>CALCIUM CHANNEL BLOCKER COMBINATIONS</b> |   |  |
| amlodipine/benazepril (generic Lotrel)                                 | amlodipine/olmesartan/HCTZ (generic Tribenzor)  |  |
| amlodipine/olmesartan (generic Azor)                                   | amlodipine/telmisartan (generic Twynsta)  |  |
| amlodipine/valsartan (generic Exforge)                                 | amlodipine/valsartan/HCTZ (generic Exforge HCT)   |  |
|  | PRESTALIA (perindopril/amlodipine)  |  |
|  | trandolapril/verapamil (generic Tarka)  |  |
|  | WIDAPLIK <sup>NR, QL</sup><br>(telmisartan/amlodipine/indapamide)   |  |
| <b>DIRECT RENIN INHIBITORS</b>   |   | <ul style="list-style-type: none"> <li>▪ <b>Direct Renin Inhibitors/Direct Renin Inhibitor Combinations:</b> May be approved with a history of TWO preferred ACE Inhibitors or Angiotensin Receptor Blockers within the last 12 months</li> </ul>  |
|  | aliskiren (generic Tekturna) <sup>QL</sup>  |  |
| <b>DIRECT RENIN INHIBITOR COMBINATIONS</b>                             |   |  |
|  | TEKTURNA/HCTZ (aliskiren/HCTZ)  |  |
| <b>NEPRILYSIN INHIBITOR COMBINATION</b>                                |   | Drug Specific Criteria   |
| ENTRESTO (sacubitril/valsartan) <sup>CL, QL</sup>                      | ENTRESTO (sacubitril/valsartan) <sup>CL, QL</sup><br><b>SPRINKLE CAP</b><br>sacubitril/valsartan (generic Entresto) <sup>CL, NR, QL</sup> | <ul style="list-style-type: none"> <li>• <b>Entresto/ sacubitril-valsartan:</b> May be approved in patients ages ≥1 years old and with a diagnosis of heart failure</li> </ul>   |

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**ANTHELMINTICS**

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|---|--|---|
| albendazole (generic Albenza)<br>BILTRICIDE (praziquantel)<br>ivermectin (generic Stromectol) | EMVERM (mebendazole) <sup>CL</sup><br>praziquantel (generic Biltricide)<br>STROMEKTOL (ivermectin) | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class within the last 6 months</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>• <b>Emverm:</b> Approval will be considered for indications not covered by preferred agents</li> </ul> |

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**ANTI-ALLERGENS, ORAL**

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|------------------|--|---|
|                  | <p>GRASTEK (timothy grass pollen allergen)<sup>AL,CL,QL</sup></p> <p>ODACTRA (Dermatophagoides farinae and Dermatophagoides pteronyssinus)<sup>AL,CL,QL</sup></p> <p>ORALAIR (sweet vernal/orchard/rye/timothy/kentucky blue grass mixed pollen allergen extract)<sup>CL</sup></p> <p>PALFORZIA (peanut allergen powder-dnfp)<sup>AL,CL</sup></p> <p>RAGWITEK (weed pollen-short ragweed)<sup>AL,CL,QL</sup></p> | <p><b>All agents require initial dose to be given in a healthcare setting</b></p> <p>Drug-specific criteria:</p> <p><b>GRASTEK</b></p> <ul style="list-style-type: none"> <li>Confirmed by positive skin test or in vitro testing for pollen specific IgE antibodies for Timothy grass or cross-reactive grass pollens.</li> <li>For use in persons 5 through 65 years of age.</li> </ul> <p><b>ODACTRA</b></p> <ul style="list-style-type: none"> <li>Confirmed by positive skin test to licensed house dust mite allergen extracts or in vitro testing for IgE antibodies to Dermatophagoides farinae and Dermatophagoides pteronyssinus house dust mite</li> <li>For use in persons 5 through 65 years of age</li> </ul> <p><b>ORALAIR</b></p> <ul style="list-style-type: none"> <li>Confirmed by positive skin test or in vitro testing for pollen specific IgE antibodies for Sweet Vernal, Orchard, Perennial Rye, Timothy, and Kentucky Blue Grass Mixed Pollens.</li> <li>For use in patients 5 through 65 years of age.</li> </ul> <p><b>PALFORZIA</b></p> <ul style="list-style-type: none"> <li>Confirmed diagnosis of peanut allergy by allergist</li> <li>For use in patients ages 4 to 17; it may be continued in patients 18 years and older with documentation of previous use within the past 90 days</li> <li>Initial dose and increase titration doses should be given in a healthcare setting</li> <li>Should not be used in patients with uncontrolled asthma or concurrently on a NSAID</li> </ul> <p><b>RAGWITEK</b></p> <ul style="list-style-type: none"> <li>Confirmed by positive skin test or in vitro testing for pollen specific IgE antibodies for short ragweed pollen.</li> <li>For use in patients 5 through 65 years of age.</li> </ul> |

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**ANTIBIOTICS, GASTROINTESTINAL**

| Preferred Agents  | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|---|--|--|
| metronidazole <b>TAB</b><br>neomycin<br>tinidazole (generic Tindamax)<br>vancomycin (generic Firvanq) <sup>QL</sup> <b>SOLN</b> | AEMCOLO (rifamycin) <b>TAB</b><br>DIFICID (fidaxomicin) <sup>CL</sup> <b>TAB, SUSP</b><br>fidaxomicin (generic Dificid) <sup>CL,NR</sup> <b>TAB</b><br>FIRVANQ (vancomycin) <sup>QL</sup> <b>SOLN</b><br>LIKMEZ (metronidazole) <b>SUSP</b><br>metronidazole <b>CAPS</b><br>metronidazole 125mg <b>TAB</b><br>nitazoxanide<br>(generic Alinia) <b>TAB</b> <sup>AL, CL, QL</sup><br>paromomycin<br>SOLOSEC (secnidazole)<br>vancomycin <b>CAPS</b> (generic Vancocin) <sup>CL</sup><br>VOWST (fecal microbiota spores) <sup>AL,QL</sup> | <ul style="list-style-type: none"> <li>▪ Note: Although azithromycin, ciprofloxacin, and trimethoprim/sulfmethoxazole are not included in this review, they are available without prior authorization</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Alinia /nitazoxanide tablet:</b> Trial and failure with metronidazole is required for a diagnosis of giardiasis</li> <li>▪ <b>Dificid®/ fidaxomicin:</b> For diagnosis of C. difficile diarrhea (pseudomembranous colitis), trial and failure or intolerance to oral vancomycin is required. For diagnosis of relapsed or recurrent C. difficile, an appropriate ICD-10 diagnosis code must be submitted for coverage.</li> <li>▪ <b>vancomycin capsules:</b> Requires patient specific documentation of why the Firvanq/vancomycin solution is not appropriate for patient</li> </ul> |

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**ANTIBIOTICS, INHALED<sup>CL</sup>**

| Preferred Agents <sup>CL</sup>   | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|--|---|---|
| BETHKIS (tobramycin)<br>KITABIS PAK (tobramycin)<br>tobramycin (generic Tobi)<br>TOBI-PODHALER (tobramycin) <sup>CL,QL</sup> | ARIKAYCE (amikacin liposomal inh) <sup>CL</sup><br><b>SUSP</b><br>CAYSTON (aztreonam lysine) <sup>CL,QL</sup><br>tobramycin (generic Bethkis) | <ul style="list-style-type: none"> <li>▪ Diagnosis of Cystic Fibrosis is required for all agents<br/>ICD10 Group = E84, ICD9 = 277.00, 277.01, 277.02, 277.03, 277.09</li> </ul> Drug-specific criteria: <ul style="list-style-type: none"> <li>▪ <b>Arikayce</b>: Requires diagnosis of refractory MAC lung disease defined as patients who did not achieve negative sputum cultures after a minimum of 6 consecutive months of a multidrug background regimen therapy</li> <li>▪ <b>Cayston</b><sup>®</sup>: Trial of tobramycin via nebulizer and demonstration of TOBI<sup>®</sup> compliance required</li> <li>▪ <b>Tobi Podhaler</b><sup>®</sup>: Requires trial of tobramycin via nebulizer or documentation of why nebulized tobramycin cannot be used</li> </ul> |

**ANTIBIOTICS, TOPICAL**

| Preferred Agents  | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|---|---|---|
| bacitracin <b>OINT</b><br>bacitracin <b>OINT OTC</b><br>bacitracin/polymyxin (generic Polysporin)<br>mupirocin <b>OINT</b> (generic Bactroban)<br>neomycin/polymyxin/bacitracin (generic Neosporin, Triple AB)<br>neomycin/polymyxin/pramoxine<br>neomycin/polymyxin/bacitracin/pramoxine | bacitracin <b>PCKT-OTC</b><br>CENTANY (mupirocin)<br>gentamicin <b>OINT, CREAM</b><br>mupirocin <b>CREAM</b> (generic Bactroban) <sup>CL</sup><br>XEPI (ozenoxacin) <sup>NR</sup> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed ALL preferred agents within this drug class within the last 12 months</li> </ul> Drug-specific criteria: <ul style="list-style-type: none"> <li>▪ <b>Mupirocin</b><sup>®</sup> <b>Cream</b>: Clinical reason the ointment cannot be used</li> </ul> |

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**ANTIBIOTICS, VAGINAL**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|--|---|--|
| CLEOCIN <b>CREAM</b> (clindamycin)<br>CLEOCIN <b>OVULES</b> (clindamycin)<br>metronidazole, vaginal<br>NUVESSA (metronidazole) | clindamycin <b>CREAM</b> (generic Cleocin) ▪<br>CLINDESSE (clindamycin)<br>metronidazole (generic Nuveessa) <sup>NR</sup><br>VANDAZOLE (metronidazole)<br>XACIATO (clindamycin phosphate) <sup>AL</sup><br><b>GEL</b> | ▪ Non-preferred agents will be approved for patients who have failed a therapeutic trial (duration = 3 days) with <b>ONE</b> preferred agent within this drug class within the last 6 months |

**ANTICOAGULANTS**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|--|---|--|
| dabigatran etexilate (generic Pradaxa) <b>CAPS</b><br>ELIQUIS (apixaban) <b>DOSE PACK, TAB</b><br>enoxaparin (generic Lovenox) <b>INJ</b><br>warfarin (generic Coumadin) <b>TAB</b><br>XARELTO (rivaroxaban) 10 mg, 15 mg, 20 mg <b>TAB</b><br>XARELTO (rivaroxaban) 2.5 mg <sup>CL,QL</sup> <b>TAB</b><br>XARELTO DOSE PACK (rivaroxaban) | ELIQUIS (apixaban) <b>SPRINKLE, SUSP</b><br>fondaparinux (generic Arixtra) <b>INJ</b><br>FRAGMIN (dalteparin) <b>INJ</b><br>PRADAXA (dabigatran) <b>CAPS, PELLETS</b><br>rivaroxaban (generic Xarelto) <sup>NR</sup> <b>TAB</b><br>rivaroxaban (generic Xarelto) <sup>AL,CL,NR</sup> <b>SUSP</b><br>SAVAYSA (edoxaban) <sup>CL,QL</sup> <b>TAB</b><br>XARELTO (rivaroxaban) <sup>CL</sup> <b>SUSP</b> | ▪ Non-preferred agents will be approved for patients who have failed <b>ONE</b> preferred agent within this drug class within the last 12 months<br><br>Drug-specific criteria:<br>▪ <b>Coumadin®</b> : Clinical reason generic warfarin cannot be used<br>▪ <b>Savaysa®</b> : Approved diagnoses include:<br>Stroke and systemic embolism (SE) risk reduction in nonvalvular atrial fibrillation (NVAf) OR<br>Treatment of deep vein thrombosis (DVT) and pulmonary embolism (PE) following 5-10 days of parenteral anticoagulant therapy<br>▪ <b>Xarelto 2.5mg</b> : Use limited to reduction of risk of major cardiovascular events (cardiovascular death, myocardial infarction, and stroke) in patients with chronic coronary artery disease or peripheral artery disease<br>▪ <b>Xarelto/ rivaroxaban Suspension</b> : Approved for patients ≤12 years of age or if there is a clinical reason why a preferred solid dosage form cannot be used. |

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**ANTIEMETICS/ANTIVERTIGO AGENTS**

| Preferred Agents   | Non-Preferred Agents   | Prior Authorization/Class Criteria  |
|--|--|---|
| <b>CANNABINOIDS</b>  |  |   |
| dronabinol (generic Marinol) <sup>AL</sup>   | dronabinol (generic Syndros) <sup>AL, NR, QL</sup><br><b>SOLN</b>  | <ul style="list-style-type: none"> <li>Non-preferred agents will be approved for patients who have failed ONE preferred agent within this drug class within the same subclass</li> </ul>  |
| <b>5HT3 RECEPTOR BLOCKERS</b>  |  |   |
| ondansetron (generic Zofran/Zofran ODT) <sup>QL</sup>  | ANZEMET (dolasetron)<br>granisetron (generic Kytril)<br>Nereus (tradipatant) <sup>NR, QL</sup> <b>CAPS</b><br>ondansetron 16mg <b>ODT (generic Zofran ODT)</b><br>SANCUSO (granisetron) <sup>CL</sup>  | <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li><b>Akynzeo®:</b> Approved for Moderately/Highly emetogenic chemotherapy with dexamethasone and a 5-HT3 antagonist</li> <li><b>Regimens include:</b> AC combination (Doxorubicin or Epirubicin with Cyclophosphamide), Aldesleukin, Amifostine, Arsenic trioxide, Azacitidine, Bendamustine, Busulfan, Carmustine, Carbplatin, Cisplatin, Clofarabine, Cyclophosphamide, Cytarabine, Dacarbazine, Dactinomycin, Daunorubicin, Epirubicin, Etoposide, Hexamethylmelamine, Idarubicin, Ifosfamide, Imatinib, Interferon α, Irinotecan, Mechlorethamine, Melphalan, Methotrexate, Oxaliplatin, Procarbazine, Streptozotocin, Temozolomide</li> <li><b>Diclegis/doxylamine-pyridoxine)/ Bonjesta:</b> Approved only for treatment of nausea and vomiting of pregnancy</li> <li><b>Sancuso®:</b> Documentation of oral dosage form intolerance</li> </ul> |
| <b>NK-1 RECEPTOR ANTAGONIST</b>  |  |   |
| aprepitant (generic Emend) <b>CAPS</b> <sup>QL</sup>   | AKYNZEO (netupitant/palonosetron) <sup>CL</sup><br>aprepitant (generic Emend) <b>PACK</b><br>EMEND (aprepitant) <b>CAPS, PACK, POWDER</b> <sup>QL</sup>  |   |
| <b>TRADITIONAL ANTIEMETICS</b>   |  |   |
| DICLEGIS (doxylamine/pyridoxine) <sup>CL, QL</sup><br>dimenhydrinate (generic Dramamine) <b>OTC</b><br>meclizine (generic Antivert)<br>metoclopramide (generic Reglan)<br>phosphoric acid/dextrose/fructose (generic Emetrol) <b>SOLN</b><br>prochlorperazine(generic Compazine)<br>promethazine (generic Phenergan) <b>SYRUP, TAB</b><br>promethazine 12.5mg, 25mg <b>SUPPOSITORY</b><br>scopolamine <b>TRANSDERMAL</b><br>TRANSDERM-SCOP (scopolamine) | BONJESTA (doxylamine/pyridoxine) <sup>CL, QL</sup><br>COMPRO (prochlorperazine)<br>doxylamine/pyridoxine (generic Diclegis) <sup>CL, QL</sup><br>prochlorperazine <b>SUPPOSITORY</b> (generic Compazine)<br>promethazine <b>SUPPOSITORY</b> 50mg<br>trimethobenzamide <b>TAB</b> (generic Tigan) |   |

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**ANTIFUNGALS, ORAL**

| Preferred Agents  | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|---|---|---|
| clotrimazole (mucous membrane, troche)<br>fluconazole <b>SUSP, TAB</b> (generic Diflucan)<br>griseofulvin <b>SUSP</b><br>griseofulvin microsize <b>TAB</b><br>nystatin <b>SUSP</b><br>terbinafine (generic Lamisil) | BREXAFEMME (ibrexafungerp) <sup>QL</sup><br>CRESEMBA (isavuconazonium) <sup>CL</sup><br>flucytosine (generic Ancobon) <sup>CL</sup><br>griseofulvin ultramicrosize (generic GRIS-PEG)<br>itraconazole (generic Sporanox) <sup>CL</sup><br>ketoconazole (generic Nizoral)<br>ORAVIG (miconazole) <sup>QL</sup> <b>BUCCAL</b><br>NOXAFIL (posaconazole) <sup>AL,CL</sup> <b>SUSP, TAB</b><br>NOXAFIL (posaconazole) <sup>AL,CL</sup> <b>POWDERMIX</b><br>nystatin <b>TAB</b><br>posaconazole (generic Noxafil) <sup>AL,CL</sup><br>TOLSURA (itraconazole) <sup>CL</sup><br>VIVJOA (oteseconazole) <b>CAPS</b><br>voriconazole (generic VFEND) <sup>CL</sup> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of TWO diagnosis-appropriate preferred agents within this drug class</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Cresemba®</b>: Approved for diagnosis of invasive aspergillosis or invasive mucormycosis</li> <li>▪ <b>Flucytosine</b>: Approved for diagnosis of: <u>Candida</u>: Septicemia, endocarditis, UTIs <u>Cryptococcus</u>: Meningitis, pulmonary infections</li> <li>▪ <b>Noxafil/ posaconazole DR tablets, oral suspension, PowderMix® for delayed oral suspension: For prophylaxis of invasive Aspergillus and Candida infections, no preferred agent</b> trial is required in severely immunocompromised patients (i.e., Neutropenia Myelodysplastic Syndrome (MDS), Neutropenic Acute Myeloid Leukemia (AML), Neutropenic hematologic malignancies, Graft vs. Host disease(GVHD), Immunosuppression secondary to hematopoietic stem cell transplant</li> <li>▪ <b>Noxafil® Powdermix</b>: pediatric patients 2 years of age and older who weigh 40 kg or less</li> <li>▪ <b>Noxafil/ posaconazole Suspension</b>:<br/>Oropharyngeal/esophageal candidiasis refractory to itraconazole and/or fluconazole and;<br/>Prophylaxis of invasive Aspergillus and Candida infections</li> <li>▪ <b>Sporanox®/itraconazole</b>: Approved for diagnosis of Aspergillosis, Blastomycosis, Histoplasmosis, Onychomycosis due to terbinafine-resistant dermatophytes, Oropharyngeal/ esophageal candidiasis refractory to fluconazole</li> <li>▪ <b>Sporanox® Liquid</b>: Clinical reason solid oral cannot be used</li> <li>▪ <b>Tolsura</b>: Approved for diagnosis of Aspergillosis, Blastomycosis, and Histoplasmosis and requires a trial and failure of generic itraconazole</li> <li>▪ <b>Vfend/voriconazole</b>: No trial for diagnosis of Myelodysplastic Syndrome (MDS), Neutropenic Acute Myeloid Leukemia (AML), Graft vs. Host disease (GVHD), Candidemia (candida krusei), Esophageal Candidiasis, Blastomycosis, <i>S. apiospermum</i> and <i>Fusarium spp.</i>, Oropharyngeal/esophageal candidiasis refractory to fluconazole</li> </ul> |

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**ANTIFUNGALS, TOPICAL**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|--|---|---|
| <b>ANTIFUNGAL</b>  |   |   |
| clotrimazole <b>CREAM</b> (generic Lotrimin) RX, OTC<br>clotrimazole (generic Lotrimin) <b>SOLN</b> -Rx<br>ketoconazole <b>CREAM, SHAMPOO</b> (generic Nizoral)<br>miconazole <b>CREAM, POWDER</b> OTC<br>nystatin <b>CREAM, POWDER</b><br>terbinafine OTC (generic Lamisil AT)<br>tolnaftate <b>AERO-POWDER OTC, CREAM-OTC, SOLN-OTC</b> (generic Tinactin) | ALEVAZOL (clotrimazole) OTC<br>ciclopirox <b>CREAM, GEL, SUSP</b> (generic Ciclodan, Loprox)<br>ciclopirox <b>SHAMPOO</b> (generic Loprox)<br>clotrimazole <b>SOLN OTC</b><br>DESENEXT <b>POWDER OTC</b> (miconazole)<br>econazole (generic Spectazole)<br>ERTACZO (sertaconazole)<br>FUNGOID (miconazole) <b>OTC</b><br>ketoconazole <b>FOAM<sup>CL</sup></b> (generic Extina, Ketodan)<br>LASOLEX AG 2% (miconazole nitrate 2%) <b>GEL<sup>NR, AL</sup></b><br>LASOLEX (clotrimazole) <sup>NR</sup> <b>SOLN (OTC)</b><br>LOPROX (ciclopirox) <b>SUSP, SHAMPOO, CREAM</b><br>LOTRIMIN AF <b>CREAM</b> OTC (clotrimazole)<br>LOTRIMIN ULTRA (butenafine)<br>miconazole OTC <b>OINT, SPRAY, SOLN</b><br>miconazole/zinc oxide/petrolatum (generic Vusion)<br>naftifine <b>CREAM, GEL</b> (generic Naftin)<br>oxiconazole (generic Oxistat)<br>tavaborole <b>SOLN<sup>CL</sup></b> (generic Kerydin)<br>tolnaftate <b>POWDER OTC, SPRAY<sup>NR</sup></b><br>TRIPENICOL (undecylenic acid) <b>CREAM-OTC</b><br>VOTRIZA-AL (clotrimazole) <b>LOTION OTC</b> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of TWO preferred agents within the last 6 months</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Extina/ Ketodan/ ketoconazole foam:</b> Requires trial and failure or contraindication to other ketoconazole forms</li> <li>▪ <b>tavaborole:</b> Approved diagnoses include Onychomycosis of the toenails due to <i>T.rubrum</i> OR <i>T. Mentagrophytes</i></li> <li>▪ <b>ciclopirox nail lacquer:</b> No trial required in diabetes, peripheral vascular disease (PVD), immunocompromised OR contraindication to oral terbinafine</li> </ul> |
| <b>ANTIFUNGAL/STEROID COMBINATIONS</b>   |   |   |
| clotrimazole/betamethasone <b>CREAM</b> (generic Lotrisone)<br>nystatin/triamcinolone (generic Mycolog) <b>CREAM, OINT</b>   | clotrimazole/betamethasone <b>LOTION</b> (generic Lotrisone)  |   |

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## ANTI-HISTAMINES, MINIMALLY SEDATING

| Preferred Agents   | Non-Preferred Agents   | Prior Authorization/Class Criteria  |
|--|--|---|
| cetirizine <b>TAB</b> (generic Zyrtec)<br>cetirizine <b>SOLN (OTC)</b> (generic Zyrtec)<br>levocetirizine <b>TAB</b> (generic Xyzal)<br>loratadine <b>TAB, SOLN</b> (generic Claritin) | cetirizine <b>CHEWABLE</b> (generic Zyrtec)<br>cetirizine <b>SOLN (Rx)</b> (generic Zyrtec)<br>desloratadine (generic Clarinex)<br>desloratadine ODT (generic Clarinex Reditabs)<br>desloratadine <sup>NR</sup> <b>SOLN</b><br>fexofenadine (generic Allegra)<br>fexofenadine 180mg (generic Allegra 180mg) <sup>QL</sup><br>levocetirizine (generic Xyzal) <b>SOLN</b><br>loratadine <b>CAPS, CHEWABLE, ODT</b> (generic Claritin Reditabs) | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed TWO preferred agents within this drug class</li> <li>▪ Combination products not covered – individual products may be covered</li> </ul> |

## ANTI-HYPERTENSIVES, SYMPATHOLYTICS

| Preferred Agents  | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|---|---|--|
| clonidine <b>TAB</b> (generic Catapres)<br>clonidine <b>TRANSDERMAL</b><br>guanfacine (generic Tenex)<br>methyldopa | clonidine 0.05mg <sup>NR</sup><br>clonidine ER (generic Nexiclon) <sup>CL</sup><br>JAVADIN (clonidine) <sup>NR</sup> <b>SOLN</b><br>methyldopa/hydrochlorothiazide<br>NEXICLON XR (clonidine ER) <sup>CL</sup> <b>TAB</b> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a 30-day trial with ONE preferred agent within this drug class</li> </ul> Drug Specific Criteria <ul style="list-style-type: none"> <li>▪ <b>Nexiclon/ clonidine ER:</b> Clinical reason why the preferred clonidine tablet or transdermal cannot be used</li> </ul> |

## ANTI-HYPERURICEMICS

| Preferred Agents  | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|---|--|--|
| allopurinol (generic Zyloprim)<br>colchicine <b>TAB</b> (generic Colcrys)<br>probenecid | allopurinol 200mg<br>colchicine <b>CAPS</b> (generic Mitigare)<br>febuxostat (generic Uloric) <sup>CL</sup><br>GLOPERBA <b>SOLN</b> (colchicine) <sup>CL,QL</sup><br>MITIGARE (colchicine)<br>probenecid/colchicine (generic Col-Probenecid) | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial with ONE preferred agent within this drug class</li> <li>▪ <b>Gloperba:</b> Approved for documented swallowing disorder</li> <li>▪ <b>Uloric/febuxostat:</b> Clinical reason why allopurinol cannot be used</li> </ul> |

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**ANTIMIGRAINE AGENTS, OTHER**

| Preferred Agents  | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|---|--|--|
| AIMOVIG (erenumab-aooe) <sup>CL,QL</sup><br>AJOVY (fremanezumab-vfrm) <sup>CL, QL</sup><br><b>PEN, Autoinjector</b><br>AJOVY (fremanezumab-vfrm)<br><b>Autoinjector 3-pack</b> <sup>CL,QL</sup><br>EMGALITY 120 mg/mL (galcanezumab-gnlm) <sup>CL, QL</sup> <b>PEN, SYRINGE</b><br>NURTEC ODT (rimegepant) <sup>AL,CL,QL</sup><br>QULIPTA (atogepant) <sup>AL,CL,QL</sup><br>UBRELVY (ubrogepant) <sup>AL,CL, QL</sup> <b>TAB</b> | BREKIYA (dihydroergotamine mesylate) <sup>NR</sup><br>diclofenac (generic Cambia) <b>POWDER</b><br>dihydroergotamine mesylate <b>NASAL</b><br>ELYXYB (celecoxib) <sup>AL,QL</sup> <b>SOLN</b><br>EMGALITY 100 mg (galcanezumab-gnlm) <sup>CL,QL</sup> <b>SYR</b><br>MIGERGOT (ergotamine/caffeine) <b>RECTAL</b><br>MIGRANAL (dihydroergotamine) <b>NASAL</b><br>REYVOW (lasmiditan) <sup>AL, CL,QL</sup> <b>TAB</b><br>ZAVZPRET (zavegepant) <sup>AL,CL,QL</sup> <b>NASAL</b> | <ul style="list-style-type: none"> <li>▪ All non-preferred agents will require a failed trial or contraindication of a preferred agent of the same indication</li> <li>▪ <b>For Acute Treatment:</b> agents will be approved for patients who have a failed trial or a contraindication to two triptans.</li> <li>▪ <b>For Prophylactic Treatment:</b> Require <math>\geq 4</math> migraines per month for <math>\geq 3</math> months and has tried and failed a <math>\geq 1</math> month trial of two medications: antidepressants (amitriptyline, venlafaxine), Beta blockers (propranolol, metoprolol, atenolol), anti-epileptics (divalproex, valproate, topiramate)</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Emgality 100mg</b> will only be approved for treatment of Episodic Cluster Headache</li> <li>▪ <b>Nurtec ODT:</b> for use in acute treatment, will be approved for patients who have a failed trial or a contraindication to two triptans. For use in preventative treatment, will be approved for patients who have a failed trial of ONE preferred injectable CGRP.</li> <li>▪ <b>Qulipta:</b> May be approved for patients who have a failed trial of ONE preferred injectable CGRP</li> <li>▪ <b>CGRP Antagonists:</b> Use will be limited to one CGRP for acute use and one CGRP for prophylactic use.</li> </ul> |

