



DEPT. OF HEALTH AND HUMAN SERVICES

Nebraska Medicaid Preferred Drug List with Prior Authorization Criteria

July 2024 PDL with May 2024 P&T changes implemented Noted in Red Font that Become Effective July 19, 2024

For the most up to date list of covered drugs consult the **Drug Lookup** on the Nebraska Medicaid website at https://ne.magellanrx.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

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

	Non Duefermed Amende	Duian Anthoniadian (Class City
Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
dapalene (generic Differin) GEL (OTC/Rx), GEL PUMP dapalene/BPO (generic Epiduo) benzoyl peroxide (BPO) WASH, LOTION benzoyl peroxide GEL OTC dindamycin/BPO (generic BenzaClin) GEL, PUMP dindamycin/BPO (generic Duac) dindamycin phosphate PLEDGET dindamycin phosphate SOLUTION erythromycin GEL erythromycin-BPO (generic for Benzamycin) RETIN-A (tretinoin) ^{AL} CREAM, GEL	adapalene (generic Differin) CREAM adapalene/BPO (generic Epiduo Forte) ALTRENO (tretinoin) ^{AL} AMZEEQ (minocycline) ARAZLO (tazarotene) ^{AL} ATRALIN (tretinoin) AVAR (sulfacetamide sodium/sulfur) AVITA (tretinoin) AZELEX (azelaic acid) BENZEFOAM (benzoyl peroxide) benzoyl peroxide CLEANSER, CLEANSING BAR OTC benzoyl peroxide FOAM (generic BenzePro) benzoyl peroxide GEL Rx benzoyl peroxide TOWELETTE OTC CABTREO (clindamycin phosphate/BPO/adapalene) ^{AL} GEL clindamycin GEL clindamycin GEL clindamycin/BPO (generic Acanya) GEL clindamycin/BPO PUMP(generic Onexton) ^{AL} clindamycin/tretinoin (generic Veltin, Ziana) dapsone (generic Aczone) erythromycin PLEDGET EVOCLIN (clindamycin)	Non-preferred agents will be approved for patients who have failed THREE preferred agents within this drug class

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

ACNE AGENTS, TOPICAL (Continued)

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
	FABIOR (tazarotene) FOAM NEUAC (clindamycin/BPO) ONEXTON (clindamycin/BPO) OVACE PLUS (sulfacetamide sodium) RETIN-A MICRO (tretinoin) sulfacetamide sulfacetamide/sulfur sulfacetamide/sulfur cLEANSER SUMADAN (sulfacetamide/sulfur) tazarotene (generic Tazorac) CREAM tazarotene FOAM (generic Fabior) tazarotene GEL (generic Tazorac) TRETIN-X (tretinoin) tretinoin (generic Avita, Retin-A) CREAM, GEL tretinoin microspheres (generic Retin-A Micro) AL GEL, GEL PUMP WINLEVI (clascoterone)AL	Non-preferred agents will be approved for patients who have failed THREE preferred agents within this drug class

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

ALZHEIMER'S AGENTS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
CHOLINESTERA	ASE INHIBITORS •	Non-preferred agents will be approved for patients who have
donepezil (generic Aricept) donepezil ODT (generic Aricept ODT) rivastigmine PATCH (generic for Exelon Patch)	ADLARITY (donepezil) PATCH ARICEPT (donepezil) donepezil 23 (generic Aricept 23) ^{CL} EXELON (rivastigmine) PATCH galantamine (generic Razadyne) SOLN, TAB galantamine ER (generic Razadyne ER) rivastigmine CAPS (generic Exelon)	failed a 120-day trial of ONE preferred agent within this drug class within the last 6 months OR Current, stabilized therapy of the non-preferred agent within the previous 45 days
	Di	rug-specific criteria:
NMDA RECEPTO	DR ANTAGONIST	Donepezil 23: Requires donepezil 10mg/day for at least 3 months
	memantine ER (generic Namenda XR) memantine SOLN (generic Namenda) NAMENDA (memantine) NAMZARIC (memantine/donepezil)	AND clinical reason as to why 5mg or 10mg tablets can't be used (to deliver 20mg or 25mg)