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**ANTIMIGRAINE AGENTS, TRIPTANS<sup>QL</sup>**

| Preferred Agents  | Non-Preferred Agents  | Prior Authorization/Class Criteria  |  |
|---|---|---|--|
| <b>ORAL</b>   |   |   |  |
| rizatriptan (generic Maxalt)<br>rizatriptan ODT (generic Maxalt MLT)<br>sumatriptan | almotriptan (generic Axert)<br>eletriptan (generic Relpax)<br>frovatriptan (generic Frova)<br>IMITREX (sumatriptan)<br>naratriptan (generic Amerge)<br>RELPAX (eletriptan) <sup>QL</sup><br>sumatriptan/naproxen (generic Treximet)<br>SYMBRAVO (rizatriptan benzoate/meloxicam) <sup>AL,NR</sup> TAB<br>zolmitriptan (generic Zomig) | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed ALL preferred agents within this drug class</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Zembrace:</b> approved for patients who have failed ALL preferred agents</li> </ul> |  |
| <b>NASAL</b>  |   |   |  |
| IMITREX (sumatriptan)<br>sumatriptan (generic Imitrex Nasal)                        | TOSYMRA (sumatriptan)<br>zolmitriptan (generic Zomig)<br>ZOMIG (zolmitriptan)   |   |  |
| <b>INJECTABLE</b>   |   |   |  |
| sumatriptan <b>SYRINGE, VIAL</b>  | sumatriptan <b>KIT</b><br>ZEMBRACE SYMTOUCH (sumatriptan) <sup>CL</sup>   |   |  |

**ANTIPARASITICS, TOPICAL**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|--|---|---|
| NATROBA (spinosad)<br>permethrin 1% OTC (generic Nix)<br>permethrin 5% RX (generic Elimite)<br>pyrethrin/piperonyl butoxide (generic RID, A-200) | CROTAN (crotamiton) <b>LOTION</b><br>EURAX (crotamiton) <b>CREAM, LOTION</b><br>ivermectin (generic Sklice) <b>LOTION</b><br>malathion (generic Ovide)<br>PRURADIK (cromtamiton) <sup>NR</sup> LOTION<br>spinosad (generic Natroba)<br>VANALICE (piperonyl butoxide/pyrethrins) | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial with ONE preferred agent within this drug class within the past 6 months</li> </ul> |

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**ANTIPARKINSON'S AGENTS, ORAL**

| Preferred Agents   | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|--|--|--|
| <b>ANTICHOLINERGICS</b>  |  | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed ONE preferred agent within the same subclass</li> </ul>  |
| benztropine (generic Cogentin)<br>trihexyphenidyl (generic Artane)   |  |  |
| <b>COMT INHIBITORS</b>   |  | <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Carbidopa/Levodopa ODT:</b> Approved for documented swallowing disorder</li> <li>▪ <b>COMT Inhibitors:</b> Approved if using as add-on therapy with levodopa-containing drug</li> <li>▪ <b>Gocovri:</b> Required diagnosis of Parkinson's disease and had trial of or is intolerant to amantadine AND must be used as an add-on therapy with levodopa-containing drug</li> <li>▪ <b>Inbrija:</b> Approval upon diagnosis of Parkinson's disease and concurrent treatment with carbidopa/levodopa agent</li> <li>▪ <b>Neupro®:</b><br/>For Parkinsons: Clinical reason required why preferred agent cannot be used<br/>For Restless Leg (RLS): Requires trial OR Contraindication to ropinirole AND pramipexole</li> </ul> |
| entacapone (generic Comtan)  | ONGENTYS (opicapone)<br>tolcapone (generic Tasmar)   |  |
| <b>DOPAMINE AGONISTS</b>   |  | <ul style="list-style-type: none"> <li>▪ <b>Nourianz:</b> Approval upon diagnosis of Parkinson's disease and concurrent treatment with carbidopa/levodopa agent</li> <li>▪ <b>Osmolex ER:</b> Required diagnosis of Parkinson's disease or drug-induced extrapyramidal reactions and had trial of or is intolerant to amantadine IR</li> <li>▪ <b>Pramipexole ER:</b> Required diagnosis of Parkinson's along with preferred agent trial</li> <li>▪ <b>Ropinerole ER:</b> Required diagnosis of Parkinson's along with preferred agent trial</li> </ul>  |
| pramipexole (generic Mirapex)<br>ropinirole (generic Requip)   | bromocriptine (generic Parlodel)<br>NEUPRO (rotigotine) <sup>CL</sup><br>pramipexole ER (generic Mirapex ER) <sup>CL</sup><br>ropinirole ER (generic Requip XL) <sup>CL</sup>  |  |
| <b>MAO-B INHIBITORS</b>  |  |  |
| selegiline <b>CAPS, TABLET</b> (generic Eldepryl)  | rasagiline (generic Azilect) <sup>QL</sup><br>XADAGO (safinamide)  |  |
| <b>OTHER ANTIPARKINSON'S DRUGS</b>   |  |  |
| amantadine <b>CAPS, SYRUP TABLET</b> (generic Symmetrel)<br>carbidopa/levodopa (generic Sinemet)<br>carbidopa/levodopa ER (generic Sinemet CR) | carbidopa (generic Lodosyn)<br>carbidopa/levodopa ODT (generic Parcopa) <sup>CL</sup><br>carbidopa/levodopa ER (generic Rytary) <sup>NR</sup><br>CREXONT (carbidopa and levodopa ER.) <sup>QL</sup> <b>CAPS</b><br>DHIVY (carbidopa/levodopa) <sup>QL</sup><br>DUOPA (carbidopa/levodopa)<br>GOCOVRI (amantadine) <sup>CL,QL</sup><br>INBRIJA (levodopa) <sup>CL,QL</sup> <b>INHALER</b><br>levodopa/carbidopa/entacapone (generic Stalevo)<br>NOURIANZ (istradefylline) <sup>CL,QL</sup><br>OSMOLEX ER (amantadine) <sup>CL,QL</sup><br>RYTARY (carbidopa/levodopa) |  |

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**ANTIPSORIATICS, ORAL**

| Preferred Agents              | Non-Preferred Agents                    | Prior Authorization/Class Criteria  |
|-------------------------------|---|---|
| acitretin (generic Soriatane) | methoxsalen (generic Oxsoresalen-Ultra) | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial with the preferred agent within this drug class</li> <li>▪ Trial of acitretin (Pregnancy category X) not required in pregnancy or while attempting or planning pregnancy</li> </ul> |

**ANTIPSORIATICS, TOPICAL**

| Preferred Agents                       | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|--|---|--|
| calcipotriene <b>CREAM, OINT, SOLN</b> | calcitriol (generic Vectical) <sup>AL</sup> <b>OINT</b><br>calcipotriene <b>FOAM</b> (generic Sorilux)<br>calcipotriene/betamethasone <b>OINT</b><br>(generic Taclonex)<br>calcipotriene/betamethasone <b>SUSP</b><br>(generic Taclonex Scalp)<br>ENSTILAR<br>(calcipotriene/betamethasone)<br>SORILUX (calcipotriene)<br>ZORYVE 0.3% (roflumilast) <sup>AL</sup> <b>CREAM</b><br>ZORYVE 0.3% (roflumilast) <sup>AL,CL</sup><br><b>FOAM</b> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial with a preferred agent within this drug class</li> </ul> <p>Drug-Specific Criteria</p> <ul style="list-style-type: none"> <li>• <b>Zoryve Foam</b>- For diagnosis of Seborrheic Dermatitis, requires trial of a topical steroid or a topical calcineurin inhibitor AND trial of a topical antifungal.</li> </ul> |

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**ANTIVIRALS, ORAL**

| Preferred Agents  | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|---|---|---|
| <b>ANTI-COVID-19 DRUGS</b>  |   | <ul style="list-style-type: none"> <li>Non-preferred agents will be approved for patients who have failed a 10-day trial of ONE preferred agent within the same subclass</li> </ul>   |
| PAXLOVID (nirmatrelvir/ritonavir) <sup>CL,QL</sup>  |   |   |
| <b>ANTI-HERPETIC DRUGS</b>  |   |   |
| acyclovir (generic Zovirax)<br>famciclovir (generic Famvir)<br>valacyclovir (generic Valtrex) | acyclovir (generic Zovirax) <sup>CL</sup> <b>SUSP</b>   | <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li><b>Acyclovir Susp:</b> Prior authorization NOT required for children ≤ 12 years old</li> <li><b>Paxlovid:</b> Requires a diagnosis of COVID-19 and is limited to 1 dose pack per 30 days</li> <li><b>Xofluza:</b> Requires clinical, patient specific reason that a preferred agent cannot be used</li> </ul> |
| <b>ANTI-INFLUENZA DRUGS</b>   |   |   |
| oseltamivir (generic Tamiflu) <sup>QL</sup> <b>CAPS, SUSP</b>                                 | rimantadine (generic Flumadine)<br>RELENZA (zanamivir) <sup>QL</sup><br>TAMIFLU (oseltamivir) <sup>QL</sup> <b>CAPS, SUSP</b><br>XOFLUZA (baloxavir marboxil) <sup>AL,CL,QL</sup> |   |

**ANTIVIRALS, TOPICAL**

| Preferred Agents                              | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|---|--|--|
| acyclovir <b>OINT</b><br>docosanol <b>OTC</b> | acyclovir CREAM, (generic Zovirax)<br>DENAVIR (penciclovir) <sup>AL</sup><br>penciclovir (generic Denavir) <sup>AL</sup> | <ul style="list-style-type: none"> <li>Non-preferred agents will be approved for patients who have failed a trial with ONE preferred ORAL Antiviral agent</li> </ul> |

**ANXIOLYTICS**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|--|---|--|
| alprazolam (generic Xanax) <b>TAB</b><br>buspirone (generic Buspar)<br>chlordiazepoxide<br>diazepam (generic Valium) <b>TAB, SOLN</b><br>lorazepam (generic Ativan) <b>INTENSOL, TAB</b> | alprazolam ER (generic Xanax XR)<br>alprazolam <b>INTENSOL</b> <sup>CL</sup><br>alprazolam ODT<br>BUCAPSOL (buspirone hcl) <sup>CL</sup> <b>CAP</b><br>clorazepate (generic Tranxene-T)<br>diazepam <b>INTENSOL</b> <sup>CL</sup><br>LOREEV XR (lorazepam) <sup>AL</sup><br>meprobamate<br>oxazepam | <ul style="list-style-type: none"> <li>Non-preferred agents will be approved for patients who have failed a trial with TWO preferred agents within this drug class</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li><b>Alprazolam IntenSol®:</b> Requires trial of diazepam solution OR lorazepam IntenSol®</li> <li><b>Bucapsol:</b> Requires clinical reason why preferred buspirone can't be used.</li> <li><b>Diazepam IntenSol®:</b> Requires clinical reason why diazepam solution cannot be used</li> </ul> |

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**BETA BLOCKERS, ORAL**

| Preferred Agents                            | Non-Preferred Agents                           | Prior Authorization/Class Criteria   |
|---|--|--|
| <b>BETA BLOCKERS</b>                        |  | <p>Non-preferred agents will be approved for patients who have failed TWO diagnosis-appropriate preferred agents .</p> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Coreg CR/carvedilol ER:</b> Requires clinical reason generic IR product cannot be used</li> <li>▪ <b>Hemangeol®:</b> Covered for diagnosis of Proliferating Infantile Hemangioma</li> <li>▪ <b>Sotylize®:</b> Covered for diagnosis of life –threatening ventricular arrhythmias OR maintenance of normal sinus rhythm in highly symptomatic atrial fibrillation/flutter (AFIB/AFL)<br/>Requires clinical reason generic sotalol cannot be used</li> </ul> |
| atenolol (generic Tenormin)                 | acebutolol (generic Sectral)                   |  |
| atenolol/chlorthalidone (generic Tenoretic) | betaxolol (generic Kerlone)                    |  |
| bisoprolol (generic Zebeta)                 | BYSTOLIC (nebivolol)                           |  |
| bisoprolol/HCTZ (generic Ziac)              | HEMANGEOL (propranolol) <sup>AL,CL</sup>       |  |
| metoprolol (generic Lopressor)              | <b>SOLN</b>                                    |  |
| metoprolol ER (generic Toprol XL)           | INDERAL/INNOPRAN XL (propranolol ER)           |  |
| nebivolol (generic Bystolic)                | KAPSPARGO SPRINKLE (metoprolol ER)             |  |
| propranolol (generic Inderal)               | LOPRESSOR (metoprolol tartrate) <sup>NR</sup>  |  |
| propranolol ER (generic Inderal LA)         | <b>SOLN</b>                                    |  |
|   | metoprolol/HCTZ (generic Lopressor HCT)        |  |
|   | nadolol (generic Corgard)                      |  |
|   | pindolol (generic Viskin)                      |  |
|   | propranolol/HCTZ (generic Inderide)            |  |
|   | timolol (generic Blocadren)                    |  |
|   | TOPROL XL (metoprolol ER)                      |  |
| <b>BETA- AND ALPHA-BLOCKERS</b>             |  |  |
| carvedilol (generic Coreg)                  | carvedilol ER <sup>CL</sup> (generic Coreg CR) |  |
| labetalol (generic Trandate)                |  |  |
| <b>ANTIARRHYTHMIC</b>                       |  |  |
| sotalol (generic Betapace)                  | SOTYLIZE (sotalol) <sup>CL</sup>               |  |

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**BILE SALTS**

| Preferred Agents  | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|---|---|--|
| ursodiol <b>CAPSULE</b> 300 mg (generic Actigall)<br>ursodiol 250 mg <b>TABLET</b> (generic URSO)<br>ursodiol 500 mg <b>TABLET</b> (generic URSO FORTE) | BYLVAY (odevixibat) <b>CAP</b> <sup>QL</sup> ,<br><b>PELLET</b> <sup>AL</sup><br>CHENODAL (chenodiol)<br>CHOLBAM (cholic acid)<br>CTEXLI (chenodiol) <b>TAB</b><br>IQIRVO (elafibranor) <sup>QL</sup> <b>TAB</b><br>LIVDELZI (seladelpar) <b>CAP</b><br>LIVMARLI (maralixibat) <sup>QL</sup><br><b>SOLN</b> <sup>AL</sup> , <b>TAB</b><br><b>LYNAVOY (linerixibat)</b> <sup>NR</sup> <b>TAB</b><br>OCALIVA (obeticholic acid)<br>RELTONE (ursodiol 200mg,400mg)<br><b>CAP</b> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial with ONE preferred agent within this drug class</li> </ul> |

**BLADDER RELAXANT PREPARATIONS**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|--|---|--|
| fesoterodine (generic Toviaz)<br>MYRBETRIQ (mirabegron) <sup>AL</sup> <b>TAB</b><br>oxybutynin IR, ER (generic Ditropan/Ditropan XL) | darifenacin ER (generic Enablex)<br>flavoxate HCL<br>GEMTESA (vibegron) <sup>AL,QL</sup><br>mirabegron ER (generic Myrbetriq) <b>TAB</b><br><b>mirabegron ER (generic Myrbetriq)</b><br><b>SUSP</b> <sup>NR</sup><br>MYRBETRIQ (mirabegron) <b>SUSP</b> <sup>AL,CL,QL</sup><br>oxybutynin 2.5mg<br>OXYTROL (oxybutynin)<br>solifenacin (generic Vesicare)<br>tolterodine IR, ER (generic Detrol/ Detrol LA)<br>TOVIAZ (fesoterodine ER)<br>trospium IR, ER (generic Sanctura/ Sanctura XR)<br>VESICARE (solifenacin)<br>VESICARE LS <b>SUSP</b> (solifenacin) <sup>AL</sup> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial with ONE preferred agent within this drug class</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Myrbetriq suspension:</b> Covered for pediatric patients ≥ 3 years old with a diagnosis of Neurogenic Detrusor Overactivity (NDO)</li> </ul> |

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**BONE RESORPTION SUPPRESSION AND RELATED DRUGS**

| Preferred Agents  | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|---|---|--|
| <b>BISPHOSPHONATES</b>  |   |  |
| alendronate (generic Fosamax) <b>TAB</b><br>ibandronate (generic Boniva) <sup>QL</sup>                  | alendronate <b>SOLN</b> (generic Fosamax) <sup>QL</sup><br>ATELVIA DR (risedronate) <sup>CL</sup><br>BINOSTO (alendronate) <sup>CL</sup><br>FOSAMAX PLUS D (alendronate sodium/ cholecalciferol) <sup>QL</sup><br>risedronate (generic Actonel) <sup>QL</sup> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within the same subclass</li> </ul> Drug-specific criteria: <ul style="list-style-type: none"> <li>▪ <b>Atelvia DR®</b>: Requires clinical reason alendronate cannot be taken on an empty stomach</li> <li>▪ <b>Binosto®</b>: Requires clinical reason why alendronate tablets OR Fosamax® solution cannot be used</li> <li>▪ <b>Forteo/ teriparatide</b>: Covered for high risk of fracture</li> </ul> High risk of fracture: <ul style="list-style-type: none"> <li>• BMD -3 or worse</li> <li>• Postmenopausal women with history of non-traumatic fractures</li> <li>• Postmenopausal women with 2 or more clinical risk factors                             <ul style="list-style-type: none"> <li>○ Family history of non-traumatic fractures</li> <li>○ DXA BMD T-score ≤ -2.5 at any site</li> <li>○ Glucocorticoid use ≥ 6 months at 7.5 dose of prednisolone equivalent</li> <li>○ Rheumatoid Arthritis</li> </ul> </li> <li>• Postmenopausal women with BMD T-score ≤ -2.5 at any site with any clinical risk factors                             <ul style="list-style-type: none"> <li>○ More than 2 units of alcohol per day</li> <li>○ Current smoker</li> </ul> </li> <li>• Men with primary or hypogonadal osteoporosis</li> <li>• Osteoporosis associated with sustained systemic glucocorticoid therapy</li> <li>• Trial of calcitonin-salmon not required</li> <li>• Maximum of 24 months treatment per lifetime</li> </ul> |
| <b>OTHER BONE RESORPTION SUPPRESSION AND RELATED DRUGS</b>  |   |  |
| calcitonin-salmon <b>NASAL</b><br>FORTEO (teriparatide) <sup>CL,QL</sup><br>raloxifene (generic Evista) | BONSTY (teriparatide) <sup>QL,NR</sup><br>EVISTA (raloxifene)<br>teriparatide (generic Forteo) <sup>CL,QL</sup><br>TYMLOS (abaloparatide)   |  |

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**BPH (BENIGN PROSTATIC HYPERPLASIA) TREATMENTS**

| Preferred Agents  | Non-Preferred Agents   | Prior Authorization/Class Criteria  |
|---|--|---|
| <b>ALPHA BLOCKERS</b>   |  | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within the same subclass.</li> </ul>   |
| alfuzosin (generic Uroxatral) <sup>CL</sup><br>doxazosin (generic Cardura)<br>tamsulosin (generic Flomax) <sup>CL</sup><br>terazosin (generic Hytrin) | CARDURA XL (doxazosin) <sup>CL</sup><br>silodosin (generic Rapaflo)<br>TEZRULY (terazosin) <sup>CL,NR</sup> <b>SOLN</b>                          |   |
| <b>5-ALPHA-REDUCTASE (5AR) INHIBITORS</b>   |  | Drug-specific criteria: <ul style="list-style-type: none"> <li>▪ <b>Alfuzosin/dutasteride/finasteride</b> <ul style="list-style-type: none"> <li>• Covered for males only</li> </ul> </li> <li>▪ <b>Cardura XL®</b>: Requires clinical reason generic IR form cannot be used</li> <li>▪ <b>Flomax/ tamsulosin</b>: Covered for males and may be covered for females for a 7-day supply with diagnosis of acute kidney stones</li> <li>▪ <b>Jalyn/ dutasteride-tamsulosin</b>: Requires clinical reason why individual agents cannot be used</li> <li>▪ <b>Tezruly</b>: Clinical reason why oral capsule is not appropriate</li> </ul> |
| dutasteride (generic Avodart) <sup>CL</sup><br>finasteride (generic Proscar) <sup>CL</sup>  | dutasteride/tamsulosin (generic Jalyn) <sup>CL</sup><br>ENTADFI (finasteride/tadalafil)<br>finasteride/tadalafil (generic Entadfi) <sup>NR</sup> |   |

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**BRONCHODILATORS, BETA AGONIST**

| Preferred Agents  | Non-Preferred Agents   | Prior Authorization/Class Criteria  |
|---|--|---|
| <b>INHALERS – Short Acting</b>  |  | <ul style="list-style-type: none"> <li>• Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within the same subclass.</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>• <b>Xopenex/levalbuterol solution:</b> Covered for cardiac diagnoses or side effect of tachycardia with albuterol product</li> </ul> |
| albuterol HFA (generic Proventil HFA)<br>VENTOLIN HFA (albuterol)<br>XOPENEX HFA (levalbuterol HFA) | albuterol HFA (generic ProAir HFA and Ventolin HFA)<br>levalbuterol HFA (generic Xopenex HFA)<br>PROAIR DIGIHALER (albuterol)<br>PROAIR RESPICLICK (albuterol)                             |   |
| <b>INHALERS – Long Acting</b>   |  |   |
| SEREVENT (salmeterol)   | STRIVERDI RESPIMAT (olodaterol)  |   |
| <b>INHALATION SOLUTION</b>  |  |   |
| albuterol (2.5mg/3ml premix or 2.5mg/0.5ml)<br>albuterol low dose (0.63mg/3ml & 1.25mg/3ml)         | arformoterol tartrate (generic Brovana)<br>BROVANA (arformoterol)<br>formoterol fumarate (generic Perforomist)<br>levalbuterol (generic Xopenex) <sup>CL</sup><br>PERFOROMIST (formoterol) |   |
| <b>ORAL</b>   |  |   |
| albuterol <b>SYRUP</b>  | albuterol <b>TAB</b><br>albuterol ER (generic Vospire ER)<br>terbutaline (generic Brethine)  |   |

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**CALCIUM CHANNEL BLOCKERS, ORAL**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|--|---|---|
| <b>SHORT-ACTING</b>  |   | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within the same subclass.</li> <li>Drug-specific criteria:               <ul style="list-style-type: none"> <li>▪ <b>Katerzia/ Norliqva:</b> May be approved with documented swallowing difficulty</li> <li>▪ <b>Nifedipine:</b> May be approved without trial for diagnosis of Preterm Labor or Pregnancy Induced Hypertension (PIH)</li> <li>▪ <b>Nimodipine:</b> Covered without trial for diagnosis of subarachnoid hemorrhage</li> <li>▪ <b>Nimodipine solution:</b> Covered without trial for diagnosis of subarachnoid hemorrhage and documented swallowing difficulty</li> </ul> </li> </ul> |
| <b>Dihydropyridines</b>  |   |   |
|  | isradipine (generic Dynacirc)<br>nicardipine (generic Cardene)<br>nifedipine (generic Procardia) <sup>CL</sup><br>nimodipine (generic Nimotop) <sup>CL</sup><br>nimodipine (generic Nymalize) <sup>CL</sup> <b>SOLN</b><br>NYMALIZE (nimodipine) <b>SOLN</b>          |   |
| <b>Non-dihydropyridines</b>  |   |   |
| diltiazem (generic Cardizem)<br>verapamil (generic Calan/Isoptin)                  |   |   |
| <b>LONG-ACTING</b>   |   |   |
| <b>Dihydropyridines</b>  |   |   |
| amlodipine (generic Norvasc)<br>nifedipine ER (generic Procardia XL/<br>Adalat CC) | felodipine ER (generic Plendil)<br>KATERZIA (amlodipine) <sup>CL,QL</sup> <b>SUSP</b><br>levamlodipine (generic Conjupri)<br>nisoldipine (generic Sular)<br>NORLIQVA (amlodipine) <sup>AL,CL,QL</sup> <b>SOLN</b><br>SDAMLO (amlodipine) <sup>NR,AL</sup> <b>SOLN</b> |   |
| <b>Non-dihydropyridines</b>  |   |   |
| diltiazem ER (generic Cardizem CD)<br>verapamil ER <b>TAB</b>                      | diltiazem ER (generic Cardizem LA)<br>MATZIM LA (diltiazem ER)<br>TIAZAC (diltiazem)<br>verapamil ER <b>CAPS</b><br>verapamil 360mg <b>CAPS</b><br>verapamil ER (generic Verelan PM)<br>verapamil SR (generic Verelan)<br><b>CAPS</b>                                 |   |

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**CEPHALOSPORINS AND RELATED ANTIBIOTICS, ORAL**

| Preferred Agents  | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|---|--|--|
| <b>BETA LACTAM/BETA-LACTAMASE INHIBITOR COMBINATIONS</b>  |  | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a 3-day trial of ONE preferred agent within the same subclass</li> <br/> <li>Drug Specific Criteria               <ul style="list-style-type: none"> <li>• <b>Cefixime</b>- May be approved for a diagnosis of gonorrhea, with an appropriate ICD-10 diagnosis code without a 3-day trial of a preferred agent</li> <li>• <b>Cefpodoxime</b>- May be approved for a diagnosis of pyelonephritis, with an appropriate ICD-10 diagnosis code without a 3-day trial of a preferred agent</li> </ul> </li> </ul> |
| amoxicillin/clavulanate <b>TAB, SUSP</b>  | amoxicillin/clavulanate <b>CHEWABLE</b><br>amoxicillin/clavulanate ER (generic Augmentin XR)<br>AUGMENTIN (amoxicillin/clavulanate) <b>SUSP, TAB</b> |  |
| <b>CEPHALOSPORINS – First Generation</b>  |  |  |
| cefadroxil <b>CAPS, SUSP</b> (generic Duricef)<br>cephalexin <b>CAPS, SUSP</b> (generic Keflex) | cefadroxil <b>TAB</b> (generic Duricef)<br>cephalexin <b>TAB</b>   |  |
| <b>CEPHALOSPORINS – Second Generation</b>   |  |  |
| cefprozil (generic Cefzil)<br>cefuroxime <b>TAB</b> (generic Ceftin)                            | cefaclor (generic Ceclor)  |  |
| <b>CEPHALOSPORINS – Third Generation</b>  |  |  |
| cefdinir (generic Omnicef)  | cefixime (generic Suprax) <sup>CL</sup> <b>CAPS, SUSP, TAB</b><br>cefpodoxime (generic Vantin) <sup>CL</sup>   |  |

**COLONY STIMULATING FACTORS**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|--|---|--|
| FULPHILA (pegfilgrastim-jmdb) <b>SUB-Q</b><br>FYLNETRA (pegfilgrastim-pbbk) <b>SUB-Q</b><br>NEUPOGEN <b>DISP SYR (SUB-Q)</b><br>NEUPOGEN (filgrastim) <b>VIAL (IV)</b> | FILKRI (filgrastim-laha) <b>SYR<sup>NR</sup></b><br>GRANIX (tbo-filgrastim) <b>SUB-Q</b><br>LEUKINE (sargramostim) <b>INTRAVENOUS</b><br>NEULASTA (pegfilgrastim) <b>SYR, VIAL<sup>NR</sup></b><br>NIVESTYM (filgrastim-aafi) <b>SYR, VIAL</b><br>NYVEPRIA (pegfilgrastim-apgf)<br>RELEUKO (filgrastim-ayow) <b>SYR</b><br>ROLVEDON (eflapeggrastim-xnst) <b>SYR</b><br>STIMUFEND (pegfilgrastim-fpgk) <b>SUB-Q</b><br>UDENYCA (pegfilgrastim-cbqv) <b>AUTOINJ</b><br>UDENYCA (pegfilgrastim-cbqv) <b>SUB-Q</b><br>ZARXIO (filgrastim-sndz) <b>IV, SUB-Q</b><br>ZIEXTENZO (pegfilgrastim-bmez) <b>SUB-Q</b> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class</li> </ul> |

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**CONTRACEPTIVES, ORAL**

| Preferred Agents   | Non-Preferred Agents   | Prior Authorization/Class Criteria |
|--|--|------------------------------------|
| <p>All reviewed agents are recommended preferred at this time<br/><i>Only those products for review are listed.</i><br/>Brand name products may be subject to Maximum Allowable Cost (MAC) pricing or require substitution with a generic equivalent</p> <p>Specific agents can be looked up using the Drug Look-up Tool at:<br/><a href="https://ne.primetherapeutics.com/drug-lookup">https://ne.primetherapeutics.com/drug-lookup</a></p> | <p>AVERI (desogestrel and ethinyl estradiol kit)<sup>NR</sup></p> <p>GALBRIELA (norethindrone/ethinyl estradiol/ferrous fumarate)<sup>NR</sup> <b>CHEW</b></p> <p>LUIZZA (norethindrone ac/eth estradiol)<sup>NR</sup></p> <p>MELEYA (norethindrone)<sup>NR</sup></p> <p>ORQUIDEA (norethindrone)<sup>NR</sup></p> <p>ROSYRAH (levonorgestrel/ ethinyl estradiol/ ethinyl estradiol kit)<sup>NR</sup></p> <p>XARAH FE (norethindrone acetate and ethinyl estradiol and ferrous fumarate)<sup>NR</sup></p> <p>XELRIA FE (norethindrone and ethinyl estradiol and ferrous fumarate)<sup>NR</sup></p> <p>YASMIN (ethinyl estradiol/drospirenone)<sup>NR</sup></p> |                                    |

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**COPD (CHRONIC OBSTRUCTIVE PULMONARY DISEASE) AGENTS**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|--|---|--|
| <b>INHALERS</b>  |   | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within the same subclass OR Patient specific documentation of inability to use traditional inhaler device.</li> <li>Drug-specific criteria:               <ul style="list-style-type: none"> <li>▪ <b>Daliresp/roflumilast:</b> <ul style="list-style-type: none"> <li>Covered for diagnosis of severe COPD associated with chronic bronchitis</li> <li>Requires trial of a bronchodilator</li> <li>Requires documentation of one exacerbation in last year upon initial review</li> </ul> </li> <li>▪ <b>Dupixent</b> (For other indications, see Immunomodulators, Atopic Dermatitis and Asthma therapeutic classes): For COPD and an Eosinophilic Phenotype: Requires documentation of inadequately controlled COPD with eosinophils <math>\geq 300</math> cells/microliter AND two exacerbations OR one exacerbation that led to hospitalization while on and adherent to a <math>\geq 90</math>-day trial of triple therapy (LABA + LAMA + ICS). Prescribed by, or in consultation with a pulmonologist, immunologist, or an allergist.</li> </ul> </li> </ul> |
| ANORO ELLIPTA<br>(umeclidinium/vilanterol)<br>ATROVENT HFA (ipratropium)<br>COMBIVENT RESPIMAT (albuterol/<br>ipratropium)<br>SPIRIVA (tiotropium)<br>SPIRIVA RESPIMAT (tiotropium)<br>STIOLTO RESPIMAT<br>(tiotropium/olodaterol) | BEVESPI AEROSPHERE<br>(glycopyrolate/formoterol)<br>DUAKLIR PRESSAIR (aclidinium br<br>and formoterol fum)<br>INCRUSE ELLIPTA (umeclidinium)<br>tiotropium (generic Spiriva)<br>TUDORZA PRESSAIR (aclidinium br<br>umeclidinium ellipta (generic Incruse) <sup>NR</sup><br>umeclidinium/vilanterol (generic Anoro<br>Ellipta)<br>ipratropium bromide HFA (Atrovent) <sup>NR</sup> |  |
| <b>INHALATION SOLUTION</b>   |   |  |
| albuterol/ipratropium (generic Duoneb)<br>ipratropium <b>SOLN</b> (generic Atrovent)   | OHTUVAYRE (ensifentrine) inhalation<br>suspension<br>YUPELRI (revedfenacin)   |  |
| <b>ORAL AGENT</b>  |   |  |
| roflumilast (generic Daliresp) <sup>CL, QL</sup>   | DALIRESP (roflumilast) <sup>CL, QL</sup>  |  |

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**COUGH AND COLD, OPIATE COMBINATION**

| Preferred Agents | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|------------------|---|--|
|                  | guaifenesin/codeine <b>LIQUID</b><br>hydrocodone/homatropine <b>SYRUP</b><br>promethazine/codeine <b>SYRUP</b><br>promethazine/phenylephrine/codeine <b>SYRUP</b> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE dextromethorphan product</li> <li>▪ All codeine or hydrocodone containing cough and cold combinations are limited to <math>\geq 18</math> years of age</li> </ul> |

**CYSTIC FIBROSIS, ORAL**

| Preferred Agents | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|------------------|---|---|
|                  | ALYFTREK (vanzacaftor; tezacaftor; deutivacaftor) <sup>AL,CL</sup> <b>TAB</b><br>BRONCHITOL (mannitol) <sup>AL,CL,QL</sup><br>KALYDECO <b>PACKET, TAB</b><br>(ivacaftor) <sup>AL,CL,QL</sup><br>ORKAMBI (lumacaftor/ivacaftor) <b>PACKET, TAB</b> <sup>AL,CL,QL</sup><br>SYMDEKO<br>(tezacaftor/ivacaftor) <sup>AL,CL,QL</sup><br>TRIKAFTA(elexacaftor, tezacaftor, ivacaftor) <sup>AL, CL</sup> <b>PACKET, TAB</b> | Drug-specific criteria: <ul style="list-style-type: none"> <li>▪ <b>Alyfrek:</b> Diagnosis of CF and documentation of at least one F508del mutation or another responsive mutation in the CFTR gene.</li> <li>▪ <b>Bronchitol:</b> Approved for diagnosis of CF and documentation that the patient has passed the BRONCHITOL Tolerance Test</li> <li>▪ <b>Kalydeco®:</b> Diagnosis of CF and documentation of the drug-specific, FDA-approved mutation of CFTR gene</li> <li>▪ <b>Orkambi®:</b> Diagnosis of CF and documentation of presence of two copies of the F580del mutation (homozygous) of CFTR gene</li> <li>▪ <b>Symdeko:</b> Diagnosis of CF and documentation of the drug specific, FDA approved mutation of CFTR gene.</li> <li>▪ <b>Trikafta:</b> Diagnosis of CF and documentation of at least one F508del mutation in the CFTR gene or a mutation that is responsive to Trikafta based on clinical and/or in vitro data</li> </ul> |

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**CYTOKINE & CAM ANTAGONISTS**

| Preferred Agents   | Non-Preferred Agents   | Prior Authorization/Class Criteria  |
|--|--|---|
| <b>IL-6 ANTAGONISTS</b>  |  | <ul style="list-style-type: none"> <li>▪ Preferred agents will be approved with FDA-approved indication – ICD-10 diagnosis code is required.</li> <li>▪ Non-preferred agents will be approved for FDA-approved indications in patients who have failed a trial of TWO preferred agents within this entire drug class, or upon diagnosis for non-preferred agent with FDA-approved indication if no preferred agent has FDA approval for diagnosis.</li> </ul>   |
|  | ACTEMRA (tocilizumab) <b>PEN,SYR</b><br>AVTOZMA (tocilizumab-anoh) <b>AUTOINJ, SYR</b> <sup>AL,NR</sup><br>ENSPRYNG (satralizumab-mwge) <b>SYR</b><br>KEVZARA (sarilumab) <b>PEN, SYR</b> TYENNE (tocilizumab-aazg) <sup>AL</sup>  |   |
| <b>IL-17 ANTAGONISTS</b>   |  |   |
| TALTZ (ixekizumab) <sup>AL</sup> <b>AUTOINJ, SYR</b>   | BIMZELX (bimekizumab-bkzx) <sup>AL</sup> <b>PEN, SYR</b><br>COSENTYX (secukinumab) <sup>AL,QL</sup> <b>PEN, SYR</b>  |   |
| <b>IL-23 ANTAGONISTS</b>   |  | Drug-specific criteria: <ul style="list-style-type: none"> <li>• <b>Cibinqo/Rinvoq/RinvoqER:</b> For diagnosis of moderate to severe Atopic Dermatitis requires treatment failure of TWO preferred systemic drug products, including biologics, within this entire drug class or the Immunomodulators, Atopic Dermatitis therapeutic class.</li> <li>• <b>JAK-Inhibitors:</b> For FDA approved indications that require a patient to have had an inadequate response to a TNF blocker, documentation of an inadequate response to TWO preferred TNF blockers of different active ingredients is required unless preferred products don't have the appropriate indication.</li> <li>• <b>Pyzchiva/Steqeyma:</b> Requires a trial and failure of a preferred TNF blocker with the same indication.</li> </ul> |
| PYZCHIVA (ustekinumab-ttwe, Stelara biosimilar) <sup>AL,CL</sup> <b>SYR</b><br>STEQEYMA (ustekinumab-stba) <sup>AL,CL</sup> <b>SYR</b> | ICOTYDE (icotrokinra) <sup>AL, NR, QL</sup> <b>TAB</b><br>ILUMYA (tildrakizumab) SUB-Q<br>IMULDOSA (ustekinumab-srlf) <sup>AL</sup> <b>SYR</b><br>OMVOH (mirikizumab-mrkz) <sup>AL</sup><br>OTULFI (ustekinumab-aauz) <sup>AL</sup> <b>SYR</b><br>100mg, 200mg,300mg <b>PEN,SYR</b><br>PYZCHIVA (ustekinumab-ttwe, Stelara biosimilar) <sup>AL,CL</sup> <b>VIAL</b><br>SELARSDI (ustekinumab-aekn) <sup>AL</sup> <b>SYR, VIAL</b><br>SKYRIZI (risankizumab-rzaa) <b>SYR</b><br>SKYRIZI (risankizumab-rzaa) <sup>QL</sup> <b>ON-BODY</b><br>SKYRIZI (risankizumab-rzaa) <sup>QL</sup> <b>PEN</b><br>STARJEMZA (ustekinumab-hmny) <sup>AL,NR</sup> <b>SYR</b><br>STARJEMZA (ustekinumab-hmny) <sup>AL,NR</sup> 45mg <b>VIAL</b><br>STELARA (ustekinumab) <sup>AL</sup> <b>SYR</b><br>STELARA (ustekinumab) <sup>AL</sup> <b>VIAL</b><br>STEQEYMA (ustekinumab-stba) <sup>AL,CL,NR</sup> <b>VIAL</b><br>TREMFYA (guselkumab) <b>AUTOINJ</b> <sup>AL,QL</sup> , <b>PEN</b> <sup>AL</sup> , <b>SYR</b> <sup>AL</sup><br>USTEKINUMAB <sup>AL</sup> <b>SYR, VIAL</b><br>USTEKINUMAB-AAUZ <b>SYR</b> (Otulfi biosimilar) <sup>AL,NR</sup><br>USTEKINUMAB-AEKN (biosimilar to Stelara) <sup>AL</sup> <b>SYR</b><br>USTEKINUMAB-TTWE <sup>AL</sup> <b>SYR, VIAL</b> <sup>NR</sup><br>YESINTEK (ustekinumab -kfce) <sup>AL</sup> <b>SYR, VIAL</b> |   |

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**CYTOKINE & CAM ANTAGONISTS, continued**

| Preferred Agents   | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|--|--|--|
| <b>JAK-INHIBITORS<sup>CL</sup></b>                                     |  | <ul style="list-style-type: none"> <li>▪ Preferred agents will be approved with FDA-approved indication – ICD-10 diagnosis code is required.</li> <li>▪ Non-preferred agents will be approved for FDA-approved indications in patients who have failed a trial of TWO preferred agents within this entire drug class, or upon diagnosis for non-preferred agent with FDA-approved indication if no preferred agent has FDA approval for diagnosis.</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>• <b>Cibinqo/Rinvoq/RinvoqER:</b> For diagnosis of moderate to severe Atopic Dermatitis requires treatment failure of TWO preferred systemic drug products, including biologics, within this entire drug class or the Immunomodulators, Atopic Dermatitis therapeutic class.</li> <li>• <b>JAK-Inhibitors:</b> For FDA approved indications that require a patient to have had an inadequate response to a TNF blocker, documentation of an inadequate response to TWO preferred TNF blockers of different active ingredients is required unless preferred products don't have the appropriate indication.</li> <li>• <b>Pyzchiva/Steqeyma:</b> Requires a trial and failure of a preferred TNF blocker with the same indication.</li> </ul> |
|  | CIBINQO (abrocitinib) <sup>AL,CL,QL</sup> <b>TAB</b><br>LEQSELVI (deuruxolitinib) <b>TAB</b><br>LITFULO (ritilecitinib) <sup>AL</sup> <b>CAPS</b><br>OLUMIANT (baricitinib) <sup>QL</sup> <b>TAB</b><br>RINVOQ ER (upadacitinib) <sup>CL,QL</sup> <b>TAB</b><br>RINVOQ (upadacitinib) <sup>AL,CL,QL</sup> LQ <b>SOLN</b><br><b>tofacitinib (generic Xeljanz)<sup>NR,QL</sup> SOLN</b><br><b>tofacitinib (generic Xeljanz)<sup>NR,QL</sup> TAB</b><br><b>tofacitinib ER (generic Xeljanz XR)<sup>NR,QL</sup></b><br>XELJANZ (tofacitinib) <b>SOLN<sup>QL</sup></b><br>XELJANZ (tofacitinib) <b>TAB<sup>QL</sup></b><br>XELJANZ XR (tofacitinib) <b>TAB<sup>QL</sup></b> |  |
| <b>TNF-BLOCKERS</b>  |  |  |
| ADALIMUMAB-ADBM(CF) <sup>AL</sup> 50mg/mL<br><b>KIT, PEN-KIT</b>       | ABRILADA (adalimumab-afzb) <sup>AL</sup> (CF)<br><b>KIT, PEN-KIT</b>   |  |
| ADALIMUMAB-ADBM(CF) <sup>AL</sup> 100mg/mL<br>KIT, PEN-KIT             | ADALIMUMAB-AACF (CF) <sup>AL</sup> <b>PEN* KIT, SYR-KIT</b>  |  |
| ENBREL (etanercept) <b>KIT, MINI CART, PEN, SYR, VIAL<sup>QL</sup></b> | ADALIMUMAB-AATY (CF) <sup>AL</sup> <b>PEN KIT</b>  |  |
| HADLIMA (adalimumab- bwwd) <sup>AL</sup><br><b>PUSHTOUCH, SYR</b>      | ADALIMUMAB-ADAZ(CF)(biosim for Hyrimoz) <sup>AL</sup> <b>KIT, PEN, SYR</b>   |  |
| HADLIMA (CF) (adalimumab- bwwd) <sup>AL</sup><br><b>PUSHTOUCH, SYR</b> | ADALIMUMAB-ADBM(CF) <sup>AL</sup><br>50mg/mL <b>KIT, PEN-KIT</b> (Quallent)  |  |
| SIMLANDI (CF) (adalimumab-ryvk) <sup>AL</sup> <b>PEN KIT</b>           | ADALIMUMAB-ADBM(CF) <sup>AL</sup><br>100mg/mL <b>KIT, PEN-KIT</b> (Quallent)   |  |
|  | ADALIMUMAB-FKJP (biosim for Hulio) <sup>AL</sup><br><b>PEN, SYR</b>  |  |
|  | ADALIMUMAB-RYVK <sup>AL</sup> (biosim for Simlandi) <b>KIT, PEN-KIT</b>  |  |
|  | AMJEVITA (adalimumab-atto) <sup>AL</sup> <b>AUTOINJ, SYR</b>   |  |
|  | AMJEVITA(adalimumab-atto) <sup>AL</sup> <b>KIT, PEN-KIT</b>  |  |
|  | CIMZIA (certolizumab pegol) <sup>QL</sup>  |  |
|  | CYLTEZO (adalimumab-adbm) <sup>AL</sup><br>50mg/mL <b>KIT, PEN-KIT</b>   |  |
|  | CYLTEZO (adalimumab-adbm) <sup>AL</sup><br>100mg/mL <b>KIT, PEN-KIT</b>  |  |

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**CYTOKINE & CAM ANTAGONISTS, continued**

| Preferred Agents                | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|---------------------------------|---|--|
| <b>TNF-BLOCKERS (continued)</b> |   |  |
|                                 | <p>HULIO (adalimumab-fkjp)<sup>AL</sup> <b>PEN, SYR</b><br/> HUMIRA (adalimumab)<sup>QL</sup><br/> HYRIMOZ(CF) (adalimumab-adaz)<sup>AL</sup><br/> <b>PEN, SYR</b><br/> IDACIO (adalimumab-aacf)<sup>AL</sup> <b>PEN, SYR</b><br/> SIMLANDI (CF) (adalimumab-ryvk)<sup>AL</sup> <b>KIT</b><br/> SIMPONI (golimumab)<br/> YUFLYMA 100mg/mL (CF) (adalimumab-aaty)<sup>AL</sup> <b>KIT,PEN-KIT</b><br/> YUFLYMA 80mg/mL (CF) (adalimumab-aaty)<sup>AL</sup> <b>AUTOINJ, PEN, KIT</b><br/> YUSIMRY (CF) (adalimumab-aqvh)<sup>AL</sup> <b>PEN KIT</b><br/> ZYMFENTRA (infliximab-dyyb) <b>PEN, SYR</b></p> | <ul style="list-style-type: none"> <li>▪ Preferred agents will be approved with FDA-approved indication – ICD-10 diagnosis code is required.</li> <li>▪ Non-preferred agents will be approved for FDA-approved indications in patients who have failed a trial of TWO preferred agents within this entire drug class, or upon diagnosis for non-preferred agent with FDA-approved indication if no preferred agent has FDA approval for diagnosis.</li> </ul> <p><b>JAK-Inhibitors:</b> For FDA approved indications that require a patient to have had an inadequate response to a TNF blocker, documentation of an inadequate response to TWO preferred TNF blockers of different active ingredients is required unless preferred products don't have the appropriate indication.</p> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>• <b>Pyzchiva/Steqeyma:</b> Requires a trial and failure of a preferred TNF blocker with the same indication.</li> <li>• <b>Cibinqo/Rinvoq/RinvoqER:</b> For diagnosis of moderate to severe Atopic Dermatitis requires treatment failure of TWO preferred systemic drug products, including biologics, within this entire drug class or the Immunomodulators, Atopic Dermatitis therapeutic class.</li> </ul> |

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**CYTOKINE & CAM ANTAGONISTS, continued**

| Preferred Agents                                   | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|--|--|--|
| <b>MISCELLANEOUS MECHANISM OF ACTION</b>           |  |  |
| <p>OTEZLA (apremilast) <b>TAB</b><sup>QL</sup></p> | <p>ARCALYST (niloncept)<br/>KINERET (anakinra)<br/>ENTYVIO (vedolizumab)<sup>AL</sup> <b>PEN</b><br/>ORENCIA (abatacept) <b>CLICKJET</b><br/>ORENCIA (abatacept) <b>SYR</b><br/>OTEZLA XR (apremilast) <b>TAB</b><br/>SOTYKTU (deucravacitinib) <b>TAB</b><br/>SPEVIGO (spesolimab-sbzo)<sup>AL</sup> <b>SYR</b><br/>VELSIPITY (etrasimod)<sup>QL</sup> <b>TAB</b></p> | <ul style="list-style-type: none"> <li>▪ Preferred agents will be approved with FDA-approved indication – ICD-10 diagnosis code is required.</li> <li>▪ Non-preferred agents will be approved for FDA-approved indications in patients who have failed a trial of TWO preferred agents within this entire drug class, or upon diagnosis for non-preferred agent with FDA-approved indication if no preferred agent has FDA approval for diagnosis.</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>• <b>Cibinqo/Rinvoq/RinvoqER:</b> For diagnosis of moderate to severe Atopic Dermatitis requires treatment failure of TWO preferred systemic drug products, including biologics, within this entire drug class or the Immunomodulators, Atopic Dermatitis therapeutic class.</li> <li>• <b>JAK-Inhibitors:</b> For FDA approved indications that require a patient to have had an inadequate response to a TNF blocker, documentation of an inadequate response to TWO preferred TNF blockers of different active ingredients is required unless preferred products don't have the appropriate indication.</li> <li>• <b>Pyzchiva/Steqeyma:</b> Requires a trial and failure of a preferred TNF blocker with the same indication.</li> </ul> |

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**DIURETICS**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|--|---|---|
| <b>SINGLE-AGENT PRODUCTS</b>   |   |   |
| amiloride <b>TAB</b><br>bumetanide <b>TAB</b><br>chlorthalidone (generic Diuril) <b>TAB</b><br>furosemide (generic Lasix) <b>SOLN, TAB</b><br>hydrochlorothiazide (generic Microzide) <b>CAPS, TAB</b><br>indapamide <b>TAB</b><br>KERENDIA (finerenone) <b>TAB</b> <sup>CL,QL</sup><br>metolazone <b>TAB</b><br>spironolactone (generic Aldactone) <sup>AL</sup> <b>TAB</b><br>torsemide <b>TAB</b> | <b>BAXFENDY (baxdrostat) TAB</b> <sup>NR,AL,QL</sup><br>CAROSPIR (spironolactone) <sup>AL</sup> <b>SUSP</b><br>ENBUMYST (bumetanide) <sup>NR</sup> <b>NASAL SPRAY</b><br>eplerenone (generic Inspra) <sup>CL</sup> <b>TAB</b><br>ethacrynic acid (generic Edecrin) <b>CAPS</b><br>HEMICLOR (chlorthalidone) <sup>NR</sup> <b>TAB</b><br>INZIRQO (hydrochlorothiazide) <sup>NR,QL</sup> <b>SUSP</b><br>spironolactone (generic Carospir) <sup>AL,CL</sup> <b>SUSP</b><br>THALITONE (chlorthalidone) <b>TAB</b><br>triamterene (generic Dyrenium) | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of <b>TWO</b> preferred agents within this drug class.</li> </ul> <p>Drug Specific Criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>eplerenone</b>: Will be approved with a failed trial or intolerance to spironolactone, a trial with two preferred agents is not required.</li> <li>▪ <b>Kerendia</b>: For diagnosis of chronic kidney disease associated with Type-II diabetes in adults. Also for diagnosis of heart failure in adults with LVEF of 40% or greater.</li> <li>▪ <b>spironolactone suspension</b>: May be approved without trial of a preferred agent if there is a clinical reason why preferred spironolactone solid dosage form cannot be used.</li> </ul> |
| <b>COMBINATION PRODUCTS</b>  |   |   |
| amiloride/HCTZ <b>TAB</b><br>spironolactone/HCTZ <b>TAB</b> (generic Aldactazide)<br>triamterene/HCTZ <b>CAPS, TAB</b> (generic Dyazide, Maxzide)  |   |   |

**ENZYME REPLACEMENT, GAUCHER'S DISEASE**

| Preferred Agents                  | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|-----------------------------------|--|--|
| ZAVESCA (miglustat) <sup>CL</sup> | CERDELGA (eliglustat)<br>miglustat (generic Zavesca) <sup>CL</sup> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents require clinical documentation on why the preferred product within this drug class is not appropriate</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Zavesca/miglustat</b>: Approved for mild to moderate type 1 Gaucher disease for whom enzyme replacement therapy is not a therapeutic option</li> </ul> |

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**EPINEPHRINE, SELF-ADMINISTERED<sup>QL</sup>**

| Preferred Agents  | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|---|---|---|
| AUVI-Q 0.1mg (epinephrine)<br>epinephrine (AUTHORIZED GENERIC<br>Epipen/ Epipen Jr.) <b>AUTOINJ</b><br>EPIPEN (epinephrine) <b>AUTOINJ</b><br>EPIPEN JR. (epinephrine) <b>AUTOINJ</b> | AUVI-Q 0.15mg (epinephrine)<br>AUVI-Q 0.3mg (epinephrine)<br>epinephrine (generic Adrenaclick)<br>epinephrine (generic Epipen/ Epipen<br>Jr.) <b>AUTOINJ</b><br>NEFFY (epinephrine) <b>NASAL</b> <sup>AL,QL</sup> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents require clinical documentation why the preferred products within this drug class are not appropriate</li> </ul> |

**ERYTHROPOIESIS STIMULATING PROTEINS**

| Preferred Agents   | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|--|--|--|
| ARANESP (darbopoetine alfa) <b>DISP<br/>SYR, VIAL</b><br>EPOGEN (rHuEPO)<br>RETACRIT (epoetin alfa-epbx) <i>Pfizer<br/>manufacturer only</i> | PROCRT (rHuEPO)<br>RETACRIT (epoetin alfa-epbx) <i>Vifor<br/>manufacturer only</i><br>VAFSEO (vadadustat) <b>TAB</b> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class</li> </ul> |

**FLUOROQUINOLONES, ORAL**

| Preferred Agents  | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|---|--|--|
| ciprofloxacin <b>TAB</b> (generic Cipro)<br>levofloxacin <b>TAB</b> (generic Levaquin)<br>moxifloxacin (generic Avelox) | BAXDELA (delafloxacin) <sup>CL</sup><br>ciprofloxacin ER<br>ciprofloxacin <b>SUSP</b> (generic Cipro) <sup>CL</sup><br>levofloxacin <b>SOLN</b> <sup>CL</sup><br>ofloxacin <sup>CL</sup> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a 3-day trial of ONE preferred agent within this drug class</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Baxdela:</b> Coverable with documented intolerance or failure of preferred MRSA agents (clindamycin, doxycycline, linezolid, sulfamethoxazole/trimethoprim)</li> <li>▪ <b>Ciprofloxacin/Levofloxacin Suspension:</b> Coverable with documented swallowing disorders</li> <li>▪ <b>Ofloxacin:</b> Trial of preferred not required for diagnoses of Pelvic Inflammatory Disease OR Acute Epididymitis (non-gonorrhea)</li> </ul> |