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

ANALGESICS, OPIOID LONG-ACTING

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
	buprenorphine BUCCAL (generic for Belbuca) AL,QL buprenorphine PATCH (generic Butrans)QL EMBEDA (morphine sulfate/ naltrexone DURAGESIC MATRIX (fentanyl)QL fentanyl 37.5/62.5/87.5 mcg PATCH QL hydrocodone ER (generic Hysingla ER)QL hydrocodone bitartrate ER (generic Zohydro ER) hydromorphone ER (generic Exalgo)CL	 does not recommend long acting opioids when beginning opioid treatment. Preferred agents require previous use of a long acting opioid or documentation of a trial on a short acting agent within 90 days Non-preferred agents will be approved with failure on, or intolerance to TWO preferred agents within this drug class Drug-specific criteria: Methadone: Will only be approved for use in pain control or end of life care. Trial of preferred agent not required for end of life

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

ANALGESICS, OPIOID SHORT-ACTING QL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
OF	RAL	Non-preferred agents will be
cetaminophen/codeine ELIXIR, TAB odeine TAB ydrocodone/APAP SOLN, TAB ydrocodone/ibuprofen ydromorphone TAB norphine CONC SOLN, SOLN, TAB xycodone TAB, SOLN xycodone/APAP ramadol 50 TAB ^{AL} (generic Ultram)	APADAZ (benzhydrocodone/APAP) ^{CL} benzhydrocodone/APAP (generic Apadaz. ^{CL} butalbital/caffeine/APAP/codeine butalbital compound w/codeine (butalbital/ASA/caffeine/codeine) carisoprodol compound-codeine (carisoprodol/ASA/codeine) dihydrocodeine/APAP/caffeine dihydrocodeine/aspirin/caffeine hydromorphone LIQUID, SUPPOSITORY (generic Dilaudid) levorphanol meperidine (generic Demerol) morphine SUPPOSITORIES NALOCET (oxycodone/APAP) NUCYNTA (tapentadol) ^{CL} oxycodone CAPS oxycodone/APAP SOLN oxycodone CONCENTRATE oxymorphone IR (generic Opana) pentazocine/naloxone PROLATE (oxycodone/APAP) SOLN,	approved for patients who have failed THREE preferred agents within this drug class within the la 12 months Note: for short acting opiate table and capsules there is a maximum quantity limit of #150 per 30 days Opiate limits for opiate naïve patients will consist of: -prescriptions limited to a 7 day supply, AND -initial opiate prescription fill limite to maximum of 50 Morphine Milligram Equivalents (MME) per day These limits may only be exceed 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 naive. Drug-specific criteria: Apadaz//benzhydrocodone-APAP: Approval for 14 days or less Nucynta®: Approved only for diagnosis of acute pain, for 30 days or less

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

ANALGESICS, OPIOID SHORT-ACTING QL (Continued)

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
NA	SAL	
	butorphanol SPRAY ^{QL} LAZANDA (fentanyl citrate)	
BUCCAL/TRANSMUCOSAL ^{CL}		TDrug-specific criteria: _• Abstral®/Actiq®/Fentora®/
	ABSTRAL (fentanyl) ^{CL} fentanyl TRANSMUCOSAL (generic Actiq) ^{CL} FENTORA (fentanyl) ^{CL}	fentanyl transmucosal/Onsolis: Approved only for diagnosis of cancer AND current use of long- acting opiate

ANDROGENIC AGENTS (TOPICAL) CL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
ANDROGEL (testosterone) PUMP CL estosterone PUMP (generic Androgel) CL ESTIM (testosterone) TRANSDERMAL	ANDRODERM (testosterone) ^{CL} NATESTO (testosterone) ^{CL} testosterone PACKET (generic Androgel) ^{CL} testosterone GEL, PACKET, PUMP (generic Vogelxo) testosterone (generic Axiron) testosterone (generic Fortesta) testosterone (generic Testim)	 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 In addition, non-preferred agents will be approved for patients who have failed ONE preferred agent within this drug class within the la 6 months Drug-specific criteria: Androderm®/Androgel®: Approved for Males only Natesto®: Approved for Males on with diagnosis of: Primary hypogonadism (congenit or acquired) OR Hypogonadotropic hypogonadism (congenital or acquired)