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**GI MOTILITY, CHRONIC**

| Preferred Agents  | Non-Preferred Agents   | Prior Authorization/Class Criteria  |
|---|--|---|
| LINZESS (linaclotide) <sup>AL,QL</sup><br>lubiprostone (generic Amitiza) <sup>AL,QL</sup> | alosetron (generic Lotronex) <sup>CL</sup><br>AMITIZA (lubiprostone) <sup>AL, QL</sup><br>IBSRELA (tenapanor) <sup>AL,CL,QL</sup><br>MOTEGRITY (prucalopride succinate)<br>MOVANTIK (naloxegol oxalate) <sup>QL</sup><br>prucalopride (generic Motegrity)<br>SYMPROIC (naldemedine) <sup>CL</sup><br>VIBERZI (eluxodoline) <sup>CL</sup> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a 30-day trial of ONE preferred agent within this drug class with the same indication</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Lotronex/ alosetron:</b> Covered for diagnosis of IBS Diarrhea Predominant type with trial and failure of loperamide AND diphenoxylate</li> <li>▪ <b>Viberzi®:</b> Covered for diagnosis of IBS Diarrhea Predominant type with trial and failure of loperamide AND diphenoxylate</li> </ul> |

**GLUCAGON AGENTS**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|--|---|--|
| BAQSIMI (glucagon) <sup>AL,QL</sup> <b>NASAL</b><br>GLUCAGON EMERGENCY<br>(glucagon) <sup>QL</sup> <b>INJ KIT</b> (Amphastar)<br>GLUCAGON EMERGENCY<br>(glucagon) <sup>QL</sup> <b>INJ KIT</b> (Fresenius)<br>GLUCAGON EMERGENCY<br>(glucagon) <sup>QL</sup> <b>INJ KIT</b> (Mylan)<br>glucagon <sup>QL</sup> <b>INJ</b><br>GVOKE (glucagon) <sup>AL,QL</sup> <b>PEN, SYR</b><br>PROGLYCEM (diazoxide) <b>SUSP</b><br>ZEGALOGUE (dasiglucagon) <sup>AL, QL</sup> | diazoxide <b>SUSP</b> (generic Proglycem)<br>GLUCAGON EMERGENCY<br>(glucagon) <sup>NR</sup> <b>INJ KIT</b> (Lupin)<br>GVOKE (glucagon) <sup>AL,QL</sup> <b>VIAL</b><br>ZEGALOGUE (dasiglucagon) <sup>AL, QL</sup><br><b>SYR</b> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class</li> </ul> |

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**GLUCOCORTICIDS, INHALED**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|--|---|---|
| <b>GLUCOCORTICIDS</b>  |   | <ul style="list-style-type: none"> <li>▪ Non-preferred agents within the Glucocorticoids and Glucocorticoid/Bronchodilator Combo groups will be approved for patients who have failed a trial of TWO preferred agents within the last 6 months</li> </ul>   |
| ARNUITY ELLIPTA (fluticasone)<br>ASMANEX (mometasone) <sup>QL,AL</sup><br>ASMANEX HFA (mometasone) <sup>QL</sup><br>fluticasone HFA (generic Flovent HFA)  | ALVESCO (ciclesonide) <sup>AL</sup><br>ARMONAIR DIGIHALER (fluticasone) <sup>AL,QL</sup><br>beclomethasone (generic Qvar) <sup>NR</sup><br>fluticasone (generic Flovent Diskus)<br>fluticasone furoate (generic Arnuity Ellipta) <sup>AL</sup><br>QVAR Redihaler (beclomethasone)   |   |
| <b>GLUCOCORTICOID/BRONCHODILATOR COMBINATIONS</b>  |   | Drug-specific criteria: <ul style="list-style-type: none"> <li>▪ <b>budesonide respules:</b> Covered without PA for age ≤ 8 years OR for diagnosis of eosinophilic esophagitis in patients ≥ 9 years, by GI biopsy or upper endoscopy. For other indications, must have failed a trial of two preferred agents within this drug class, within the last 6 months.</li> </ul> |
| ADVAIR DISKUS (fluticasone/salmeterol) <sup>QL</sup><br>ADVAIR HFA (fluticasone/salmeterol) <sup>QL</sup><br>DULERA (mometasone/formoterol)<br>SYMBICORT (budesonide/ formoterol)<br>TRELEGY ELLIPTA (fluticasone/umeclidinium/vilanterol) | AIRDUO DIGIHALER (fluticasone/salmeterol) <sup>AL,QL</sup><br>AIRSUPRA HFA (albuterol and budesonide) <sup>AL</sup><br>BREO ELLIPTA (fluticasone/vilanterol)<br>BREZTRI (budesonide/formoterol/glycopyrrolate) <sup>QL</sup><br>budesonide/formoterol (generic Symbicort)<br>fluticasone/salmeterol (generic Advair Diskus) <sup>QL</sup><br>fluticasone/salmeterol (generic Advair HFA) <sup>QL</sup><br>fluticasone/salmeterol (generic Airduo Respiclick)<br>fluticasone/vilanterol (Breo Ellipta) |   |
| <b>INHALATION SOLUTION</b>   |   |   |
|  | budesonide <b>RESPULES</b> (generic Pulmicort) <sup>CL</sup>  |   |

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**GLUCOCORTICOIDS, ORAL**

| Preferred Agents  | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|---|---|---|
| budesonide EC <b>CAPS</b> (generic Entocort EC)<br>dexamethasone <b>ELIXIR, SOLN</b><br>dexamethasone <b>TAB</b><br>hydrocortisone <b>TAB</b><br>methylprednisolone tablet (generic Medrol)<br>prednisolone <b>SOLN</b><br>prednisolone sodium phosphate<br>prednisone <b>DOSE PAK</b><br>prednisone <b>TAB</b> | ALKINDI (hydrocortisone) <b>GRANULES<sup>AL</sup></b><br>CORTEF (hydrocortisone)<br>cortisone <b>TAB</b><br>dexamethasone <sup>CL</sup> <b>INTENSOL</b><br><b>DEXLYT (dexamethasone)<sup>NR</sup> TAB</b><br>EOHILIA (budesonide) <sup>AL,QL</sup> <b>SUSP</b><br>HEMADY (dexamethasone)<br>KHINDIVI (hydrocortisone) <sup>AL</sup> <b>SOLN</b><br>methylprednisolone 8mg, 16mg, 32mg<br>prednisolone sodium phosphate (generic Millipred/Veripred)<br>prednisolone sodium phosphate <b>ODT</b><br>prednisone <b>SOLN</b><br>prednisone <sup>CL</sup> <b>INTENSOL</b><br>prednisone DR <sup>NR</sup> <b>TAB</b><br>RAYOS DR (prednisone) <b>TAB</b> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class within the last 6 months</li> </ul> Drug-specific criteria: <ul style="list-style-type: none"> <li>▪ <b>Intensol Products:</b> Patient specific documentation of why the less concentrated solution is not appropriate for the patient</li> </ul> |

**GROWTH HORMONES**

| Preferred Agents                                    | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|---|---|---|
| GENOTROPIN (somatropin)<br>NORDITROPIN (somatropin) | HUMATROPE (somatropin)<br>NGENLA (somatrogon-ghla) <sup>AL</sup><br>NUTROPIN AQ (somatropin)<br>OMNITROPE (somatropin)<br>SEROSTIM (somatropin)<br>SKYTROFA (lonapegsomatropin-tcgd)<br>SOGROYA (somapacitan-beco)<br>ZOMACTON (somatropin) | <p style="text-align: center;"><a href="#">Growth Hormone PA Form</a><br/><a href="#">Growth Hormone Criteria</a></p> |

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**H. PYLORI TREATMENTS**

| Preferred Agents  | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|---|---|--|
| PYLERA (bismuth, metronidazole, tetracycline) <sup>QL</sup> | bismuth,metronidazole,tetracycline (generic Pylera) <sup>QL</sup><br>lansoprazole/amoxicillin/clarithromycin (generic Prevpac) <sup>QL</sup><br>OMECLAMOX-PAK (omeprazole, clarithromycin, amoxicillin) <sup>QL</sup><br>TALICIA (omeprazole/amoxicillin/rifabutin)<br>VOQUEZNA (vonoprazan) <sup>CL,QL</sup> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class</li> </ul> <p>Drug-Specific Criteria</p> <ul style="list-style-type: none"> <li>• <b>Voquezna:</b> For the diagnosis of erosive esophagitis or heartburn associated with non-erosive GERD, will require confirmation by initial endoscopy and a trial/failure or a contraindication of two different PPIs (8 weeks each) up to maximally indicated doses in the past 180days. Length of therapy for erosive esophagitis is 240 days max per calendar year and 4 weeks for heartburn associated with non-erosive GERD.</li> </ul> |

**HAE TREATMENTS<sup>CL</sup>**

| Preferred Agents  | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|---|---|--|
| BERINERT (C1 esterase inhibitor, human) <b>INTRAVENOUS</b>            | ANDEMBRY (garadacimab) <sup>AL,NR,QL</sup> <b>AUTOINJECTOR</b>                              | <p style="text-align: center;"><a href="#">HAE Treatments PA Form</a></p> <ul style="list-style-type: none"> <li>• All agents require documentation of diagnosis of Type I or Type II HAE and deficient or dysfunctional C1 esterase inhibitor enzyme. Concomitant use with ACE inhibitors, NSAIDs, or estrogen-containing products is contraindicated</li> <li>• Non-preferred agents will be approved for patients who have a failed trial or a contraindication to ONE preferred agent within this drug class with the same indication.</li> </ul> <p>Drug-Specific Criteria</p> <ul style="list-style-type: none"> <li>• <b>Cinryze, Haegarda, Orladeyo, and Takhzyro,</b> require a history of two or more HAE attacks monthly</li> </ul> |
| HAEGARDA (C1 esterase inhibitor, human) <sup>AL,CL</sup> <b>SUB-Q</b> | CINRYZE (C1 esterase inhibitor, human) <sup>AL,CL,QL</sup> <b>INTRAVENOUS</b>               |  |
| icatibant acetate (generic FIRAZYR) <sup>AL</sup> <b>SUB-Q</b>        | DAWNZERA <sup>AL,NR,QL</sup> (donidalorsen)   |  |
| TAKHZYRO (lanadelumab-flyo) <sup>AL,CL,QL</sup> <b>SYRINGE</b>        | EKTERLY (sebetralstat) <sup>AL,NR</sup> <b>TAB</b>  |  |
|   | FIRAZYR (icatibant acetate) <sup>AL</sup> <b>SUB-Q</b>                                      |  |
|   | ORLADEYO (berotralstat) <b>CAP</b> <sup>AL,CL,QL</sup> <b>PELLET</b> <sup>AL,CL,NR,QL</sup> |  |
|   | RUCONEST (recombinant human C1 inhibitor) <sup>AL</sup> <b>INTRAVENOUS</b>                  |  |
|   | SAJAZIR (icatibant) <sup>AL, NR</sup> <b>SUB-Q</b>  |  |

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**HEMOPHILIA TREATMENTS**

| Preferred Agents  | Non-Preferred Agents                        | Prior Authorization/Class Criteria  |  |
|---|---|---|--|
| <b>BISPECIFIC FACTORS</b>                                       |   | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this entire drug class and with the same indication.</li> </ul> |  |
| HEMLIBRA  |   |   |  |
| <b>FACTOR VIII</b>  |   |   |  |
| ALPHANATE   | ADVATE                                      |   |  |
| HUMATE-P  | ADYNOVATE                                   |   |  |
| KOVALTRY  | AFSTYLA                                     |   |  |
| NOVOEIGHT   | ALTUVIIIIO                                  |   |  |
| NUWIQ   | ELOCTATE                                    |   |  |
| XYNTHA KIT, SOLOFUSE  | ESPEROCT                                    |   |  |
|   | HEMOFIL-M                                   |   |  |
|   | JIVI <sup>AL</sup>                          |   |  |
|   | KOATE-DVI KIT                               |   |  |
|   | KOATE-DVI VIAL                              |   |  |
|   | KOGENATE FS                                 |   |  |
|   | OBIZUR                                      |   |  |
|   | RECOMBINATE                                 |   |  |
| <b>FACTOR IX</b>  |   |   |  |
| ALPROLIX  | ALPHANINE SD                                |   |  |
| BENEFIX   | IDELVION                                    |   |  |
|   | IXINITY                                     |   |  |
|   | PROFILNINE SD                               |   |  |
|   | REBINYN                                     |   |  |
|   | RIXUBIS                                     |   |  |
| <b>FACTOR VIIa AND PROTHROMBIN COMPLEX-PLASMA DERIVED</b>       |   |   |  |
| NOVOSEVEN RT  | FEIBA NF                                    |   |  |
|   | SEVENFACT <sup>AL</sup>                     |   |  |
| <b>FACTOR X AND XIII PRODUCTS</b>                               |   |   |  |
| COAGADEX  | TRETTEN                                     |   |  |
| CORIFACT  |   |   |  |
| <b>TISSUE FACTOR PATHWAY INHIBITOR/ ANTITHROMBIN INHIBITORS</b> |   |   |  |
|   | ALHEMO <sup>AL</sup>                        |   |  |
|   | HYMPAVZI <sup>AL</sup>                      |   |  |
|   | QFITLIA (fitusiran) <sup>AL</sup> PEN, VIAL |   |  |
| <b>VON WILLEBRAND PRODUCTS</b>                                  |   |   |  |
| WILATE  | VONVENDI                                    |   |  |

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**HEPATITIS B AND RELATED AGENTS**

| Preferred Agents     | Non-Preferred Agents   | Prior Authorization/Class Criteria  |
|----------------------|--|---|
| entecavir <b>TAB</b> | adefovir dipivoxil<br>BARACLUDE (entecavir) <b>SOLN,</b><br><b>TAB</b><br>HEPCLUDEX (bulevirtide) <b>SUB-Q<sup>NR</sup></b><br>lamivudine hbv <b>TAB</b><br>VEMLIDY (tenofovir alafenamide fumarate) | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class</li> </ul> Drug Specific Criteria <ul style="list-style-type: none"> <li>▪ <b>tenofovir disoproxil fumarate (generic Viread) tablet:</b> Diagnosis for use required. May be indicated for chronic hepatitis B or HIV-1 infection.                             <ul style="list-style-type: none"> <li>○ See HIV/AIDS class for drug listing and placement</li> </ul> </li> </ul> |

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**HEPATITIS C TREATMENTS**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|--|---|--|
| <b>DIRECT ACTING ANTI-VIRAL</b>  |   | <a href="#">Hepatitis C Treatments PA Form</a><br><a href="#">Hepatitis C Criteria</a> <ul style="list-style-type: none"> <li>▪ Non-preferred products require trial of preferred agents within the same subclass and/or will only be considered with documentation of why the preferred product within this drug class is not appropriate for patient</li> <li>▪ Patients newly eligible for Medicaid will be allowed to complete treatment with the original that treatment was initially authorized by another payor</li> </ul> |
| MAVYRET (glecaprevir/pibrentasvir)<br><b>TAB<sup>CL</sup>, PELLETT<sup>AL,CL</sup></b><br>sofosbuvir/velpatasvir (generic Epclusa) <sup>CL</sup><br>VOSEVI (sofosbuvir/velpatasvir/voxilaprevir) <sup>CL</sup> | HARVONI 200/45MG (ledipasvir/sofosbuvir) <sup>CL</sup> <b>TAB</b><br>HARVONI (ledipasvir/sofosbuvir) <sup>CL</sup> <b>PELLET</b><br>ledipasvir/sofosbuvir (generic Harvoni) <sup>CL</sup><br>SOVALDI (sofosbuvir) <sup>CL</sup> <b>PELLET</b><br>SOVALDI <b>TAB</b> (sofosbuvir) <sup>CL</sup><br>ZEPATIER (elbasvir/grazoprevir) <sup>CL</sup> |  |
| <b>RIBAVIRIN</b>   |   |  |
| ribavirin 200mg <b>CAPSULE, TAB</b>  |   | Drug-specific criteria:<br>Trial with with a preferred agent not required in the following: <ul style="list-style-type: none"> <li>▪ <b>Harvoni/ ledipasvir-sofosbuvir:</b> <ul style="list-style-type: none"> <li>○ Post liver transplant for genotype 1 or 4</li> </ul> </li> <li>▪ <b>Vosevi:</b> Requires documentation of non-response after previous treatment course of Direct Acting Anti-viral agent (DAA) for genotype 1-6 without cirrhosis or with compensated cirrhosis</li> </ul>                                    |
| <b>INTERFERON</b>  |   |  |
| PEGASYS (pegylated interferon alfa-2a) <sup>CL</sup>   |   |  |

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**HISTAMINE II RECEPTOR BLOCKERS**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|--|---|--|
| famotidine <b>TAB</b> (generic Pepcid)<br>famotidine <b>SUSP</b> | cimetidine (generic Tagamet) <sup>CL</sup><br><b>SOLN, TAB</b><br>famotidine <b>CHEW-TAB</b><br>nizatidine <b>CAPS</b> (generic Axid)<br>ranitidine <b>SYRUP, TAB</b> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class</li> </ul> Drug-specific criteria: <ul style="list-style-type: none"> <li>▪ <b>Cimetidine:</b> Approved for viral M. contagiosum or common wart V. Vulgaris treatment</li> </ul> |

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**HIV / AIDS<sup>CL</sup>**

| Preferred Agents  | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|---|---|---|
| <b>CAPSID INHIBITOR</b>   |   | <ul style="list-style-type: none"> <li>▪ All agents require:               <ul style="list-style-type: none"> <li>○ Diagnosis of HIV/AIDS required, OR</li> <li>○ Diagnosis of Pre and Post Exposure Prophylaxis</li> </ul> </li> </ul>   |
|   | SUNLENCA (lenacapavir) <sup>QL</sup><br>YEZTUGO (lenacapavir) <sup>NR,QL</sup> <b>TAB</b>   |   |
| <b>CCR5 ANTAGONISTS</b>   |   | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have a diagnosis of HIV/AIDS and patient specific documentation of why the preferred products within this drug class are not appropriate for the patient, including, but not limited to, drug resistance or concomitant conditions not recommended with preferred agents</li> </ul> |
| SELZENTRY <b>SOLN, TAB</b> (maraviroc)  | maraviroc (generic Selzentry)   |   |
| <b>FUSION INHIBITORS</b>  |   | <ul style="list-style-type: none"> <li>▪ Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy</li> </ul>  |
| FUZEON <b>SUB-Q</b> (enfuvirtide) <sup>QL</sup>   |   |   |
| <b>HIV-1 ATTACHMENT INHIBITOR</b>   |   | <ul style="list-style-type: none"> <li>▪ Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy</li> </ul>  |
|   | RUKOBIA ER (fostemsavir) <sup>AL,QL</sup>   |   |
| <b>INTEGRASE STRAND TRANSFER INHIBITORS (INSTIs)</b>  |   | <ul style="list-style-type: none"> <li>▪ Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy</li> </ul>  |
| ISENTRESS (raltegravir) <sup>QL</sup><br>ISENTRESS HD (raltegravir)<br>TIVICAY (dolutegravir)   | TIVICAY PD (dolutegravir)   |   |
| <b>NON-NUCLEOSIDE REVERSE TRANSCRIPTASE INHIBITORS (NNRTIs)</b>   |   | <ul style="list-style-type: none"> <li>▪ Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy</li> </ul>  |
| EDURANT (rilpivirine)<br>efavirenz <b>CAPS, TABLET</b> (generic Sustiva)<br>INTELENCE (etravirine) <sup>QL</sup><br>PIFELTRO (doravirine) <sup>QL</sup>   | etravirine (generic Intelence) <sup>QL</sup><br>nevirapine IR, ER (generic Viramune/Viramune XR)<br>rilpivirine (generic Edurant) <sup>NR</sup><br>SUSTIVA <b>CAPS, TABLET</b> (efavirenz)<br>VIRAMUNE (nevirapine) <b>SUSP</b> |   |
| <b>NUCLEOSIDE REVERSE TRANSCRIPTASE INHIBITORS (NRTIs)</b>  |   | <ul style="list-style-type: none"> <li>▪ Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy</li> </ul>  |
| abacavir <b>SOLN, TABLET</b> (generic Ziagen)<br>EMTRIVA <b>CAPS, SOLN</b> (emtricitabine)<br>lamivudine <b>SOLN, TABLET</b> (generic EpiVir)<br>zidovudine <b>CAPS, SYRUP, TABLET</b> (generic Retrovir) | didanosine DR (generic Videx EC)<br>emtricitabine <b>CAPS</b> (generic Emtriva)<br>EPIVIR (lamivudine)<br>RETROVIR (zidovudine)<br>stavudine <b>CAPS</b> (generic Zerit)<br>ZIAGEN (abacavir)                                   |   |
| <b>NUCLEOTIDE REVERSE TRANSCRIPTASE INHIBITORS (NRTIs)</b>  |   | <ul style="list-style-type: none"> <li>▪ Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy</li> </ul>  |
| tenofovir <b>TABLET</b> (generic Viread)  | VIREAD (tenofovir) <b>POWDER</b>  |   |
| <b>PHARMACOKINETIC ENHANCER</b>   |   | <ul style="list-style-type: none"> <li>▪ Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy</li> </ul>  |
|   | TYBOST (cobicistat) <sup>QL</sup>   |   |

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**HIV / AIDS<sup>CL</sup> (Continued)**

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|--|---|--|
| <b>PROTEASE INHIBITORS</b>   |   | <ul style="list-style-type: none"> <li>▪ All agents require:               <ul style="list-style-type: none"> <li>○ Diagnosis of HIV/AIDS required, OR</li> <li>○ Diagnosis of Pre and Post Exposure Prophylaxis</li> </ul> </li> <li>▪ Non-preferred agents will be approved for patients who have a diagnosis of HIV/AIDS and patient specific documentation of why the preferred products within this drug class are not appropriate for patients, including, but not limited to, drug resistance or concomitant conditions not recommended with preferred agents</li> <li>▪ Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy.</li> </ul> |
| atazanavir <b>CAPS</b> (generic Reyataz)<br>NORVIR (ritonavir) <b>TAB</b><br>PREZISTA (darunavir) <b>TAB</b><br>ritonavir TAB (generic Norvir)   | APTIVUS <b>CAPS, SOLN</b> (tipranavir)<br>CRIXIVAN (indinavir)<br>DARUNAVIR PROPYLENE GLYCOLATE <sup>AL</sup> <b>TAB</b><br>darunavir ethanolate (generic Prezista) <sup>AL</sup> <b>TAB</b><br>fosamprenavir <b>TAB</b> (generic Lexiva)<br>LEXIVA <b>SUSP</b> (fosamprenavir)<br>LEXIVA <b>TAB</b> (fosamprenavir)<br>NORVIR <b>POWDER, SOLN</b> (ritonavir)<br>PREZISTA (darunavir) <b>SUSP</b><br>REYATAZ <b>POWDER</b> (atazanavir)<br>VIRACEPT (nelfinavir) |  |
| <b>COMBINATION PROTEASE INHIBITORS (PIs) or PIs plus PHARMACOKINETIC ENHANCER</b>  |   |  |
| EVOTAZ (atazanavir/cobicistat) <sup>QL</sup><br>lopinavir/ritonavir <b>SOLN, TAB</b><br>(generic Kaletra)  | KALETRA <b>SOLN</b> (lopinavir/ritonavir)<br>KALETRA <b>TAB</b> (lopinavir/ritonavir)<br>PREZCOBIX (darunavir/cobicistat) <sup>QL</sup>   |  |
| <b>COMBINATION NUCLEOS(T)IDE REVERSE TRANSCRIPTASE INHIBITORS</b>  |   |  |
| abacavir/lamivudine (generic Epzicom)<br>CIMDUO (lamivudine/tenofovir) <sup>QL</sup><br>DESCOVY (emtricitabine/tenofovir) <sup>QL</sup><br>emtricitabine/tenofovir (generic Truvada)<br>lamivudine/zidovudine (generic Combivir) | abacavir/lamivudine/zidovudine (generic Trizivir)<br>COMBIVIR (lamivudine/zidovudine)<br>EPZICOM (abacavir sulfate/lamivudine)<br>TRIZIVIR<br>(abacavir/lamivudine/zidovudine)<br>TRUVADA (emtricitabine/tenofovir)   |  |

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**HIV / AIDS<sup>CL</sup> (Continued)**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|--|---|---|
| <b>COMBINATION PRODUCTS – MULTIPLE CLASSES</b>   |   |   |
| BIKTARVY (bictegravir/emtricitabine/tenofovir) <sup>QL</sup><br>COMPLERA (rilpivirine/emtricitabine/tenofovir)<br>DELSTRIGO (doravirine/lamivudine/tenofovir) <sup>QL</sup><br>DOVATO (dolutegravir/lamivudine) <sup>QL</sup><br>efavirenz/emtricitabine/tenofovir (generic Atripla) <sup>CL</sup><br>GENVOYA (elvitegravir/cobicistat/emtricitabine/tenofovir) <sup>QL, AL</sup><br>JULUCA (dolutegravir/rilpivirine) <sup>QL</sup><br>ODEFSEY (emtricitabine/rilpivirine/tenofovir) <sup>QL</sup><br>STRIBILD (elvitegravir/cobicistat/emtricitabine/tenofovir) <sup>QL</sup><br>SYMFI (efavirenz/lamivudine/tenofovir) <sup>QL</sup><br>SYMFI LO (efavirenz/lamivudine/tenofovir) <sup>QL</sup><br>SYMTUZA (darunavir/cobicistat/emtricitabine/tenofovir) <sup>QL</sup><br>TRIUMEQ (dolutegravir/abacavir/lamivudine) | efavirenz/lamivudine/tenofovir (generic Symfi) <sup>QL</sup><br>efavirenz/lamivudine/tenofovir (generic Symfi Lo) <sup>QL</sup><br>IDVYNZO (doravirine/islatravir) <sup>AL, NR</sup><br>rilpivirine/emtricitabine/tenofovir (Complera) <sup>NR</sup><br>TRIUMEQ PD (abacavir/dolutegravir/lamivudine) <b>SUSP</b> | <ul style="list-style-type: none"> <li>▪ All agents require:               <ul style="list-style-type: none"> <li>○ Diagnosis of HIV/AIDS required, OR</li> <li>○ Diagnosis of Pre and Post Exposure Prophylaxis</li> </ul> </li> <li>▪ Non-preferred agents will be approved for patients who have a diagnosis of HIV/AIDS and patient specific documentation of why the preferred products within this drug class are not appropriate for patients, including, but not limited to, drug resistance or concomitant conditions not recommended with preferred agents</li> <li>▪ Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy</li> </ul> |

**HYPOGLYCEMICS, ALPHA-GLUCOSIDASE INHIBITORS**

| Preferred Agents           | Non-Preferred Agents      | Prior Authorization/Class Criteria   |
|----------------------------|---------------------------|--|
| acarbose (generic Precose) | miglitol (generic Glyset) | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class</li> </ul> |

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**HYPOGLYCEMICS, INCRETIN MIMETICS/ENHANCERS**

| Preferred Agents  | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|---|---|---|
| <b>GLUCAGON-LIKE PEPTIDE-1 RECEPTOR AGONIST (GLP-1 RA)<sup>AL,CL,QL</sup></b>   |   | <b>GLP-1 RA Criteria</b>  |
| OZEMPIC (semaglutide) <sup>AL,QL</sup><br>TRULICITY (dulaglutide) <sup>AL,QL</sup><br>VICTOZA (liraglutide) <sup>AL,QL</sup>                                    | BYDUREON BCISE <b>PEN</b> (exenatide) <sup>AL,QL</sup><br>BYETTA (exenatide) <sup>AL,QL</sup> subcutaneous<br>exenatide (generic Byetta) <sup>AL,QL</sup><br>liraglutide (generic Victoza) <sup>AL,QL</sup><br>MOUNJARO (tirzepatide) <sup>AL,QL</sup> <b>PEN</b><br>OZEMPIC (semaglutide) <sup>AL, NR, QL</sup> <b>TAB</b><br>RYBELSUS (semaglutide) <sup>AL,QL</sup> 1.5mg, 3mg, 4mg, 7mg, 9mg, 14mg <b>TAB</b>   | Preferred agents require a diagnosis of Type II diabetes AND a trial and failure or intolerance to metformin <b>OR</b><br>A diagnosis of ASCVD associated with a diagnosis of Type II diabetes (no metformin trial required)<br><br>Non-preferred agents will be approved for patients who have:<br><ul style="list-style-type: none"> <li>• Failed a trial of TWO preferred agents within GLP-1 RA</li> </ul> <b>AND</b><br><ul style="list-style-type: none"> <li>• Diagnosis of diabetes with HbA1C ≥ 7 <b>AND</b></li> <li>• Trial of metformin, or contraindication or intolerance to metformin</li> </ul> |
| <b>INSULIN/GLP-1 RA COMBINATIONS</b>  |   |   |
|   | SOLIQUA (insulin glargine/lixisenatide)<br>XULTOPHY (insulin degludec/liraglutide)  |   |
| <b>AMYLIN ANALOG<sup>CL</sup></b>   |   | <b>Amylin Analog Criteria</b>   |
|   | SYMLIN (pramlintide) subcutaneous   | ALL criteria must be met <ul style="list-style-type: none"> <li>▪ Concurrent use of short-acting mealtime insulin</li> <li>▪ Current therapy compliance</li> <li>▪ No diagnosis of gastroparesis</li> <li>▪ HbA1C ≤ 9% within last 90 days</li> <li>▪ Monitoring of glucose during initiation of therapy</li> </ul>   |
| <b>DIPEPTIDYL PEPTIDASE-4 (DPP-4) INHIBITOR<sup>AL,CL,QL</sup></b>  |   | <b>DPP-4 Inhibitor Criteria</b>   |
| JANUMET (sitagliptin/metformin)<br>JANUMET XR (sitagliptin/metformin)<br>JANUVIA (sitagliptin)<br>JENTADUETO (linagliptin/metformin)<br>TRADJENTA (linagliptin) | alogliptin (generic Nesina)<br>alogliptin/metformin (generic Kazano)<br>alogliptin/pioglitazone (generic Oseni)<br>BRYNOVIN (sitagliptin) <sup>NR,QL</sup> SOLN<br>dapagliflozin/saxagliptin (generic Qtern) <sup>AL,NR, QL</sup><br>GLYXAMBI (empagliflozin/linagliptin)<br>JENTADUETO XR (linagliptin/metformin)<br>KOMBIGLYZE XR (saxagliptin/metformin)<br>linagliptin (generic Tradjenta) <sup>NR</sup><br>ONGLYZA (saxagliptin)<br>QTERN (dapagliflozin/saxagliptin) <sup>AL, QL</sup><br><b>sitagliptin (generic Januvia)<sup>NR</sup></b> | Preferred agents require a diagnosis of Type II diabetes AND a trial and failure or intolerance to metformin.<br><br>Non-preferred DPP-4s will be approved for patients who have failed a trial of ONE preferred agent within the DPP-4 inhibitor class   |

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**HYPOGLYCEMICS, INCRETIN MIMETICS/ENHANCERS (Continued)**

| Preferred Agents | Non-Preferred Agents   | Prior Authorization/Class Criteria |
|------------------|--|------------------------------------|
|                  | <b>DIPEPTIDYL PEPTIDASE-4 (DPP-4) INHIBITOR<sup>AL,CL,QL</sup></b> |                                    |
|                  | saxagliptin (generic Onglyza)                                      |                                    |
|                  | saxagliptin/metformin ER (generic Kombiglyze ER)                   |                                    |
|                  | sitagliptin (generic Zituvio)                                      |                                    |
|                  | sitagliptin/metformin (generic Janumet) <sup>NR</sup>              |                                    |
|                  | sitagliptin/ metformin (Zituvimet)                                 |                                    |
|                  | sitagliptin/ metformin ER (Zituvimet XR) <sup>NR</sup>             |                                    |
|                  | STEGLUJAN (ertugliflozin/sitagliptin)                              |                                    |
|                  | TRIJARDY XR<br>(empagliflozin/linagliptin/metformin)               |                                    |
|                  | ZITUVIMET (sitagliptin/metformin)<br><b>TAB<sup>QL</sup></b>       |                                    |
|                  | ZITUVIMET XR (sitagliptin/ metformin ER) <b>TAB<sup>QL</sup></b>   |                                    |
|                  | ZITUVIO (sitagliptin)  |                                    |

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**HYPOGLYCEMICS, INSULIN AND RELATED DRUGS**

| Preferred Agents  | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|---|--|--|
| <p>HUMULIN (insulin) <b>VIAL</b><br/> HUMULIN 70/30 <b>VIAL</b><br/> HUMULIN U-500 <b>VIAL</b><br/> HUMULIN U-500 <b>PEN</b><sup>CL</sup><br/> HUMULIN OTC <b>PEN</b><br/> HUMULIN 70/30 OTC <b>PEN</b><br/> insulin aspart/insulin aspart protamine <b>PEN, VIAL</b>(generic Novolog Mix)<br/> insulin lispro (generic Humalog) <b>PEN, VIAL, JR KWIKPEN</b><br/> insulin lispro/lispro protamine <b>KWIKPEN</b> (Humalog Mix Kwikpen)<br/> LANTUS SOLOSTAR <b>PEN</b> (insulin glargine)<br/> LANTUS (insulin glargine) <b>VIAL</b></p> | <p>ADMELOG (insulin lispro) <b>PEN, VIAL</b><br/> AFREZZA (regular insulin)<sup>CL</sup><br/> <b>INHALATION</b><br/> APIDRA (insulin glulisine)<br/> <b>SOLOSTAR, VIAL</b><br/> BASAGLAR (insulin glargine, rec)<br/> <b>PEN, TEMPO PEN</b><br/> FIASP (insulin aspart) <b>CARTRIDGE, PEN, VIAL</b><br/> HUMALOG U-100 <b>TEMPO PEN</b><br/> HUMALOG (insulin lispro)<sup>CL</sup> U-200 <b>KWIKPEN</b><br/> HUMALOG (insulin lispro) U-100 <b>CARTRIDGE, PEN, VIAL</b><br/> HUMALOG JR. (insulin lispro) U-100 <b>KWIKPEN</b><br/> HUMALOG MIX <b>VIAL</b> (insulin lispro/lispro protamine)<br/> HUMALOG MIX <b>KWIKPEN</b> (insulin lispro/lispro protamine)<br/> insulin degludec (generic Tresiba) 100U/mL <b>PEN, VIAL</b><br/> insulin degludec (generic Tresiba) 200U/mL <b>PEN</b><br/> insulin glargine <b>PEN, VIAL</b><br/> insulin glargine (Toujeo)<br/> insulin glargine max (Toujeo Max)<br/> insulin glargine-YFGN <b>PEN, VIAL</b> (generic Semglee-YFGN)<br/> LEVEMIR (insulin detemir) <b>PEN, VIAL</b><br/> LYUMJEV <b>KWIKPEN, VIAL</b>(insulin lispro-aabc)<br/> LYUMJEV (insulin lispro-aabc) <b>TEMPO PEN</b></p> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Afrezza®</b>: Approved for T1DM on long-acting insulin with no current history of smoking or chronic lung disease</li> <li>• <b>Humulin® R U-500 Kwikpen</b>: May be approved for patients who require &gt;200 units/day</li> <li>▪ <b>Humalog U-200 Pen</b>: May be approved for patients who require &gt; 100 units/day</li> </ul> |

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**HYPOGLYCEMICS, INSULIN AND RELATED DRUGS, continued**

| Preferred Agents | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|------------------|--|--|
|                  | MERILOG (insulin aspart-szjj) <sup>NR</sup><br><b>SOLOSTAR PEN</b><br>MERILOG (insulin aspart-szjj) <sup>NR</sup><br><b>VIAL</b><br>NOVOLIN (insulin) <b>PEN-OTC, VIAL-OTC</b><br>NOVOLIN 70/30 VIAL (insulin)<br>NOVOLOG (insulin aspart)<br><b>CARTRIDGE, PEN, VIAL</b><br>NOVOLOG MIX (insulin aspart/aspart protamine) <b>PEN, VIAL</b><br>REZVOGLAR (insulin glargine-aglr)<br><b>KWIKPEN</b><br>SEMGLEE (insulin glargine) <b>PEN, VIAL</b><br>SEMGLEE YFGN (insulin glargine)<br><b>PEN, VIAL</b><br>TOUJEO SOLOSTAR (insulin glargine) | <ul style="list-style-type: none"> <li>Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class</li> </ul> |

**HYPOGLYCEMICS, MEGLITINIDES**

| Preferred Agents              | Non-Preferred Agents          | Prior Authorization/Class Criteria   |
|-------------------------------|-------------------------------|--|
| repaglinide (generic Prandin) | nateglinide (generic Starlix) | <ul style="list-style-type: none"> <li>Non-preferred agents will be approved for patients with:<br/>Failure of a trial of ONE preferred agent in another Hypoglycemic class OR<br/>T2DM and inadequate glycemic control</li> </ul> |

**HYPOGLYCEMICS, METFORMINS**

| Preferred Agents                                     | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|--|---|--|
| metformin IR & ER (generic Glucophage/Glucophage XR) | metformin IR 750 mg<br>metformin ER (generic Fortamet/Glumetza) <sup>CL</sup><br>metformin <b>SOLN</b> (generic Riomet) <sup>CL</sup><br>RIOMET ER (metformin ER) <sup>AL</sup> | <ul style="list-style-type: none"> <li><b>Metformin ER (generic Fortamet®)/Glumetza®:</b> Requires clinical reason why generic Glucophage XR® cannot be used</li> <li><b>Metformin solution:</b> Prior authorization not required for age &lt;7 years</li> </ul> |

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**HYPOGLYCEMICS, SGLT2<sup>CL</sup>**

| Preferred Agents  | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|---|---|--|
| FARXIGA (dapagliflozin) <sup>QL</sup><br>JARDIANCE (empagliflozin) <sup>QL</sup><br>SYNJARDY<br>(empagliflozin/metformin) <sup>AL,QL</sup><br>XIGDUO XR (dapagliflozin/metformin) <sup>QL</sup> | BRENZAVVY (bexagliflozin) <sup>NR</sup><br>dapagliflozin <sup>CL,NR,QL</sup> (generic Farxiga)<br>dapagliflozin/metformin <sup>QL</sup> (generic Xigduo)<br>INPEFA (sotagliflozin) <sup>QL</sup> <b>TAB</b><br>INVOKAMET (canagliflozin/metformin) <sup>QL</sup><br>INVOKAMET XR<br>(canagliflozin/metformin) <sup>QL</sup><br>INVOKANA (canagliflozin)<br>SEGLUROMET<br>(ertugliflozin/metformin) <sup>QL</sup><br>STEGLATRO (ertugliflozin) <sup>QL</sup><br>SYNJARDY XR (empagliflozin/metformin) <sup>AL,QL</sup> | Preferred agents require a diagnosis of Type II diabetes AND a trial and failure or intolerance to metformin,<br><b>OR</b><br>A diagnosis of ASCVD or Heart Failure, or Chronic Kidney Disease associated with a diagnosis of Type II diabetes (no metformin trial required)<br><br>▪ Non-preferred agents will be approved for patients who have failed a trial with ONE preferred agent within this drug class<br><br>Drug Specific Criteria:<br>• <b>Farxiga/ dapagliflozin:</b> May be approved for a diagnosis of Heart Failure without a diagnosis of diabetes<br>• May be approved for a diagnosis of chronic kidney disease at risk of progression without a diagnosis of diabetes<br>• <b>Jardiance:</b> May be approved for a diagnosis of Heart Failure without a diagnosis of diabetes |

**HYPOGLYCEMICS, SULFONYLUREAS**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|--|---|--|
| glimepiride 1mg, 2mg, 4mg, 6mg, 8mg<br>(generic Amaryl)<br>glipizide IR & ER (generic Glucotrol/<br>Glucotrol XL)<br>glyburide (generic Diabeta/Glynase) | chlorpropamide<br>glimepiride 3mg (generic Amaryl)<br>glipizide 15mg <sup>NR</sup><br>tolazamide<br>tolbutamide | ▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class |
| SULFONYLUREA COMBINATIONS  |   |  |
| glipizide/metformin<br>glyburide/metformin (generic Glucovance)  |   |  |

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**HYPOGLYCEMICS, TZD**

| Preferred Agents                     | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|--------------------------------------|---|--|
| <b>THIAZOLIDINEDIONES (TZDs)</b>     |   | <ul style="list-style-type: none"> <li>Non-preferred agents will be approved for patients who have failed a trial of the preferred agent within this drug class</li> </ul> |
| pioglitazone (generic Actos)         |   |  |
| <b>TZD COMBINATIONS<sup>CL</sup></b> |   | <ul style="list-style-type: none"> <li><b>Combination products:</b> Require clinical reason why individual ingredients cannot be used</li> </ul>                           |
|                                      | pioglitazone/glimepiride (generic Duetact)<br>pioglitazone/metformin (generic Actoplus Met) |  |

**IDIOPATHIC PULMONARY FIBROSIS**

| Preferred Agents                            | Non-Preferred Agents   | Prior Authorization/Class Criteria  |
|---|--|---|
| pirfenidone (generic Esbriet) <sup>QL</sup> | ESBRIET (pirfenidone) <sup>QL</sup><br>JASCAYD (nerandomilast) <sup>NR</sup><br>nintedanib (generic Ofev) <sup>NR, QL</sup> <b>CAPS</b><br>OFEV (nintedanib esylate) <sup>QL</sup> <b>CAPS</b> | <ul style="list-style-type: none"> <li>Non-preferred agent requires trial of preferred agent within this drug class with the same indication</li> <li>FDA approved indication required – ICD-10 diagnosis code</li> </ul> |

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**IMMUNOMODULATORS, ASTHMA<sup>CL</sup>**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|--|---|--|
| FASENRA (benralizumab) <sup>AL</sup> <b>PEN</b><br>XOLAIR (omalizumab)<br><b>AUTO-INJ<sup>AL,QL</sup>, SYR<sup>AL,QL</sup></b> | EXDENSUR (depemokimab-ulaa) <sup>AL,NR</sup><br><b>SYR</b><br>NUCALA (mepolizumab) <sup>AL</sup> <b>AUTO-INJ,</b><br><b>SYR</b><br>TEZSPIRE (tezepelumab-ekko) <sup>AL,CL</sup><br><b>PEN</b> | <p style="text-align: center;"><a href="#">Immunomodulators Self-Injectable PA Form</a><br/> <a href="#">Immunomodulators: Self-Administered Injectables Criteria</a></p> <ul style="list-style-type: none"> <li>▪ All agents require prior authorization AND an FDA-approved diagnosis for approval</li> <li>▪ Non-preferred agents require a trial of a preferred agent within this drug class with the same indication</li> <li>▪ For asthma indications: All agents must be prescribed by or in consultation with an allergist, immunologist, or pulmonologist</li> <li>▪ Agents listed may have other FDA approved indications, and will be subject to prior authorization</li> </ul> <p><b>Drug Specific Criteria:</b></p> <ul style="list-style-type: none"> <li>▪ <b>Dupixent:</b> (For other indications, see Immunomodulators, Atopic Dermatitis and COPD therapeutic classes). For Eosinophilic Asthma or Corticosteroid Dependent Asthma: Patients must be ages 6 and older. Documentation of moderate to severe asthma with either eosinophils <math>\geq 150</math> + 1 exacerbation OR oral corticosteroid dependency AND prior drug therapy of med-high or max-tolerated inhaled corticosteroid + controller OR max-tolerated inhaled corticosteroid / long-acting beta agonist combo</li> <li>▪ <b>Tezspire Pen:</b> Does not require a trial of a preferred agent for severe asthma that is non-Type 2 (EOS and IgE asthma). Prior authorization still applies for approval (see also <a href="#">Immunomodulators Self-Injectable PA Form</a>)</li> </ul> |

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**IMMUNOMODULATORS, ATOPIC DERMATITIS<sup>AL</sup>**

| Preferred Agents  | Non-Preferred Agents   | Prior Authorization/Class Criteria  |
|---|--|---|
| <p>ADBRY (tralokinumab-ldrm)<sup>AL,CL,QL</sup><br/><b>SUB-Q</b><br/>ADBRY 300mg/2mL<br/>(tralokinumab-ldrm)<sup>AL,CL,QL</sup> <b>AUTOINJ</b><br/>DUPIXENT (dupilumab)<sup>AL,CL</sup> <b>PEN,SYR</b><br/>EBGLYSS (lebrikizumab-lbkz)<sup>AL,CL,QL</sup><br/><b>PEN, SYRINGE</b><br/>ELIDEL (pimecrolimus)<br/>EUCRISA (crisaborole)<sup>CL,QL</sup><br/>pimecrolimus (generic Elidel)<br/>tacrolimus (generic Protopic)</p> | <p>ANZUPGO (delgocitinib)<sup>QL</sup><br/><b>CREAM</b><br/>NEMLUVIO (nemolizumab-ilto)<sup>AL</sup><br/>OPZELURA (ruxolitinib phosphate) •<br/><b>CREAM</b><sup>AL,CL,QL</sup><br/>VTAMA (tapinarof)<sup>AL</sup> <b>CREAM</b><br/>ZORYVE 0.15% (roflumilast)<sup>AL</sup><br/><b>CREAM</b><br/>ZORYVE 0.05% (roflumilast)<sup>AL,NR</sup><br/><b>CREAM</b></p> | <p><a href="#">Immunomodulators Self-Injectable PA Form</a><br/><a href="#">Immunomodulators: Self-Administered Injectables Criteria</a></p> <p>Non-preferred agents require: Trial of a topical steroid AND Trial of one preferred product within this drug class with same indication. Non-preferred biologics also require treatment failure of a preferred biologic with the same indication.</p> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>• <b>Adbry/Ebglyss:</b> May be approved after a trial or failure of a topical corticosteroid AND a topical calcineurin inhibitor</li> <li>• <b>Dupixent:</b> (For other indications, see Immunomodulators, Asthma and COPD therapeutic classes, <b>and links above</b>): <ul style="list-style-type: none"> <li>○ <b>Atopic Dermatitis:</b> May be approved after a minimum of a 14-day trial of a topical corticosteroid AND a 6-week topical calcineurin</li> </ul> </li> <li>• <b>Eucrisa:</b> May be approved after a 30 day trial failure of a preferred topical corticosteroid (TCS) or topical calcineurin inhibitor (TCI) within the past 180 days; Maximum of 300 grams per year</li> <li>• <b>Opzelura:</b> May be approved for a diagnosis of Atopic Dermatitis and after a trial/failure of a topical steroid and trial of a preferred agent</li> </ul> |

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**IMMUNOMODULATORS, TOPICAL**

| Preferred Agents           | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|----------------------------|--|--|
| imiquimod (generic Aldara) | HYFTOR (sirolimus) <sup>AL</sup> <b>GEL</b><br>imiquimod (generic Zyclara)<br>podofilox (generic Condylox) <b>GEL, SOLN</b><br>VEREGEN (sinecatechins) | <ul style="list-style-type: none"> <li>▪ Non-preferred agents require clinical reason why preferred agent within this drug class cannot be used</li> </ul> |

**IMMUNOSUPPRESSIVES, ORAL**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|--|---|--|
| azathioprine (generic Imuran)<br>azathioprine (generic Azasan) <sup>NR</sup><br>cyclosporine, modified (generic Neoral) <b>CAPS</b><br>everolimus (generic Zortress) <sup>AL</sup><br>mycophenolate (generic Cellcept) <b>CAPS, TAB</b><br>mycophenolic acid<br>RAPAMUNE (sirolimus) <b>SOLN</b><br>RAPAMUNE (sirolimus) <b>TAB</b><br>sirolimus (generic Rapamune) <b>SOLN, TAB</b><br>tacrolimus | ASTAGRAF XL (tacrolimus)<br>AZASAN (azathioprine)<br>cyclosporine <b>CAP, SOFTGEL</b><br>cyclosporine, modified (generic Neoral) <b>SOLN</b><br>ENVARSUS XR (tacrolimus)<br>GENGRAF (cyclosporine, modified) <b>CAP, SOLN</b><br>mycophenolate (generic Cellcept) <b>SUSP</b><br>MYFORTIC (mycophenolate sodium)<br>MYHIBBIN (mycophenolate) <sup>AL</sup> <b>SUSP</b><br>PROGRAF (tacrolimus) <b>CAPS, PACKET</b><br>REZUROCK (belumosudil) <sup>AL,QL</sup> <b>TAB</b><br>SANDIMMUNE (cyclosporine) <b>CAPS, SOLN</b><br>TAVNEOS (avacopan) <sup>CL,QL</sup> <b>CAPS</b><br>ZORTRESS (everolimus) <sup>AL</sup> | Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class<br><br><ul style="list-style-type: none"> <li>▪ Patients established on existing therapy will be allowed to continue</li> </ul> Drug Specific Criteria<br><ul style="list-style-type: none"> <li>▪ <b>Tavneos</b> (avacopan)                             <ul style="list-style-type: none"> <li>▪ No trial of a preferred agent required with appropriate FDA indications with concurrent use of standard therapy, including glucocorticoids</li> </ul> </li> </ul> |

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**INTRANASAL RHINITIS AGENTS**

| Preferred Agents                        | Non-Preferred Agents   | Prior Authorization/Class Criteria  |
|---|--|---|
| <b>ANTICHOLINERGICS</b>                 |  | Non-preferred agents will be approved for patients who have failed a 30-day trial of ONE preferred agent within the same subclass.<br><br>Drug-specific criteria: <ul style="list-style-type: none"> <li>▪ <b>mometasone:</b> Prior authorization NOT required for children ≤ 12 years</li> <li>▪ <b>budesonide:</b> Approved for use in Pregnancy (Pregnancy Category B)</li> <li>▪ <b>Xhance:</b> Indicated for treatment of nasal polyps in ≥ 18 years only</li> </ul> |
| <b>ANTI-HISTAMINES</b>                  |  |   |
| azelastine 0.1% (generic Astelin)       | azelastine 0.15% (generic Astepro)<br>azelastine/fluticasone (generic Dymista)<br>olopatadine (generic Patanase)<br>RYALTRIS (olopatadine/mometasone) <sup>AL</sup>  |   |
| <b>CORTICOSTEROIDS</b>                  |  |   |
| fluticasone <b>Rx</b> (generic Flonase) | BECONASE AQ (beclomethasone)<br>budesonide (Rhinocort) <sup>CL</sup> <b>OTC</b><br>flunisolide (generic Nasalide)<br>fluticasone (generic Flonase) <b>OTC</b><br>mometasone (generic Nasonex) <sup>CL</sup> <b>OTC, RX</b><br>NASONEX (mometasone) <b>OTC</b><br>OMNARIS (ciclesonide)<br>QNASL 40 & 80 (beclomethasone)<br>triamcinolone (generic Nasacort) <b>OTC</b><br>XHANCE (fluticasone) <sup>CL</sup><br>ZETONNA (ciclesonide) |   |

**LEUKOTRIENE MODIFIERS**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|--|---|---|
| montelukast (generic Singulair)<br><b>TAB<sup>QL</sup>/CHEWABLE<sup>AL</sup></b> | montelukast <b>GRANULES</b> (generic Singulair) <sup>CL, AL</sup><br>zafirlukast (generic Accolate)<br>zileuton ER (generic Zyflo CR)<br>ZYFLO (zileuton) | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a 30-day trial of the preferred agent within this drug class</li> </ul> Drug-specific criteria: <ul style="list-style-type: none"> <li>▪ <b>montelukast granules:</b><br/>PA not required for age &lt; 2 years</li> </ul> |

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**LINCOSAMIDES / OXAZOLIDINONES / STREPTOGRAMINS**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|--|---|--|
| clindamycin <b>CAPS</b><br>clindamycin palmitate <b>SOLN</b><br>linezolid <b>TAB</b> | CLEOCIN (clindamycin) <b>CAPS</b><br>CLEOCIN PALMITATE (clindamycin)<br>CLEOCIN PEDIATRIC (clindamycin) <sup>NR</sup><br><b>SOLN</b><br>linezolid <b>SUSP</b><br>SIVEXTRO (tedizolid phosphate)<br>ZYVOX (linezolid) <b>SUSP, TAB</b> | <ul style="list-style-type: none"> <li>Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class</li> </ul> |

**LIPOTROPICS, OTHER**

| Preferred Agents   | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|--|--|--|
| <b>BILE ACID SEQUESTRANTS</b>  |  | <ul style="list-style-type: none"> <li>Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within the same subclass.</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li><b>Colesevelam:</b> Trial not required for diabetes control and monotherapy with metformin, sulfonyleurea, or insulin has been inadequate</li> <li><b>Juxtapid/ Kynamro:</b> <ul style="list-style-type: none"> <li>Approved for diagnosis of homozygous familial hypercholesterolemia (HoFH) OR</li> <li>Treatment failure/maximized dosing/contraindication to ALL the following: statins, ezetimibe, niacin, fibric acid derivatives, omega-3 agents, bile acid sequestrants</li> <li>Require faxed copy of REMS PA form</li> </ul> </li> <li><b>Tryngolza:</b> Approved for diagnosis of familial chylomicronemia syndrome and fasting triglycerides equal to or greater than 880 mg/dL within the past 90 days and used in combination with a low-fat diet of 20 gm or less of fat per day</li> </ul> |
| cholestyramine (generic Questran)<br>colestipol <b>TAB</b> (generic Colestid)                | colesevelam (generic Welchol) <sup>CL</sup> <b>TAB, PACKET</b><br>colestipol (generic Colestid) <b>GRANULES</b><br>QUESTRAN LIGHT (cholestyramine) |  |
| <b>TREATMENT OF HOMOZYGOUS FAMILIAL HYPERCHOLESTEROLEMIA</b>                                 |  |  |
|  | JUXTAPID (lomitapide) <sup>CL</sup><br>KYNAMRO (mipomersen) <sup>CL</sup>  |  |
| <b>TREATMENT OF FAMILIAL CHYLOMICRONEMIA SYNDROME (FCS)</b>                                  |  |  |
|  | TRYNGOLZA (olezarsen) <sup>AL,CL,QL</sup> <b>INJ</b><br>REDEMPLO (plozasiran) <b>SYR</b>   |  |
| <b>FIBRIC ACID DERIVATIVES</b>   |  |  |
| fenofibrate (generic Tricor)<br>fenofibrate (generic Lofibra)<br>gemfibrozil (generic Lopid) | fenofibric acid (generic Fibracor/Trilipix)<br>fenofibrate (generic Antara/Fenoglide/<br>Lipofen/Triglide)   |  |
| <b>NIACIN</b>  |  |  |
| niacin ER (generic Niaspan)  | NIACOR (niacin IR)   |  |
| <b>OMEGA-3 FATTY ACIDS</b>   |  |  |
| omega-3 fatty acids (generic Lovaza)   | icosapent (generic Vascepa) <sup>CL</sup><br>omega-3 <b>OTC</b>  |  |
| <b>CHOLESTEROL ABSORPTION INHIBITORS</b>   |  |  |
| ezetimibe (generic Zetia)  | NEXLETOL (bempedoic acid)<br>NEXLIZET (bempedoic acid/<br>ezetimibe) <sup>QL</sup>   |  |

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**LIPOTROPICS, OTHER (Continued)**

| Preferred Agents   | Non-Preferred Agents                                    | Prior Authorization/Class Criteria   |
|--|---|--|
| <b>PROTEIN CONVERTASE SUBTILISIN/KEXIN TYPE 9 (PCSK9)<br/>INHIBITORS</b>                           |   | <p><b>Drug-Specific Criteria</b><br/><b>Praluent and Repatha:</b> May be approved for diagnoses of:</p> <ul style="list-style-type: none"> <li>• Atherosclerotic cardiovascular disease (ASCVD) in adults</li> <li>• Heterozygous familial hypercholesterolemia (HeFH)               <ul style="list-style-type: none"> <li>○ Praluent ≥ 8 years of age</li> <li>○ Repatha ≥ 10 years of age</li> </ul> </li> <li>• Homozygous familial hypercholesterolemia (HoFH)               <ul style="list-style-type: none"> <li>○ Praluent ≥ 18 years of age</li> <li>○ Repatha ≥ 10 years of age</li> </ul> </li> </ul> <p><b>AND</b></p> <ul style="list-style-type: none"> <li>• Trial and failure or intolerance to a statin for 8 continuous weeks</li> <li>• Concurrent use of a maximally tolerated statin must continue, except for statin-induced rhabdomyolysis or a contraindication to a statin</li> <li>• Failure to reach target LDL-C levels:               <ul style="list-style-type: none"> <li>○ ASCVD – &lt; 70 mg/dL</li> <li>○ Very high risk ASCVD- &lt; 55mg/dL</li> <li>○ HeFH – &lt; 100 mg/dL</li> </ul> </li> </ul> |
| PRALUENT (alirocumab) <sup>CL</sup><br>REPATHA (evolocumab) <sup>CL</sup><br><b>SURECLICK, SYR</b> | REPATHA (evolocumab) <sup>CL</sup><br><b>PUSHTRONEX</b> |  |

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**LIPOTROPICS, STATINS**

| Preferred Agents   | Non-Preferred Agents   | Prior Authorization/Class Criteria  |
|--|--|---|
| <b>STATINS</b>   |  | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of TWO preferred agents within this drug class, within the last 12 months</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Altprev®:</b> One of the TWO trials must be IR lovastatin</li> <li>▪ <b>Combination products:</b> Require clinical reason why individual ingredients cannot be used</li> <li>▪ <b>fluvastatin ER:</b> Requires trial of TWO preferred agents AND trial of IR fluvastatin OR clinical reason IR cannot be used</li> <li>▪ <b>simvastatin/ezetimibe:</b> Approved for 3-month continuous trial of ONE standard dose statin</li> </ul> |
| atorvastatin (generic Lipitor) <sup>QL</sup><br>lovastatin (generic Mevacor)<br>pravastatin (generic Pravachol)<br>rosuvastatin (generic Crestor)<br>simvastatin (generic Zocor) | ALTOPREV (lovastatin ER) <sup>CL</sup><br>ATORVALIQ (atorvastatin) <sup>QL</sup> <b>SUSP</b><br>EZALLOR SPRINKLE (rosuvastatin) <sup>QL</sup><br>fluvastatin ER (generic Lescol XL) <sup>CL</sup><br>fluvastatin IR (generic Lescol)<br>LIVALO (pitavastatin) <sup>AL,QL</sup><br>pitavastatin (generic Livalo) <sup>AL,QL</sup><br>ZYPITAMAG (pitavastatin) |   |
| <b>STATIN COMBINATIONS<sup>CL</sup></b>  |  |   |
|  | atorvastatin/amlodipine (generic Caduet)<br>simvastatin/ezetimibe (generic Vytorin) <sup>CL</sup>  |   |

**MACROLIDES AND KETOLIDES, ORAL**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|--|---|--|
| <b>MACROLIDES</b>  |   | <ul style="list-style-type: none"> <li>▪ Non-preferred agents require clinical reason why preferred products within this drug class cannot be used AND ≥ 3-day trial on a preferred product</li> </ul> |
| azithromycin (generic Zithromax)<br>clarithromycin <b>SUSP, TAB</b> (generic Biaxin)<br>E.E.S. <b>SUSP</b> (erythromycin ethylsuccinate) | clarithromycin ER (generic Biaxin XL)<br>E.E.S. <b>TAB</b> (erythromycin ethylsuccinate)<br>ERY-TAB (erythromycin)<br>ERYPED <b>SUSP</b> (erythromycin)<br>erythromycin ethylsuccinate <b>SUSP</b><br>ERYTHROCIN (erythromycin)<br>erythromycin base <b>CAPS, TAB</b> |  |

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**METHOTREXATE**

| Preferred Agents                          | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|---|---|---|
| methotrexate <b>PF VIAL, TABLET, VIAL</b> | JYLAMVO (methotrexate) <b>SOLN</b><br>OTREXUP (methotrexate) <b>SUB-Q</b><br>RASUVO (methotrexate) <b>SUB-Q</b><br>TREXALL (methotrexate) <b>TAB</b><br>XATMEP (methotrexate) <sup>CL</sup> <b>SOLN</b> | Non-preferred agents require a trial of the preferred agent AND will be approved for an FDA-approved indication<br><br>Drug-specific criteria:<br><ul style="list-style-type: none"> <li>▪ <b>Xatmep™</b>: Indicated for pediatric patients only</li> </ul> |

**MOVEMENT DISORDERS<sup>CL</sup>**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|--|---|--|
| AUSTEDO (deutetrabenazine) <sup>AL,CL,QL</sup><br>AUSTEDO XR (deutetrabenazine) <sup>AL,CL,QL</sup><br>AUSTEDO XR Titration Pack (deutetrabenazine) <sup>AL,CL</sup><br>INGREZZA (valbenazine) <sup>AL,CL,QL</sup><br><b>CAPS, SPRINKLES</b><br>tetrabenazine (generic Xenazine) <sup>CL</sup> | INGREZZA (valbenazine) <sup>AL,CL</sup><br><b>INITIATION PACK</b><br>XENAZINE (tetrabenazine) <sup>CL</sup> | All drugs require an FDA approved indication – ICD-10 diagnosis code required.<br><br>Non-preferred agents require a trial and failure of a preferred agent with the same indication or a clinical reason why a preferred agent in this class cannot be used.<br><br>Drug-specific criteria:<br><ul style="list-style-type: none"> <li>▪ <b>Austedo/Austedo XR/Ingrezza</b>: Diagnosis of Tardive Dyskinesia or chorea associated with Huntington’s Disease; Requires a Step through tetrabenazine with the diagnosis of chorea associated with Huntington’s Disease</li> <li>▪ <b>tetrabenazine</b>: Diagnosis of chorea with Huntington’s Disease</li> </ul> |

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**MULTIPLE SCLEROSIS DRUGS**

| Preferred Agents  | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|---|---|--|
| AVONEX (interferon beta-1a) <sup>QL</sup><br>COPAXONE 20mg (glatiramer) <sup>QL</sup><br>dimethyl fumarate (generic Tecfidera)<br>fingolimod (generic Gilenya) <sup>QL</sup><br>KESIMPTA (Ofatumumab) <sup>CL,QL</sup><br>teriflunomide (generic Aubagio) <sup>QL</sup> | AUBAGIO (teriflunomide) <sup>QL</sup><br>BAFIERTAM (monomethyl fumarate) <sup>QL</sup><br>BETASERON (interferon β -1b) <sup>QL</sup><br>cladribine (generic Mavenclad) <sup>NR</sup><br>dalfampridine (generic Ampyra) <sup>CL,QL</sup><br>dimethyl fumarate (generic Tecfidera Starter Pack) <b>Starter Pack</b><br>EXTAVIA (interferon β -1b) <sup>QL</sup><br>GILENYA (fingolimod) <sup>QL</sup><br>glatiramer (generic Copaxone) <sup>QL</sup><br>MAVENCLAD (cladribine) <b>TAB</b><br>MAYZENT (siponimod) <sup>QL</sup><br>PLEGRIDY (peginterferon β-1a) <sup>CL,QL</sup><br>PONVORY (ponesimod)<br>REBIF (interferon β -1a) <sup>QL</sup><br>TASCENSO ODT (fingolimod) <b>TAB<sup>AL</sup></b><br>TECFIDERA (dimethyl fumarate)<br>VUMERITY (diroximel) <sup>QL</sup><br>ZEPOSIA (ozanimod) <sup>AL,CL,QL</sup> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of TWO preferred agents within this drug class</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Ampyra/ dalfampridine:</b> Approved for diagnosis of gait disorder associated with MS AND EDSS score ≤ 7</li> <li>▪ <b>Kesimpta:</b> Approved for patients who have failed a trial of a preferred injectable agent within this class</li> <li>▪ <b>Plegridy:</b> Approved for diagnosis of relapsing MS</li> <li>▪ <b>Zeposia:</b> Approved for a diagnosis of relapsing forms of multiple sclerosis (MS) with trial of TWO preferred agents OR a diagnosis of moderately to severely active ulcerative colitis and treatment failure of a preferred adalimumab product.</li> </ul> |

**NITROFURAN DERIVATIVES**

| Preferred Agents   | Non-Preferred Agents                            | Prior Authorization/Class Criteria  |
|--|---|---|
| nitrofurantoin macrocrystals <b>CAPS</b> (generic Macrochantin)<br>nitrofurantoin monohydrate-macrocrystals <b>CAPS</b> (generic Macrobid) | nitrofurantoin <b>SUSP</b> (generic Furadantin) | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of <b>ONE</b> preferred agent within this drug class</li> </ul> |

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**NSAIDs, ORAL**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|--|---|--|
| <b>COX-I SELECTIVE</b>   |   | <ul style="list-style-type: none"> <li>▪ Non-preferred agents within COX-1 SELECTIVE subclass will be approved for patients who have failed no less than 30-day trial of TWO preferred agents within the same subclass.</li> <li>▪ Drug-specific criteria:               <ul style="list-style-type: none"> <li>▪ <b>meclofenamate:</b> Approvable without trial of preferred agents for menorrhagia</li> <li>▪ <b>Sprix/ketoralac Nasal:</b> Approved for patients unable to tolerate, swallow OR absorb oral NSAIDs OR contraindication OR trial of TWO preferred oral NSAIDs</li> </ul> </li> </ul> |
| diclofenac sodium (generic Voltaren)<br>ibuprofen OTC, Rx (generic Advil, Motrin) <b>CHEW, DROPS, SUSP, TAB</b><br>ibuprofen OTC (generic Advil, Motrin) <b>CAPS</b><br>indomethacin (generic Indocin) <b>CAPS</b><br>ketorolac (generic Toradol)<br>meloxicam (generic Mobic) <b>TAB</b><br>nabumetone (generic Relafen)<br>naproxen Rx, OTC (generic Naprosyn)<br>naproxen enteric coated<br>sulindac (generic Clinoril) | diclofenac potassium (generic Cataflam, Zipsor)<br>diclofenac SR (generic Voltaren-XR)<br>diflunisal (generic Dolobid)<br>etodolac & SR (generic Lodine/XL)<br>fenoprofen (generic Nalfon)<br>flurbiprofen (generic Ansaid)<br>ibuprofen/famotidine (generic Duexis)<br>ibuprofen 300mg <b>TAB</b><br>indomethacin ER (generic Indocin)<br>ketoprofen & ER (generic Orudis)<br>ketorolac (generic Sprix Nasal) <sup>CL, QL</sup><br><b>NASAL</b><br>meclofenamate (generic Meclomen) <sup>CL</sup><br>mefenamic acid (generic Ponstel)<br>meloxicam (generic Vivlodex) <sup>QL</sup> <b>CAP</b><br>meloxicam (generic Mobic) <b>SUSP</b><br>naproxen CR (generic Naprelan)<br>naproxen (generic Naprosyn) <b>SUSP</b><br>naproxen sodium (generic Anaprox)<br>naproxen-esomeprazole (generic Vimovo)<br>ORUDIS (ketoprofen) <sup>NR</sup> <b>CAP</b><br>oxaprozin (generic Daypro)<br>piroxicam (generic Feldene)<br>tolmetin (generic Tolectin)<br>ZYBIC (meloxicam) <sup>NR, AL</sup> <b>SUSP</b> |  |

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**NSAIDs, ORAL (Continued)**

| Preferred Agents                                     | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|--|---|---|
| <b>COX-I SELECTIVE (continued)</b>                   |   | <ul style="list-style-type: none"> <li>▪ All combination agents require a clinical reason why individual agents can't be used separately</li> </ul> |
|  | <b>ALL BRAND NAME NSAIDs including:</b><br>DOLOBID (diflunisal) <sup>AL</sup> 250mg <b>TAB</b><br>DUEXIS (ibuprofen/famotidine) <sup>CL</sup><br>NALFON (fenoprofen)<br>RELAFEN DS (nabumetone) |   |
| <b>NSAID/GI PROTECTANT COMBINATIONS<sup>CL</sup></b> |   |   |
|  | diclofenac/misoprostol (generic Arthrotec)  |   |
| <b>COX-II SELECTIVE</b>                              |   |   |
| celecoxib (generic Celebrex)                         | VYSCOXA SUSP (celecoxib) <sup>AL,NR</sup>   |   |

**NSAIDs, TOPICAL**

| Preferred Agents                        | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|---|---|--|
| diclofenac sodium <b>GEL (OTC only)</b> | diclofenac <b>PUMP</b> (generic Pennsaid)<br>diclofenac <b>SOLN</b> (generic Pennsaid)<br>FLECTOR <b>PATCH</b> (diclofenac)<br>LICART <b>PATCH</b> (diclofenac) | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed ONE preferred agent within this drug class AND a clinical reason why patient cannot use oral dosage form.</li> </ul> |

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**ONCOLOGY AGENTS, ORAL, BREAST**

| Preferred Agents  | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|---|---|--|
| <b>CDK 4/6 INHIBITOR</b>  |   | <ul style="list-style-type: none"> <li>▪ Non-preferred agents <b>DO NOT</b> require a trial of a preferred agent, but <b>DO</b> require an FDA-approved indication <b>OR</b> documentation submitted supporting off-label use from current treatment guidelines</li> <li>▪ Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy</li> </ul>   |
|   | IBRANCE (palbociclib)<br>KISQALI (ribociclib)<br>KISQALI FEMARA <b>CO-PACK</b><br>VERZENIO (abemaciclib)  |  |
| <b>CHEMOTHERAPY</b>   |   |  |
| capecitabine (generic Xeloda)<br>cyclophosphamide   | XELODA (capecitabine)   |  |
| <b>HORMONE BLOCKADE</b>   |   | Drug-specific criteria <ul style="list-style-type: none"> <li>▪ <b>anastrozole:</b> May be approved for malignant neoplasm of male breast (male breast cancer)</li> <li>▪ <b>Fareston/toremifene:</b> Require clinical reason why tamoxifen cannot be used</li> <li>▪ <b>letrozole:</b> Approved for diagnosis of breast cancer with day supply greater than 12 – NOT approved for short term use</li> <li>▪ <b>Soltamox:</b> May be approved with documented swallowing difficulty</li> </ul> |
| anastrozole (generic Arimidex) <sup>CL</sup><br>exemestane (generic Aromasin)<br>letrozole (generic Femara) <sup>CL</sup><br>tamoxifen citrate (generic Nolvadex) | INLURIYO (imlunestrant) <sup>NR</sup><br>ORSERDU (elacestrant)<br>SOLTAMOX <b>SOLN</b> (tamoxifen) <sup>CL</sup><br>toremifene (generic Fareston) <sup>CL</sup>   |  |
| <b>OTHER</b>  |   |  |
|   | ITOVEBI (inavolisib)<br>NERLYNX (neratinib)<br>PIQRAY (alpelisib)<br>lapatinib (generic Tykerb)<br>TALZENNA (talazoparib tosylate) <sup>QL</sup><br>TUKYSA (tucatinib) <sup>QL</sup><br>TRUQAP (capivasertib) |  |

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**ONCOLOGY AGENTS, ORAL, HEMATOLOGIC**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|--|---|--|
| <b>ALL</b>   |   | <ul style="list-style-type: none"> <li>▪ Non-preferred agents DO NOT require a trial of a preferred agent, but DO require an FDA-approved indication OR documentation submitted supporting off-label use from current treatment guidelines</li> <br/> <li>▪ Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy</li> <br/> <li>Drug-specific criteria               <ul style="list-style-type: none"> <li>▪ <b>Hydrea®:</b> Requires clinical reason why generic cannot be used</li> <li>▪ <b>Purixan/ mercaptopurine:</b> Prior authorization not required for age &lt;12 or for documented swallowing disorder</li> <li>▪ <b>Xpovio:</b> Indicated for relapsed or refractory multiple myeloma. Requires concomitant therapy with dexamethasone</li> </ul> </li> </ul> |
| mercaptopurine   | PURIXAN (mercaptopurine) <sup>AL,CL</sup><br>mercaptopurine (generic Purixan) <sup>AL,CL</sup><br><b>SUSP</b>   |  |
| <b>AML</b>   |   |  |
|  | DAURISMO (glasdegib maleate) <sup>QL</sup><br>IDHIFA (enasidenib)<br>REZLIDHIA (olutasidenib) <sup>QL</sup><br>RYDAPT (midostaurin)<br>TIBSOVO (ivosidenib) <sup>QL</sup><br>VANFLYTA (quizartinib)<br>XOSPATA (gilteritinib) <sup>QL</sup>   |  |
| <b>CLL</b>   |   |  |
|  | COPIKTRA (duvelisib) <sup>QL</sup><br>IMBRUVICA (ibrutinib)<br>VENCLEXTA (venetoclax)<br>ZYDELIG (idelalisib)   |  |
| <b>CML</b>   |   |  |
| hydroxyurea (generic Hydrea)<br>imatinib (generic Gleevec) | BOSULIF (bosutinib)<br><b>bosutinib (generic Bosulif) TAB<sup>NR</sup></b><br>DANZITEN (nilotinib)<br>dasatinib (generic Sprycel)<br>GLEEVEC (imatinib)<br>HYDREA (hydroxyurea) <sup>CL</sup><br>ICLUSIG (ponatinib)<br>IMKELDI (imatinib)<br>nilotinib HCL (generic Tassigna)<br>nilotinib TARTRATE (generic Dansiten)<br>nilotinib TARTRATE <sup>NR</sup> (AG) <b>CAPS</b><br>PHYRAGO (dasatinib) <sup>NR</sup><br>SCEMBLIX (asciminib)<br>SPRYCEL (dasatinib)<br>TASIGNA (nilotinib) |  |

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**ONCOLOGY AGENTS, ORAL, HEMATOLOGIC, continued**

| Preferred Agents  | Non-Preferred Agents   | Prior Authorization/Class Criteria  |
|---|--|---|
| <b>MPN</b>  |  | <ul style="list-style-type: none"> <li>▪ Non-preferred agents <b>DO NOT</b> require a trial of a preferred agent, but <b>DO</b> require an FDA-approved indication <b>OR</b> documentation submitted supporting off-label use from current treatment guidelines</li> <li>▪ Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy</li> </ul>            |
|   | JAKAFI (ruxolitinib) <sup>AL</sup><br>JAKAFI XR (ruxolitinib) <sup>AL,NR</sup>   |   |
| <b>MYELOMA</b>  |  |   |
| REVLIMID <sup>QL</sup> (lenalidomide)                                 | lenalidomide <sup>QL</sup> (generic Revlimid)<br>NINLARO (ixazomib)<br>pomalidomide (generic Pomalyst) <sup>NR</sup><br>POMALYST (pomalidomide)<br>THALOMID (thalidomide)<br>XPOVIO (70envatini) <sup>CL</sup>   | Drug-specific criteria <ul style="list-style-type: none"> <li>▪ <b>Hydrea®:</b> Requires clinical reason why generic cannot be used</li> <li>▪ <b>Purixan/ mercaptopurine:</b> Prior authorization not required for age ≤12 or for documented swallowing disorder</li> <li>▪ <b>Xpovio:</b> Indicated for relapsed or refractory multiple myeloma. Requires concomitant therapy with dexamethasone</li> </ul> |
| <b>OTHER</b>  |  |   |
| MATULANE (procarbazine)<br>tretinoin (generic Vesanoid) <sup>AL</sup> | BRUKINSA (zanubrutinib) <sup>QL</sup><br>CALQUENCE (acalabrutinib) <sup>QL</sup><br>INREBIC (fedratinib dihydrochloride) <sup>QL</sup><br>INQOVI (decitabine/cedazuridine)<br>OJJAARA (mometotinib)<br>REVUFORJ (revumenib) <b>TAB</b><br>VONJO (pacritinib) <sup>QL</sup><br>ZOLINZA (vorinostat) |   |

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**ONCOLOGY AGENTS, ORAL, LUNG**

| Preferred Agents            | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|-----------------------------|---|---|
| <b>ALK</b>                  |   | <ul style="list-style-type: none"> <li>▪ Non-preferred agents DO NOT require a trial of a preferred agent, but DO require an FDA-approved indication OR documentation submitted supporting off-label use from current treatment guidelines</li> <li>▪ Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy</li> </ul> |
|                             | ALECENSA (alectinib)<br>ALUNBRIG (brigatinib) <sup>QL</sup><br>ENSACOVE (ensartinib) <sup>NR</sup><br>LORBRENA (lorlatinib) <sup>QL</sup><br>ZYKADIA (ceritinib) <b>CAPS, TAB</b>   |   |
| <b>ALK / ROS1 / NTRK</b>    |   |   |
|                             | AUGTYRO (reprotectinib) <b>CAPS</b><br>ROZLYTREK (entrectinib) <sup>QL</sup> <b>CAPS, PELLETS</b><br>XALKORI (crizotinib) <b>CAPS, PELLETS</b>  |   |
| <b>EGFR</b>                 |   |   |
| erlotinib (generic Tarceva) | gefitinib (generic Iressa)<br>GILOTRIF (afatinib)<br>IRESSA (gefitinib)<br>LAZCLUZE (71envatinib)<br>TAGRISSO (71envatinib71)<br>TARCEVA (erlotinib)<br>VIZIMPRO (dacomitinib) <sup>QL</sup>  |   |
| <b>HER2/ TKD</b>            |   |   |
|                             | HERNEXEOS (zongertinib) <sup>NR</sup><br>HYRNUO (sevabertinib) <sup>NR</sup>  |   |
| <b>OTHER</b>                |   |   |
|                             | GAVRETO (pralsetinib) <sup>QL</sup><br>HYCAMTIN (topotecan)<br>KRAZATI (adagrasib)<br>LUMAKRAS (sotrasib) <sup>QL</sup><br>RETEVMO (selpercatinib) <sup>AL</sup><br>TABRECTA (capmatinib) <sup>QL</sup><br>TEPMETKO (tepotinib) <sup>QL</sup> |   |

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**ONCOLOGY AGENTS, ORAL, OTHER**

| Preferred Agents               | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|--------------------------------|---|--|
| temozolomide (generic Temodar) | AVMAPKI-FAKZYNJA (avutometinib/defactinib) <b>Combo-Pack</b><br>AYVAKIT (avapritinib) <sup>AL,QL</sup><br>BALVERSA (erdafitinib)<br>CAPRELSA (vandetanib)<br>COMETRIQ (cabozantinib)<br>FRUZAQLA (fruquintinib) <b>CAPS</b><br>GOMEKLI (mirdametinib) <sup>AL</sup> <b>CAPS, TABS FOR ORAL SUSP</b><br>IWILFIN (eflornithine)<br>JAYPIRCA (pirtobrutinib)<br>KOSELUGO (selumetinib) <sup>AL</sup><br>LIFYORLI (relacorilant) <sup>NR</sup> <b>CAPS</b><br>LONSURF (trifluridine/tipiracil)<br>LYNPARZA (olaparib)<br>LYTGOBI (futibatinib)<br>MODEYSO (dordaviprone) <sup>NR</sup><br>OGSIVEO (nirogacestat) <b>TAB</b><br>PEMAZYRE (pemigatinib) <sup>QL</sup><br>QINLOCK (ripretinib)<br>ROMVIMZA (vimseltinib) <b>CAPS</b><br>RUBRACA (rucaparib)<br>STIVARGA (regorafenib)<br>TAZVERIK (tazemetostat) <sup>AL</sup><br>TURALIO (pexidartinib) <sup>QL</sup><br>VITRAKVI (72envatinib72ib) <b>CAPS, SOLN</b><br>VORANIGO (vorasidenib) <sup>AL</sup> <b>TAB</b><br>ZEJULA (niraparib) <b>TAB</b> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents <b>DO NOT</b> require a trial of a preferred agent, but <b>DO</b> require an FDA-approved indication <b>OR</b> documentation submitted supporting off-label use from current treatment guidelines</li> <li>▪ Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy</li> </ul> |

**ONCOLOGY AGENTS, ORAL, PROSTATE**

| Preferred Agents  | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|---|--|--|
| abiraterone (generic Zytiga) <sup>AL,QL</sup><br>bicalutamide (generic Casodex) | AKEEGA (niraparib/abiraterone)<br>ERLEADA (apalutamide) <sup>QL</sup><br>nilutamide (generic Nilandron)<br>NUBEQA (darolutamide) <sup>QL</sup><br>ORGOVYX (relugolix) <sup>AL</sup><br>XTANDI (enzalutamide) <sup>AL,QL</sup><br>YONSA (abiraterone acetone, submicronized)<br>ZYTIGA (abiraterone) <sup>AL,QL</sup> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents <b>DO NOT</b> require a trial of a preferred agent, but <b>DO</b> require an FDA-approved indication <b>OR</b> documentation submitted supporting off-label use from current treatment guidelines</li> <li>▪ Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy</li> </ul> |

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# Nebraska Medicaid Preferred Drug List with Prior Authorization Criteria

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## ONCOLOGY AGENTS, ORAL, RENAL

| Preferred Agents  | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|---|---|---|
| everolimus (generic Afinitor) <b>TAB</b><br>sunitinib malate (generic Sutent) <b>CAPS</b><br>VOTRIENT (pazopanib) | AFINITOR DISPERZ (everolimus) <sup>C</sup><br>CABOMETYX (cabozantinib)<br>everolimus <b>SUSP</b> (generic Afinitor Disperz)<br>everolimus <b>TAB</b> (generic Yulithira) <sup>NR</sup><br>FOTIVDA (tivozanib)<br>INLYTA (axitinib)<br>LENVIMA (73envatinib)<br>NEXAVAR (sorafenib)<br>pazopanib (generic Votrient) <b>TAB</b><br>sorafenib (generic Nexavar)<br>SUTENT (sunitinib) <b>CAPS</b><br>TORPENZ (generic everolimus) <b>TAB</b><br>WELIREG (belzutifan) <sup>QL</sup> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents DO NOT require a trial of a preferred agent, but DO require an FDA-approved indication OR documentation submitted supporting off-label use from current treatment guidelines</li> <li>▪ Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy</li> </ul> |

## ONCOLOGY AGENTS, ORAL, SKIN

| Preferred Agents                               | Non-Preferred Agents   | Prior Authorization/Class Criteria  |
|--|--|---|
| <b>BASAL CELL</b>                              |  | <ul style="list-style-type: none"> <li>▪ Non-preferred agents DO NOT require a trial of a preferred agent, but DO require an FDA-approved indication OR documentation submitted supporting off-label use from current treatment guidelines</li> </ul> |
|  | ERIVEDGE (vismodegib)<br>ODOMZO (sonidegib)  |   |
| <b>BRAF MUTATION</b>                           |  | <ul style="list-style-type: none"> <li>▪ Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy</li> </ul>  |
| MEKINIST (trametinib)<br>TAFINLAR (dabrafenib) | BRAFTOVI (encorafenib)<br>COTELLIC (cobimetinib)<br>MEKINIST (trametinib) <b>SOLN</b><br>MEKTOVI (binimetinib)<br>OJEMDA (tovorafenib)<br><b>SUSP<sup>AL</sup>, TAB</b><br>TAFINLAR (dabrafenib) <b>SUSP</b><br>ZELBORAF (vemurafenib) |   |

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**OPHTHALMICS, ALLERGIC CONJUNCTIVITIS**

| Preferred Agents  | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|---|---|---|
| ALREX (loteprednol 0.2%)<br>ketotifen <b>OTC</b> (generic Zaditor)<br>olopatadine <b>OTC</b> (Pataday once daily)<br>olopatadine <b>OTC</b> (Pataday twice daily) | ALOCRIL (nedocromil)<br>ALOMIDE (Iodoxamide)<br>azelastine (generic Optivar)<br>BEPREVE (bepotastine besilate)<br>bepotastine besilate (generic Bepreve)<br>epinastine (generic Elestat)<br>LASTACAFT (alcaftadine) <b>OTC</b><br>loteprednol 0.2% (generic Alrex)<br>olopatadine <b>DROPS</b> (generic Pataday)<br>olopatadine 0.1% (generic Patanol)<br>olopatadine <b>OTC</b> (PATADAY XS) <sup>NR</sup><br>PATADAY OTC (olopatadine 0.2%)<br>PATADAY XS (olopatadine 0.7%)<br>ZERVIAE (certirizine) <sup>AL</sup> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of TWO preferred agents within this drug class</li> </ul> |

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**OPHTHALMICS, ANTIBIOTICS**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|--|---|---|
| <b>FLUOROQUINOLONES</b>  |   | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a one-month trial of TWO preferred agents within the same subclass</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Azasite®</b>: Approval only requires trial of erythromycin</li> <li>▪ <b>Natacyn®</b>: Approved for documented fungal infection</li> </ul> |
| ciprofloxacin <b>SOLN</b> (generic Ciloxan)<br>ofloxacin (generic Ocuflox)                 | besifloxacin (generic Besivance) <sup>NR</sup><br>BESIVANCE (besifloxacin)<br>CILOXAN (ciprofloxacin)<br>gatifloxacin 0.5% (generic Zymaxid)<br>moxifloxacin (generic Vigamox)<br>moxifloxacin (generic Moxeza)<br>VIGAMOX (moxifloxacin) |   |
| <b>MACROLIDES</b>  |   |   |
| erythromycin   | AZASITE (azithromycin) <sup>CL</sup>  |   |
| <b>AMINOGLYCOSIDES</b>   |   |   |
| gentamicin <b>SOLN</b><br>tobramycin (generic Tobrex drops)                                | TOBREX <b>OINT</b> (tobramycin)   |   |
| <b>OTHER OPHTHALMIC AGENTS</b>   |   |   |
| bacitracin/polymyxin B (generic Polysporin)<br>polymyxin B/trimethoprim (generic Polytrim) | bacitracin<br>NATACYN (natamycin) <sup>CL</sup><br>neomycin/bacitracin/polymyxin B <b>OINT</b><br>neomycin/polymyxin B/gramicidin<br>sulfacetamide <b>SOLN</b> (generic Bleph-10)<br>sulfacetamide <b>OINT</b>                            |   |

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**OPHTHALMICS, ANTIBIOTIC-STEROID COMBINATIONS**

| Preferred Agents  | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|---|---|---|
| neomycin/polymyxin/dexamethasone (generic Maxitrol)<br>sulfacetamide/prednisolone<br>TOBRADEX <b>OINT</b> (tobramycin and dexamethasone)<br>tobramycin/dexamethasone <b>SUSP</b> (generic TobraDex) | neomycin/bacitracin/poly/HC<br>neomycin/polymyxin/HC<br>TOBRADEX S.T. (tobramycin and dexamethasone)<br>ZYLET (loteprednol, tobramycin) | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of TWO preferred agents within this drug class</li> </ul> |

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**OPHTHALMICS, ANTI-INFLAMMATORIES**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|--|---|---|
| <b>CORTICOSTEROIDS</b>   |   | <ul style="list-style-type: none"> <li>▪ ALL sub-classes unless listed below: Non-preferred agents will be approved for patients who have failed a trial of TWO preferred agents within this drug class</li> <li>▪ <b>NSAID class:</b> Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within the same sub-class</li> </ul> |
| fluorometholone 0.1% (generic FML)<br><b>OINT</b><br>MAXIDEX (dexamethasone)<br>PRED MILD (prednisolone 0.12%)<br>prednisolone acetate 1% (generic Omnipred, Pred Forte) | BYQLOVI (clobetasol) <sup>NR</sup><br>dexamethasone (generic Maxidex)<br>difluprednate (generic Durezol)<br>DUREZOL (difluprednate)<br>FLAREX (fluorometholone)<br>FML (fluorometholone 0.1% <b>SOLN</b> )<br>FML FORTE (fluorometholone 0.25%)<br>INVELTYS (loteprednol etabonate)<br>LOTEMAX <b>OINT, GEL</b> (loteprednol)<br>loteprednol <b>GEL</b> (generic Lotemax Gel)<br>loteprednol 0.5% <b>SOLN</b> (generic Lotemax SOLN)<br>prednisolone sodium phosphate<br>prednisolone sodium phosphate 1% |   |
| <b>NSAID</b>   |   |   |
| diclofenac (generic Voltaren)<br>ketorolac 0.5% (generic Acular)   | ACUVAIL (ketorolac 0.45%)<br>bromfenac 0.09% (generic Bromday)<br>bromfenac (generic Bromsite)<br>bromfenac 0.07% (generic Prolensa)<br>BROMSITE (bromfenac)<br>flurbiprofen (generic Ocufen)<br>ILEVRO (nepafenac 0.3%)<br>ketorolac LS 0.4% (generic Acular LS)<br>NEVANAC (nepafenac)<br>PROLENSA (bromfenac 0.07%)  |   |

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**OPHTHALMICS, DRY EYE AGENTS**

| Preferred Agents   | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|--|--|--|
| RESTASIS (cyclosporine)<br>RESTASIS MULTIDOSE (cyclosporine)<br>XIIDRA (lifitegrast) | CEQUA (cyclosporine) <sup>QL</sup><br>cyclosporine (generic Restasis)<br>EYSUVIS (loteprednol etabonate) <sup>QL</sup><br>MIEBO (perfluorohexyloctane)<br>TRYPTYR (acoltremon) <b>SOLN</b><br>TYRVAYA (varenicline tartrate) <sup>QL</sup><br>VERKAZIA (cyclosporine emulsion)<br>VEVYE (cyclosporine) | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class</li> </ul> |

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**OPHTHALMICS, GLAUCOMA**

| Preferred Agents   | Non-Preferred Agents   | Prior Authorization/Class Criteria  |
|--|--|---|
| <b>MIOTICS</b>   |  | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within the same subclass.</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Rhopressa and Rocklatan:</b> Electronically approved for patients who have a trial of ONE generic agent, within ophthalmic – glaucoma within 180 days</li> </ul> |
| pilocarpine  | VUITY (pilocarpine)  |   |
| <b>SYMPATHOMIMETICS</b>  |  |   |
| ALPHAGAN P (brimonidine 0.15%)<br>brimonidine 0.2% (generic Alphagan)  | ALPHAGAN P (brimonidine 0.1%)<br>apraclonidine (generic Iopidine)<br>brimonidine P 0.15% (generic Alphagan P 0.15%)<br>brimonidine 0.1% (generic Alphagan P 0.1%)  |   |
| <b>BETA BLOCKERS</b>   |  |   |
| levobunolol (generic Betagan)<br>timolol (generic Timoptic)            | betaxolol (generic Betoptic)<br>BETIMOL (timolol)<br>BETOPTIC S (betaxolol)<br>carteolol (generic Ocupress)<br>timolol (generic Betimol)<br>timolol (generic Istalol)<br>timolol (generic Timoptic Ocudose)<br>TIMOPTIC (timolol) OCUDOSE  |   |
| <b>CARBONIC ANHYDRASE INHIBITORS</b>                                   |  |   |
| dorzolamide (generic Trusopt)  | AZOPT (brinzolamide)<br>brinzolamide (generic Azopt)   |   |
| <b>PROSTAGLANDIN ANALOGS</b>   |  |   |
| latanoprost (generic Xalatan)<br>TRAVATAN Z (travoprost)               | bimatoprost (generic Lumigan) <sup>NR</sup> 0.01%<br>bimatoprost (generic Lumigan) 0.03%<br>IYUZEH (latanoprost)<br>OMLONTI (omidenepag isopropyl) <sup>NR</sup><br>tafluprost (generic Zioptan)<br>travoprost (generic Travatan Z)<br>VYZULTA (latanoprostene)<br>XALATAN (latanoprost)<br>ZIOPTAN (tafluprost)<br>ZOLYMBUS GEL (bimatoprost) <sup>NR</sup> |   |
| <b>COMBINATION DRUGS</b>   |  |   |
| COMBIGAN (brimonidine/timolol)<br>dorzolamide/timolol (generic Cosopt) | brimonidine/timolol (generic Combigan)<br>COSOPT (dorzolamide/timolol)<br>dorzolamide/timolol PF (generic Cosopt PF)<br>SIMBRINZA (brinzolamide/brimonidine)<br>YUVEZZI (carbachol/brimonidine tartrate) <sup>NR</sup>   |   |

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**OPHTHALMICS, GLAUCOMA (Continued)**

| Preferred Agents   | Non-Preferred Agents | Prior Authorization/Class Criteria   |
|--|----------------------|--|
| <b>OTHER</b>   |                      | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within the same subclass.</li> </ul> Drug-specific criteria: <ul style="list-style-type: none"> <li>▪ <b>Rhopressa and Rocklatan:</b> Electronically approved for patients who have a trial of ONE generic agent, within the ophthalmic - glaucoma class within 180 days</li> </ul> |
| RHOPRESSA (netarsudil) <sup>CL</sup><br>ROCKLATAN (netarsudil and latanoprost) <sup>CL</sup> |                      |  |

**OPIOID DEPENDENCE TREATMENTS**

| Preferred Agents  | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|---|--|--|
| buprenorphine <sup>AL,QL</sup> <b>SL</b><br>buprenorphine/naloxone <sup>AL,QL</sup> <b>TAB (SL)</b><br>naltrexone <b>TAB</b><br>SUBOXONE (buprenorphine/naloxone) <sup>AL,QL</sup><br><b>FILM</b> | buprenorphine/naloxone <sup>AL,QL</sup> <b>FILM</b><br>lofexidine (generic Lucemyra) <sup>CL,QL</sup><br>LUCEMYRA (lofexidine) <sup>CL,QL</sup><br>ZUBSOLV (buprenorphine/naloxone) <sup>AL,QL</sup> | <p style="text-align: center;"><a href="#">Opioid Dependence Treatment PA Form</a><br/><a href="#">Opioid Dependence Treatment Informed Consent</a></p> <ul style="list-style-type: none"> <li>▪ Non-preferred agents require a treatment failure of a preferred drug or patient-specific documentation of why a preferred product is not appropriate for the patient.</li> </ul> Drug-specific criteria: <ul style="list-style-type: none"> <li>▪ <b>Lucemyra/ lofexidine:</b> Approved for FDA approved indication and dosing per label. Trial of preferred product not required.</li> </ul> |

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**OPIOID-REVERSAL TREATMENTS**

| Preferred Agents  | Non-Preferred Agents   | Prior Authorization/Class Criteria  |
|---|--|---|
| naloxone <b>NASAL(Rx)</b> , <b>VIAL</b><br>NARCAN (naloxone) <b>NASAL (OTC)</b> | KLOXXADO (naloxone) <b>NASAL</b><br>naloxone (generic Narcan) <b>OTC</b><br><b>NASAL</b><br>naloxone (generic Narcan) <b>(Rx) SYR</b><br>NARCAN (naloxone) <b>NASAL (Rx)</b><br>OPVEE (nalmeffene) <sup>AL</sup> <b>NASAL</b><br>REXTOVY (naloxone) <b>NASAL</b><br>ZIMHI (naloxone) <b>SYR</b><br>ZURNAL (nalmeffene injection) <sup>NR</sup> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved with documentation of why preferred products within this drug class are not appropriate for the patient</li> </ul> |

**OTIC ANTI-INFECTIVES & ANESTHETICS**

| Preferred Agents            | Non-Preferred Agents                          | Prior Authorization/Class Criteria   |
|-----------------------------|---|--|
| acetic acid (generic Vosol) | acetic acid/hydrocortisone (generic Vosol HC) | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of the preferred agent within this drug class</li> </ul> |

**OTIC ANTIBIOTICS**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|--|---|--|
| CIPRO HC (ciprofloxacin/<br>hydrocortisone)<br>ciprofloxacin/dexamethasone (generic Ciprodex)<br>neomycin/polymyxin/hydrocortisone (generic Cortisporin)<br>ofloxacin (generic Floxin) | ciprofloxacin<br>ciprofloxacin/fluocinolone (generic Otovel)<br>ciprofloxacin/ hydrocortisone (generic Cipro HC) <sup>NR</sup><br>CORTISPORIN TC (colistin/neomycin thonzonium/hydrocortisone)<br>OTOVEL (ciprofloxacin/fluocinolone) | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class</li> </ul> |

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**PAH (PULMONARY ARTERIAL HYPERTENSION AGENTS), ORAL AND INHALED**

| Preferred Agents  | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|---|--|--|
| ambrisentan (generic Letairis) <b>TAB</b><br>REVATIO (sildenafil) <sup>CL,QL</sup> <b>TAB</b><br>sildenafil (generic Revatio) <sup>CL</sup> <b>SUSP</b><br>tadalafil (generic Adcirca) <sup>CL</sup> <b>TAB</b><br>TRACLEER (bosentan) <b>TAB</b> | ADEMPAS (riociguat) <sup>CL</sup> <b>TAB</b><br>ADCIRCA (tadalafil) <sup>CL</sup> <b>TAB</b><br>bosentan (generic Tracleer) <b>TAB</b> ,<br><b>TAB for SUSPENSION</b> <sup>NR</sup><br>LETAIRIS (ambrisentan) <b>TAB</b><br>LIQREV (sildenafil) <sup>CL</sup> <b>SUSP</b><br>macitentan (generic Opsumit) <sup>NR</sup> <b>TAB</b><br>OPSUMIT (macitentan) <b>TAB</b><br>OPSYNVI (macitentan and tadalafil)<br><b>TAB</b><br>ORENITRAM ER (treprostinil) <b>TAB</b> ,<br><b>TITRATION KIT</b><br>REVATIO (sildenafil) <sup>CL</sup> <b>SUSP</b><br>sildenafil (generic Revatio) <sup>CL</sup> <b>TAB</b><br>TADLIQ (tadalafil) <sup>CL</sup> <b>SUSP</b><br>TRACLEER (bosentan) <b>TAB FOR</b><br><b>SUSPENSION</b><br>TYVASO <b>DPI</b> (treprostinil) <b>INHL</b><br>TYVASO (treprostinil) <b>INHL PWDR</b><br>UPTRAVI (selexipag) <b>TAB</b><br>VENTAVIS (iloprost) <b>INHALATION</b><br>YUTREPIA (treprostinil) <sup>NR</sup> <b>INHAL CAP</b> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class within the last 6 months</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Adcirca/Liqrev/ Revatio/sildenafil tablets and suspension/Tadliq/tadalafil:</b> Approved for diagnosis of Pulmonary Arterial Hypertension (PAH)</li> <li>▪ <b>Adempas®:</b><br/>PAH: Requires clinical reason preferred agent cannot be used<br/>CTEPH: Approved for persistent/recurrent diagnosis after surgical treatment or inoperable CTEPH<br/>NOT for use in Pregnancy</li> <li>▪ <b>Liqrev/ Revatio/Tadliq suspension:</b> Requires clinical reason why preferred sildenafil suspension cannot be used</li> </ul> |

**PANCREATIC ENZYMES**

| Preferred Agents               | Non-Preferred Agents                             | Prior Authorization/Class Criteria  |
|--------------------------------|--|---|
| CREON<br>ZENPEP (pancrelipase) | PERTZYE (pancrelipase)<br>VIOKACE (pancrelipase) | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of TWO preferred agents within this drug class</li> </ul> |

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**PEDIATRIC VITAMIN PREPARATIONS**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|--|---|---|
| CHILD MVI (mvi, ped mvi no. 19/FA, CHILD MVI (mvi, ped mvi no. 19/FA, ped mvi no. 17) <b>OTC CHEW</b>    | DEKAs PLUS <sup>AL, CL</sup><br>FLORAFOL (pedi mv no.257/sodium fluoride) <b>PEDIATRIC DROPS OTC</b>                                      | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of TWO preferred agents within this drug class</li> </ul> <p>Drug specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>DEKAs Plus:</b> Approved for diagnosis of Cystic Fibrosis and does not require a trial of a preferred agent</li> </ul> |
| CHILDREN'S CHEWABLES <b>OTC</b><br>(ped mvi no. 25/FA, ped mvi no. 31 /iron/FA, ped mvi no.17/iron sulf) | FLORAFOL FE PEDIATRIC (ped multivit 257/fluoride/iron) <b>DROPS OTC</b>   |   |
| FLUORIDE/VITAMINS A,C,AND D <b>DROPS</b> (ped mvi A,C,D3 no.21/fluoride)                                 | MULTI-VIT-FLOR (pedi multivit no.228/fluoride, pedi multivit no.219/fluoride) <b>CHEW</b>   |   |
| MULTIVITAMINS W/ FLUORIDE (PEDI MVI NO.2 W-FLUORIDE) <b>DROPS</b>  | PEDI MULTIVIT A,C,AND D3 NO.21 <b>DROPS OTC</b>   |   |
| MULTIVITS W/ IRON & FLUORIDE <b>DROPS</b> (ped mvi no. 45/fluoride/iron)                                 | PEDI MVI NO.22 WITH FLUORIDE <sup>NR</sup> <b>DROPS-OTC</b>   |   |
| PED MVI NO.17 W/ FLUORIDE <b>CHEW</b>  | PEDI MVI NO.242/FLUORIDE <b>CHEW-OTC</b>  |   |
| POLY-VITAMIN (ped mvi no. 212) <b>DROPS OTC</b>  | POLY-VI-FLOR (ped mvi no.217/fluoride, ped mvi no. 205/fluoride) <b>CHEW</b>  |   |
| POLY-VITAMIN W/ IRON (ped mvi no. 207 w/ferrous sulf) <b>DROPS OTC</b>                                   | POLY-VI-FLOR (ped mvi no.213 w/fluoride) <b>DROPS</b>   |   |
| TRI-VITAMIN W/ FLUORIDE<br>(ped mvi A,C, D3 no. 21/fluoride)   | POLY-VI-FLOR W/ IRON (ped mvi no. 205/fluoride/iron) <b>CHEW</b><br><br>POLY-VI-FLOR W/ IRON (ped mvi no. 214/fluoride/iron) <b>DROPS</b> |   |

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**PEDIATRIC VITAMIN PREPARATIONS, continued**

| Preferred Agents | Non-Preferred Agents   | Prior Authorization/Class Criteria  |
|------------------|--|---|
|                  | <p>QUFLORA (ped mvi no.84/fluoride, ped mvi no. 63/fluoride, ped mvi no. 83/fluoride)</p> <p>QUFLORA FE (ped mvi 142/iron/fluoride, ped mvi no. 151/iron/fluoride) <b>CHEW</b></p> <p>QUFLORA (ped mvi no.157/ fluoride) <b>OTC</b></p> <p>SOLUVITA A,C,D WITH FLUORIDE <b>DROPS OTC</b></p> <p>TRI-VI-FLOR (ped mvi A,C,D3 no.38/fluoride) <b>DROPS</b></p> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of TWO preferred agents within this drug class</li> </ul> <p>Drug specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>DEKAs Plus:</b> Approved for diagnosis of Cystic Fibrosis and does not require a trial of a preferred agent</li> </ul> |

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## PENICILLINS

| Preferred Agents   | Non-Preferred Agents                                 | Prior Authorization/Class Criteria   |
|--|--|--|
| amoxicillin <b>CAPS, CHEWABLE TAB, SUSP, TAB</b><br>ampicillin <b>CAPS</b><br>dicloxacillin<br>penicillin VK | PIVYA (pivmecillinam) <sup>AL,CL,NR</sup> <b>TAB</b> | Drug Specific Criteria <ul style="list-style-type: none"> <li><b>Pivya tablets:</b> Approved for treatment of female patients 18 years of age and older with uncomplicated urinary tract infections (uUTI) caused by E. coli, Proteus mirabilis and Staphylococcus saprophyticus.</li> </ul> |

## PHOSPHATE BINDERS

| Preferred Agents  | Non-Preferred Agents   | Prior Authorization/Class Criteria  |
|---|--|---|
| calcium acetate <b>TAB</b><br>sevelamer carbonate (generic Renvela)<br><b>PWD PACK, TAB</b> | AURYXIA (ferric citrate)<br>calcium acetate <b>CAPS</b><br>CALPHRON <b>OTC</b> (calcium acetate)<br>ferric citrate (generic Auryxia) <sup>NR</sup><br>lanthanum (generic FOSRENOL)<br>PHOSLYRA (calcium acetate)<br>RENAGEL (sevelamer HCl)<br>RENVELA (sevelamer carbonate)<br><b>PWD PACK, TAB</b><br>sevelamer HCl (generic Renagel)<br>VELPHORO (sucroferric oxyhydroxide)<br>XPHOZAH (tenapanor) <b>TAB</b> | <ul style="list-style-type: none"> <li>Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class within the last 6 months</li> </ul> |

## PLATELET AGGREGATION INHIBITORS

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|--|---|---|
| aspirin<br>BRILINTA (ticagrelor)<br>clopidogrel (generic Plavix)<br>dipyridamole (generic Persantine)<br>prasugrel (generic Effient) | aspirin/dipyridamole (generic Aggrenox)<br>ticlopidine (generic Ticlid)<br>ticagrelor (generic Brilinta) <sup>NR</sup><br>YOSPRALE (aspirin/omeprazole) | <ul style="list-style-type: none"> <li>Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class OR documented clopidogrel resistance</li> </ul> |

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Additional covered agents can be looked up using the Drug Look-up Tool at:

<https://ne.primetherapeutics.com/drug-lookup>

**PRENATAL VITAMINS**

| Preferred Agents  | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|---|--|--|
| CONCEPT OB<br>CONCEPT DHA<br>FE C/FA<br>PNV 2/IRON B-G SUC-P/FA/OMEGA-3<br>PNV NO.118/IRON FUMARATE/FA <b>CHEW TAB</b><br>PNV NO.15/IRON FUM & PS CMP/FA<br>PNV WITH CA, NO.72/IRON/FA<br>PNV WITH CA, NO.74/IRON/FA <b>OTC</b><br>PNV#16/IRON FUM & PS/FA/OM-3<br>PNV119/IRON FUMARATE/FA/DSS<br>PRENATAL MULTI <b>OTC</b><br>PRENATAL VIT #76/IRON, CARB/FA<br>PRENATAL VIT/FE FUMARATE/FA <b>OTC</b><br>SELECT-OB + DHA<br>STUART ONE <b>OTC</b><br>TRICARE<br>TRINATAL RX 1<br>VITAFOL <b>CHEW TAB</b><br>VITAFOL FE+<br>VITAFOL ULTRA<br>VITAFOL-OB<br>VITAFOL-OB+DHA<br>VITAFOL-ONE | ALTRIXA OB <sup>NR</sup> <b>OTC</b><br>CITRANATAL B-CALM<br>COMPLETENATE <b>CHEW TAB</b><br>ENBRACE HR<br>MARNATAL-F<br>MATRONEX TABLET <b>OTC</b> <sup>NR</sup><br>MULTI-MAC <b>OTC</b><br>NATAL PNV (pvn no.164/iron/folate no.6)<br>NEO-VITAL RX TAB <b>OTC</b><br>NESTABS<br>NESTABS ABC<br>NESTABS DHA<br>NESTABS ONE<br>OB COMPLETE ONE<br>OB COMPLETE PETITE<br>OB COMPLETE PREMIER<br>OB COMPLETE <b>TAB</b><br>OB COMPLETE WITH DHA <b>OTC</b><br>ONE NATAL RX <b>TAB</b> <b>OTC</b> <sup>NR</sup><br>PNV 11-IRON FUM-FOLIC ACID-OM3<br>PNV COMBO#47/IRON/FA #1/DHA<br>PNV W-CA NO.40/IRON FUM/FA CMB NO.1<br>PNV WITH CA NO.68/IRON/FA NO.1/DHA<br>PRENATAL + DHA <b>OTC</b><br>PRENATE AM<br>PRENATE <b>CHEW TAB</b><br>PRENATE DHA<br>PRENATE ELITE<br>PRENATE ENHANCE<br>PRENATE ESSENTIAL<br>PRENATE MINI<br>PRENATE PIXIE<br>PRENATE RESTORE<br>PRENATE STAR<br>PRIMACARE<br>PROVIDA OB<br>SELECT-OB <b>CHEW TAB</b><br>TRISTART DHA<br>VITAFOL NANO<br>WESTGEL DHA | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of or are intolerant to TWO preferred agents within this drug class</li> </ul> |

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**PROTON PUMP INHIBITORS**

| Preferred Agents   | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|--|--|--|
| esomeprazole magnesium (generic Nexium) <b>RX</b> <sup>QL</sup><br>omeprazole (generic Prilosec) <b>RX</b><br>pantoprazole (generic Protonix) <sup>QL</sup><br>PROTONIX <b>SUSP</b> (pantoprazole) | DEXILANT (dexlansoprazole)<br>dexlansoprazole (generic Dexilant)<br>esomeprazole magnesium (generic Nexium) <b>OTC</b> <sup>QL</sup><br>esomeprazole strontium<br>KONVOMEP (omeprazole/sodium bicarb) <b>SUSP</b><br>lansoprazole (generic Prevacid) <sup>QL</sup><br>NEXIUM <b>SUSP</b> (esomeprazole)<br>omeprazole/sodium bicarbonate (generic Zegerid RX)<br>pantoprazole <b>GRANULES</b> <sup>QL</sup><br>rabeprazole (generic Aciphex) | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed an 8-week trial of THREE preferred agents.</li> </ul> <p><b>Pediatric Patients:</b><br/>                     Patients ≤ 4 years of age – No PA required for Prevacid 30mg/ lansoprazole 30mg capsules (used to compound suspensions).</p> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Omeprazole OTC/ Prilosec® OTC:</b> EXCLUDED from coverage<br/>                     Acceptable as trial instead of Omeprazole 20mg</li> <li>▪ <b>Prevacid (lansoprazole) Solutab:</b> may be approved after trial of compounded suspension.<br/>                     Patients ≥ 5 years of age- Only approve non-preferred for GI diagnosis if:                     <ul style="list-style-type: none"> <li>▪ Child can not swallow whole generic omeprazole capsules OR,</li> <li>▪ Documentation that contents of capsule may not be sprinkled in applesauce</li> </ul> </li> </ul> |

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**SEDATIVE HYPNOTICS**

| Preferred Agents  | Non-Preferred Agents   | Prior Authorization/Class Criteria  |
|---|--|---|
| <b>BENZODIAZEPINES</b>  |  |   |
| temazepam 15 mg, 30 mg (generic Restoril)   | estazolam (generic ProSom)<br>quazepam (generic Doral)<br>temazepam (generic Restoril) <sup>CL</sup><br>7.5 mg, 22.5 mg<br>triazolam (generic Halcion)   | <p><b>Benzodiazepines Criteria</b></p> <ul style="list-style-type: none"> <li>▪ Non-preferred agents require a trial of the preferred benzodiazepine agent</li> <li>▪ <b>temazepam 7.5/22.5 mg:</b> Requires clinical reason why 15 mg/30 mg cannot be used</li> </ul>  |
| <b>OTHERS<sup>CL</sup></b>  |  |   |
| eszopiclone (generic Lunesta) <sup>AL</sup><br>zaleplon (generic Sonata)<br>zolpidem (generic Ambien) <sup>CL</sup> | BELSOMRA (suvorexant) <sup>AL,QL</sup><br>DAYVIGO (lemborexant) <sup>AL,QL</sup><br>doxepin (generic Silenor) <sup>CL</sup><br>EDLUAR (zolpidem sublingual)<br>HETLIOZ (tasimelteon) <sup>AL</sup><br>HETLIOZ LQ (tasimelteon)<br><b>SUSP</b> <sup>AL,QL</sup><br>QUVIVIQ (daridorexant) <sup>QL</sup><br>ramelteon (generic Rozerem) <sup>AL</sup><br>tasimelteon (generic HetlioZ) <sup>AL</sup><br>zolpidem <sup>QL</sup> <b>CAP</b><br>zolpidem ER (generic Ambien CR) <sup>CL</sup><br>zolpidem SL (generic Intermezzo) <sup>CL</sup> | <p><b>Others Criteria</b></p> <ul style="list-style-type: none"> <li>▪ Non-preferred agents require a trial of TWO preferred agents in the OTHERS sub-category</li> <li>▪ <b>Silenor/doxepin Tablet:</b> Must meet ONE of the following:                             <ul style="list-style-type: none"> <li>○ Contraindication to all of the preferred oral sedative hypnotics agents in the OTHERS sub-category</li> <li>○ Medical necessity for doxepin dose &lt; 10 mg</li> <li>○ Age greater than 65 years old or hepatic impairment (3 mg dose will be approved if this criterion is met)</li> </ul> </li> <li>▪ <b>zolpidem/zolpidem ER:</b> Maximum daily dose for females: zolpidem 5 mg; zolpidem ER 6.25 mg</li> <li>▪ <b>zolpidem SL:</b> Requires clinical reason why half of zolpidem tablet cannot be used or documented swallowing disorder</li> </ul> |

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**SICKLE CELL ANEMIA TREATMENT<sup>AL</sup>**

| Preferred Agents                   | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|------------------------------------|---|--|
| ENDARI (L-glutamine) <sup>CL</sup> | GLUTAMINE POWD PACK (generic Endari)<br>OXBRYTA (voxelotor) <sup>CL</sup><br>SIKLOS (hydroxyurea) <sup>CL</sup> <b>TAB</b><br>XROMI (hydroxyurea) <sup>CL</sup> <b>SOLN</b> | <p>Drug-Specific Criteria</p> <ul style="list-style-type: none"> <li>▪ <b>Endari:</b> Patient must have documented two or more hospital admissions per year due to sickle cell crisis despite maximum hydroxyurea dosage.</li> <li>▪ <b>Oxbryta:</b> Not indicated for sickle cell crisis. Patient must have had at least one sickle cell-related vaso-occlusive event within the past 12 months; AND baseline hemoglobin is 5.5 g/dL ≤ 10.5 g/dL; AND patient is not receiving concomitant, prophylactic blood transfusion therapy</li> <li>▪ <b>Siklos:</b> May be approved for use in patients ages 2 to 17 years old.</li> <li>▪ <b>Xromi</b> solution: May be approved for use in patients ages 6 months to 2 years old.</li> </ul> |

**SINUS NODE INHIBITORS**

| Preferred Agents | Non-Preferred Agents   | Prior Authorization/Class Criteria  |
|------------------|--|---|
|                  | CORLANOR <b>SOLN, TAB</b> (ivabradine)<br>ivabradine (generic Corlanor) <b>TAB</b> | <ul style="list-style-type: none"> <li>▪ Diagnosis of Chronic Heart Failure with left ventricular ejection fraction less than or equal to 35%, AND</li> <li>▪ Sinus rhythm with resting heart rate greater than or equal to 70 beats per minute, AND</li> <li>▪ On maximally tolerated doses of beta-blockers OR have a contraindication to beta-blocker use</li> </ul> |

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**SKELETAL MUSCLE RELAXANTS**

| Preferred Agents  | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|---|---|--|
| baclofen (generic Lioresal)<br>chlorzoxazone (generic Parafon Forte)<br>cyclobenzaprine (generic Flexeril) <sup>QL</sup><br>methocarbamol (generic Robaxin)<br>tizanidine <b>TAB</b> (generic Zanaflex) | ATMEKSI (methocarbamol) <sup>NR, AL</sup> <b>SUSP</b> ▪<br>baclofen (generic Fleqsuvy) <sup>QL</sup> <b>SUSP</b><br>baclofen (generic Ozobax) <sup>QL</sup> <b>SOLN</b><br>baclofen (generic Ozobax DS) <b>SUSP</b><br>carisoprodol (generic Soma) <sup>CL, QL</sup><br>carisoprodol compound<br>cyclobenzaprine ER (generic Amrix) <sup>CL</sup><br>dantrolene (generic Dantrium) <sup>CL</sup><br>FEXMID (cyclobenzaprine ER)<br>FLEQSUVY (baclofen) <sup>QL</sup> <b>SUSP</b><br>LORZONE (chlorzoxazone) <sup>CL</sup><br>LYVISPAH (baclofen) <sup>QL</sup> <b>GRANULES</b><br>metaxalone (generic Skelaxin)<br>NORGESIC FORTE (orphenadrine/ASA/caffeine)<br>ONTRALFY (tizanidine) <sup>NR</sup> <b>SOLN</b><br>orphenadrine ER<br>PARAFON FORTE (chlorzoxazone)<br>TANLOR (methocarbamol) <b>TAB</b><br>tizanidine <sup>CL</sup> <b>CAPS</b><br>ZANAFLEX (tizanidine) <sup>CL</sup> <b>CAPS, TAB</b> | ▪ Non-preferred agents will be approved for patients who have failed a 1-week trial of TWO preferred agents within this drug class<br><br>Drug-specific criteria:<br>▪ <b>cyclobenzaprine ER:</b> <ul style="list-style-type: none"> <li>○ Requires clinical reason why IR cannot be used</li> <li>○ Approved only for acute muscle spasms</li> <li>○ NOT approved for chronic use</li> </ul> ▪ <b>carisoprodol:</b> <ul style="list-style-type: none"> <li>○ Approved for Acute, musculoskeletal pain - NOT for chronic pain</li> <li>○ Use is limited to no more than 30 days</li> <li>○ Additional authorizations will not be granted for at least 6 months following the last day of previous course of therapy</li> </ul> ▪ <b>dantrolene:</b> Trial NOT required for treatment of spasticity from spinal cord injury<br>▪ <b>Lorzone®:</b> Requires clinical reason why chlorzoxazone cannot be used<br>▪ <b>Soma® 250 mg:</b> Requires clinical reason why 350 mg generic strength cannot be used<br>▪ <b>Zanaflex/tizanidine capsules:</b> Requires clinical reason generic cannot be used |

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**STEROIDS, TOPICAL**

| Preferred Agents   | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|--|--|--|
| <b>LOW POTENCY</b>   |  | <ul style="list-style-type: none"> <li>▪ Low Potency Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class</li> </ul>     |
| DERMA-SMOOTH FS (fluocinolone) hydrocortisone OTC & RX <b>CREAM, LOTION, OINT (Rx only)</b>                                  | alclometasone dipropionate (generic Acloivate)<br>desonide <b>LOTION</b> (generic Desowen)<br>desonide <b>CREAM, OINT</b> (generic Desowen, Tridesilon)<br>fluocinolone 0.01% <b>OIL</b> (generic DERMA-SMOOTH-FS)<br>hydrocortisone/aloe <b>CREAM</b><br>hydrocortisone <b>OTC OINT</b><br>hydrocortisone <b>SOLN</b> (generic Texacort)<br>HYDROXYM (hydrocortisone) <b>GEL</b><br>TEXACORT (hydrocortisone)       |  |
| <b>MEDIUM POTENCY</b>  |  | <ul style="list-style-type: none"> <li>▪ Medium Potency Non-preferred agents will be approved for patients who have failed a trial of TWO preferred agents within this drug class</li> </ul> |
| fluticasone propionate <b>CREAM, OINT</b> (generic Cutivate)<br>mometasone furoate <b>CREAM, OINT, SOLN</b> (generic Elocon) | betamethasone valerate (generic Luxiq)<br>clocortolone (generic Cloderm)<br>fluocinolone acetonide (generic Synalar)<br>flurandrenolide (generic Cordran)<br>fluticasone propionate <b>LOTION</b> (generic Cutivate)<br>hydrocortisone butyrate (generic Locoid)<br>hydrocortisone butyrate/emoll (generic Locoid Lipocream)<br>hydrocortisone valerate (generic Westcort)<br>PANDEL (hydrocortisone probutate 0.1%) |  |

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**STEROIDS, TOPICAL (Continued)**

| Preferred Agents   | Non-Preferred Agents   | Prior Authorization/Class Criteria  |
|--|--|---|
| <b>HIGH POTENCY</b>  |  | <ul style="list-style-type: none"> <li>▪ High Potency Non-preferred agents will be approved for patients who have failed a trial of TWO preferred agents within this drug class</li> </ul>      |
| triamcinolone acetonide <b>OINTMENT, CREAM</b><br>triamcinolone <b>LOTION</b>  | amcinonide <b>CREAM</b><br>betamethasone dipropionate<br>betamethasone / propylene glycol<br>betamethasone valerate<br>clobetasol propionate (generic Impoyz) <b>CREAM</b><br>desoximetasone<br>diflorasone diacetate<br>fluocinonide <b>CREAM, GEL, OINT</b><br>fluocinonide emollient<br>fluocinonide <b>SOLN</b><br>halcinonide <b>CREAM, SOLN</b> (generic Halog)<br>HALOG (halcinonide) <b>CREAM, OINT, SOLN</b><br>KENALOG AEROSOL (triamcinolone)<br>triamcinolone <b>SPRAY</b> (generic Kenalog spray)<br>VANOS (fluocinonide) |   |
| <b>VERY HIGH POTENCY</b>   |  | <ul style="list-style-type: none"> <li>▪ Very High Potency Non-preferred agents will be approved for patients who have failed a trial of TWO preferred agents within this drug class</li> </ul> |
| clobetasol emollient (generic Temovate-E)<br>clobetasol propionate <b>CREAM, OINT, SOLN</b><br>halobetasol propionate (generic Ultravate) <b>CREAM</b> | APEXICON-E (diflorasone)<br>clobetasol <b>LOTION, SHAMPOO</b><br>clobetasol propionate <b>GEL, FOAM, SPRAY</b><br>halobetasol propionate (generic Lexette) <sup>AL,QL</sup> <b>FOAM</b><br>halobetasol propionate (generic Ultravate) <sup>NR</sup> <b>LOTION</b>  |   |

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**STIMULANTS AND RELATED ADHD DRUGS <sup>AL</sup>**

| Preferred Agents  | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|---|---|---|
| <b>CNS STIMULANTS</b>   |   | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this subclass.</li> </ul>   |
| <b>Amphetamine type</b>   |   |   |
| amphetamine salt combination ER (generic Adderall XR)             | ADDERALL XR (amphetamine salt combo)  | <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Procentra/ dextroamphetamine soln:</b> May be approved with documentation of swallowing disorder</li> <li>▪ <b>Zenzedi®:</b> Requires clinical reason generic dextroamphetamine IR cannot be used</li> </ul> |
| amphetamine salt combination IR                                   | ADZENYS XR (amphetamine) <b>ODT</b><br>amphetamine salt combination ER (generic Mydayis) <b>CAP</b> |   |
| DYANAVEL XR (amphetamine) <sup>QL</sup>                           | amphetamine ER ODT <sup>AL, NR</sup> (generic Adzenys XR ODT)                                       |   |
| lisdexamfetamine (generic Vyvanse Chew) <sup>QL</sup> <b>CHEW</b> | amphetamine sulfate (generic Evekeo)  |   |
| lisdexamfetamine (generic Vyvanse) <sup>QL</sup> <b>CAP</b>       | ARYNTA (lisdexamfetamine dimesylate) <sup>AL, NR</sup> <b>SOLN</b>                                  |   |
| VYVANSE (lisdexamfetamine) <sup>QL</sup> <b>CAPS, CHEWABLE</b>    | dextroamphetamine (generic Dexedrine) <b>TAB</b>  |   |
|   | dextroamphetamine (generic Procentra) <sup>CL</sup> <b>SOLN</b>                                     |   |
|   | dextroamphetamine ER (generic Dexedrine ER Spansule) <b>CAPS</b>                                    |   |
|   | EVEKEO ODT (amphetamine sulfate)  |   |
|   | methamphetamine (generic Desoxyn)   |   |
|   | MYDAYIS (amphetamine salt combo) <sup>QL</sup>  |   |
|   | XELSTRYM (detroamphetamine) <sup>QL</sup> <b>PATCH</b>  |   |
|   | ZENZEDI (dextroamphetamine) <sup>CL</sup>   |   |

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**STIMULANTS AND RELATED ADHD DRUGS (Continued)<sup>AL</sup>**

| Preferred Agents  | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|---|---|--|
| <b>Methylphenidate type</b>   |   |  |
| CONCERTA (methylphenidate ER) <sup>QL</sup><br>18 mg, 27 mg, 36 mg, 54 mg | APTENSIO XR (methylphenidate)<br>AZSTARYS (serdexmethylphenidate and dexamethylphenidate) <sup>QL</sup>   | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of TWO preferred agents within this subclass.</li> <li>▪ Maximum accumulated dose of 108mg per day for ages &lt; 18</li> <li>▪ Maximum accumulated dose of 72mg per day for ages &gt; 19</li> <li>Drug-specific criteria:               <ul style="list-style-type: none"> <li>▪ <b>Daytrana/methylphenidate patch:</b> May be approved in history of substance use disorder by parent, caregiver, or patient. May be approved with documentation of difficulty swallowing</li> <li>▪ <b>QuilliChew ER:</b> May be approved for children ≤ 12 years of age OR with documentation of difficulty swallowing</li> </ul> </li> </ul> |
| DAYTRANA <b>PATCH</b><br>(methylphenidate) <sup>CL,QL</sup>               | COTEMPLA XR-ODT<br>(methylphenidate) <sup>QL</sup>  |  |
| dexamethylphenidate (generic Focalin IR)                                  | FOCALIN IR (dexamethylphenidate)<br>FOCALIN XR (dexamethylphenidate)<br>JORNAY PM (methylphenidate) <sup>QL</sup><br>methylphenidate <b>CHEW</b>  |  |
| dexamethylphenidate ER (generic Focalin XR)                               | methylphenidate ER (45 mg and 63 mg) <sup>QL</sup>  |  |
| METHYLIN <b>SOLN</b> (methylphenidate)                                    | methylphenidate 50/50 (generic Ritalin LA)<br>methylphenidate 30/70 (generic Metadate CD)   |  |
| methylphenidate TD24 <sup>AL,CL</sup> <b>PATCH</b><br>(generic Daytrana)  | methylphenidate ER 18 mg, 27 mg, 36 mg, 54 mg (generic Concerta) <sup>QL</sup>  |  |
| methylphenidate (generic Ritalin)   | methylphenidate ER <b>CAP</b> (generic Aptensio XR) <sup>QL</sup>   |  |
| methylphenidate <b>SOLN</b> (generic Methylin)                            | methylphenidate ER (generic Metadate ER)<br>methylphenidate ER 72 mg (generic RELEXXII) <sup>QL</sup>   |  |
| QUILLICHEW ER <b>CHEWTAB</b><br>(methylphenidate) <sup>CL</sup>           | methylphenidate ER (generic Ritalin LA)   |  |
| QUILLIVANT XR<br>(methylphenidate) <b>SUSP</b>                            | methylphenidate TD24 <sup>AL,CL</sup> <b>PATCH (AG)</b><br>(generic Daytrana) <b>Padagis Manf. only</b><br>RELEXXII ER (methylphenidate 45mg and 63mg) <sup>AL,QL</sup> <b>TAB</b><br>RITALIN (methylphenidate) |  |

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| Preferred Agents   | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|--|--|--|
| <b>MISCELLANEOUS</b>   |  | <p><b>Note:</b> generic guanfacine IR and clonidine IR are available without prior authorization</p> <ul style="list-style-type: none"> <li>• Non-preferred agents will be approve for patients who have failed a trial of ONE preferred agent within this subclass</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Wakix and Sunosi:</b> Require trial of armodafinil or modafinil</li> <li>▪ <b>armodafinil and modafinil:</b> approved only for:               <ul style="list-style-type: none"> <li>○ Sleep Apnea with documentation/confirmation via sleep study and documentation that C-PAP has been maxed</li> <li>○ Narcolepsy with documentation of diagnosis via sleep study</li> <li>○ Shift Work Sleep Disorder (only approvable for 6 months) with work schedule verified and documented. Shift work is defined as working the all night shift</li> </ul> </li> <li>▪ <b>Sunosi</b> approved only for:               <ul style="list-style-type: none"> <li>○ Sleep Apnea with documentation/confirmation via sleep study and documentation that C-PAP has been maxed</li> <li>○ Narcolepsy with documentation of diagnosis via sleep study</li> </ul> </li> <li>▪ <b>Wakix:</b> approved only for excessive daytime sleepiness in adults with narcolepsy with documentation of narcolepsy diagnosis via sleep study</li> </ul> |
| atomoxetine (generic Strattera) <sup>QL</sup><br>guanfacine ER (generic Intuniv) <sup>QL</sup><br>QELBREE (viloxazine) <sup>QL</sup> | <b>ATONCY SOLN (atomoxetine)<sup>AL, NR</sup></b><br>clonidine ER (generic Kapvay) <sup>QL</sup><br>INTUNIV (guanfacine)<br>Onyda XR (clonidine suspension, extended release) <sup>QL</sup><br>STRATTERA (atomoxetine) |  |
| <b>ANALEPTICS</b>  |  |  |
|  | armodafinil (generic Nuvigil) <sup>CL</sup><br>modafanil (generic Provigil) <sup>CL</sup><br>SUNOSI (solriamfetol) <sup>CL, QL</sup><br>WAKIX (pitolisant) <sup>CL, QL</sup>   |  |

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**TETRACYCLINES**

| Preferred Agents   | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|--|---|--|
| doxycycline hyclate IR (generic Vibramycin) <b>CAPS</b><br>doxycycline monohydrate (generic Vibramycin) <sup>CL</sup> <b>SUSP</b><br>doxycycline monohydrate <b>TAB</b> (generic Vibramycin)<br>minocycline HCl (generic Dynacin/Myrac) <b>TAB</b><br>tetracycline | demeclocycline (generic Declomycin) <sup>CL</sup><br>DORYX MPC DR (doxycycline pelletized)<br>doxycycline hyclate IR <b>TAB</b> (generic Vibramycin)<br>doxycycline hyclate DR (generic Doryx)<br>doxycycline monohydrate <b>50MG, 100MG CAPS</b><br>doxycycline monohydrate 40MG, 75MG and 150MG <b>CAP</b> (generic Adoxa/Monodox/ Oracea)<br>minocycline HCl <b>CAPS</b> (generic Dynacin/ Minocin/Myrac)<br>minocycline HCl ER (generic Solodyn)<br>NUZYRA (omadacycline) | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a sequential 3-day trial of TWO preferred agents within this drug class</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>demeclocycline:</b> Approved for diagnosis of SIADH</li> <li>▪ <b>doxycycline suspension:</b> May be approved with documented swallowing difficulty</li> </ul> |

**THROMBOPOIESIS STIMULATING PROTEINS<sup>CL</sup>**

| Preferred Agents                  | Non-Preferred Agents  | Prior Authorization/Class Criteria   |
|-----------------------------------|---|--|
| PROMACTA (eltrombopag) <b>TAB</b> | ALVAIZ (eltrombopag choline) <sup>AL</sup><br>DOPTELET (avatrombopag) <sup>AL</sup> <b>TAB, SPRINKLES<sup>AL,NR</sup></b><br>eltrombopag (generic Promacta) <b>SUSP, TAB</b><br>MULPLETA (lusutrombopag) <sup>CL</sup><br>PROMACTA (eltrombopag) <b>SUSP</b><br>TAVALISSE (fostamatinib)<br>WAYRILZ (rilzabrutinib) <sup>NR</sup> | <ul style="list-style-type: none"> <li>▪ All agents will be approved with FDA-approved indication, ICD-10 code is required.</li> <li>▪ Non-preferred agents require a trial of a preferred agent with the same indication or a contraindication.</li> </ul> <p>Drug-Specific Criteria</p> <ul style="list-style-type: none"> <li>▪ <b>Mulpleta:</b> Approved for one course of therapy for a scheduled procedure with a risk of bleeding for treatment of thrombocytopenia in adult patients with chronic liver disease</li> </ul> |

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**THYROID HORMONES**

| Preferred Agents   | Non-Preferred Agents   | Prior Authorization/Class Criteria   |
|--|--|--|
| levothyroxine (generic Synthroid) <b>TAB</b><br>liothyronine (generic Cytomel) <b>TAB</b><br>thyroid, pork <b>TAB</b><br>UNITHROID (levothyroxine) | ADTHYZA (thyroid, pork)<br>ERMEZA (levothyroxine) <b>SOLN</b><br>EUTHYROX (levothyroxine)<br>LEVO-T (levothyroxine)<br>levothyroxine <b>CAPS</b> (generic Tirosint)<br>RENTHYROID (thyroid, pork) <sup>NR</sup> <b>TAB</b><br>SYNTHROID (levothyroxine) <b>TAB</b><br>THYQUIDITY (levothyroxine) <b>SOLN</b> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class</li> </ul> |

**ULCERATIVE COLITIS**

| Preferred Agents  | Non-Preferred Agents  | Prior Authorization/Class Criteria  |
|---|---|---|
| <b>ORAL</b>   |   |   |
| PENTASA (mesalamine)<br>mesalamine (generic Lialda)<br>sulfasalazine IR, DR (generic Azulfidine)      | balsalazide (generic Colazal)<br>budesonide DR (generic Uceris)<br>DIPENTUM (olsalazine)<br>LIALDA (mesalamine)<br>mesalamine ER (generic Apriso)<br>mesalamine ER (generic Pentasa)<br>mesalamine (generic Asacol HD/Delzicol) <sup>CL</sup> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class</li> </ul> <p>Drug-specific criteria:</p> <ul style="list-style-type: none"> <li>▪ <b>Asacol HD®/Delzicol DR®:</b><br/>Requires clinical reason why preferred mesalamine products cannot be used</li> </ul> |
| <b>RECTAL</b>   |   |   |
| mesalamine (generic Canasa)<br><b>SUPPOSITORY</b><br>Sulfite-Free ROWASA (mesalamine)<br><b>ENEMA</b> | CANASA (mesalamine)<br>mesalamine (generic Rowasa) <b>ENEMA</b><br>UCERIS (budesonide)  |   |

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**UTERINE DISORDER TREATMENT**

| Preferred Agents   | Non-Preferred Agents | Prior Authorization/Class Criteria  |
|--|----------------------|---|
| MYFEMBREE (relugolix/ estradiol/<br>norethindrone acetate) <sup>AL, CL, QL</sup><br>ORIAHNN (elagolix/ estradiol/<br>norethindrone) <sup>AL, CL</sup><br>ORLISSA (elagolix sodium) <sup>QL, CL</sup> |                      | Drug-specific criteria: <ul style="list-style-type: none"> <li>• <b>Myfembree, Orilissa, and Oriahnn:</b> Requires an FDA approved indication, must follow FDA dosing guidelines, and have had a trial and failure of an NSAID and oral contraceptive                             <ul style="list-style-type: none"> <li>○ Total duration of treatment is max of 24 months</li> </ul> </li> </ul> |

**VASODILATORS, CORONARY**

| Preferred Agents   | Non-Preferred Agents   | Prior Authorization/Class Criteria  |
|--|--|---|
| isosorbide dinitrate <b>TAB</b><br>isosorbide dinitrate/hydralazine (Bidil) <sup>CL</sup><br>isosorbide mono IR/SR <b>TAB</b><br>nitroglycerin <b>SUBLINGUAL, TRANSDERMAL</b><br>nitroglycerin ER <b>TAB</b> | BIDIL (isosorbide dinitrate/<br>hydralazine) <sup>CL</sup><br>GONITRO (nitroglycerin)<br>isosorbide dinitrate <b>TAB (Oceanside Pharm MFR only)</b><br>NITRO-BID <b>OINT</b> (nitroglycerin)<br>NITRO-DUR (nitroglycerin)<br>nitroglycerin <b>TRANSLINGUAL</b><br>(generic Nitrolingual)<br>VERQUVO (vericiguat) <sup>AL, CL, QL</sup> | <ul style="list-style-type: none"> <li>▪ Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class</li> </ul> Drug-specific criteria: <ul style="list-style-type: none"> <li>▪ <b>BiDil/ isosorbide dinitrate-hydralazine:</b> Approved for the treatment of heart failure as an adjunct therapy to standard therapy in self-identified black patients</li> <li>▪ <b>Verquvo:</b> Approved for use in patients following a recent hospitalization for HF within the past 6 months OR need for outpatient IV diuretics, in adults with symptomatic chronic HF and EF less than 45%</li> </ul> |

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