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

ANGIOTENSIN MODULATORS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
benazepril (generic Lotensin) enalapril (generic Vasotec) lisinopril (generic Prinivil, Zestril) quinapril (generic Accupril) ramipril (generic Altace)	captopril (generic Capoten) EPANED (enalapril) ^{CL} ORAL SOLN enalapril (generic for Epaned) ^{CL} ORAL SOLN fosinopril (generic Monopril) moexepril (generic Univasc) perindopril (generic Aceon) QBRELIS (lisinopril) ^{CL} ORAL SOLN	 Non-preferred agents will be approved for patients who have failed TWO preferred agents within this drug class within the last 12 months Non-preferred combination products may be covered as individual prescriptions without prior authorization Drug-specific criteria: Epaned/enalapril oral solution: Clinical reason why oral tablet is not appropriate
irbesartan (generic Avapro) losartan (generic Cozaar) olmesartan (generic Benicar) valsartan (generic Diovan)	candesartan (generic Atacand) EDARBI (azilsartan) eprosartan (generic Teveten) telmisartan (generic Micardis)	

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

ANGIOTENSIN MODULATORS (Continued)

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
ANGIOTENSIN RECEPTOR BLO	ANGIOTENSIN RECEPTOR BLOCKER/DIURETIC COMBINATIONS	
irbesartan/HCTZ (generic Avalide) losartan/HCTZ (generic Hyzaar) olmesartan/HCTZ (generic Benicar- HCT) valsartan/HCTZ (generic Diovan-HCT)	candesartan/HCTZ (generic Atacand- HCT) EDARBYCLOR (azilsartan/ chlorthalidone) telmisartan/HCTZ (generic Micardis- HCT)	 approved for patients who have failed TWO preferred agents within this drug class within the last 12 months Non-preferred combination products may be covered as individual prescriptions without prior authorization
	I MODULATOR/ OCKER COMBINATIONS	
amlodipine/benazepril (generic Lotrel) amlodipine/olmesartan (generic Azor) amlodipine/valsartan (generic Exforge) amlodipine/valsartan/HCTZ (generic Exforge HCT)	amlodipine/olmesartan/HCTZ (generic Tribenzor) amlodipine/telmisartan (generic Twynsta) PRESTALIA (perindopril/amlodipine) trandolapril/verapamil (generic Tarka)	
DIRECT RENI	N INHIBITORS	
	aliskiren (generic Tekturna) ^{QL}	
DIRECT RENIN INHIB	ITOR COMBINATIONS	Direct Renin Inhibitors/Direct
	TEKTURNA/HCT (aliskiren/HCTZ)	Renin Inhibitor Combinations: May be approved with history of
NEPRILYSIN INHIBI	TOR COMBINATION	TWO preferred ACE Inhibitors or Angiotensin Receptor Blockers
ENTRESTO (sacubitril/valsartan) ^{CL,QL}		within the last 12 months
ANGIOTENSIN RECEPTOR BLOCKE	ER/BETA-BLOCKER COMBINATIONS	Drug Specific Criteria
	BYVALSON (nevibolol/valsartan)	Entresto: May be approved in patients ages >1 years old and with a diagnosis of heart failure

ANTHELMINTICS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
albendazole (generic Albenza) BILTRICIDE (praziquantel) ivermectin (generic Stromectol)	EMVERM (mebendazole) ^{CL} praziquantel (generic Biltricide) STROMECTOL (ivermectin)	 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 Drug-specific criteria: Emverm: Approval will be considered for indications not covered by preferred agents

Unless otherwise specified, the listing of a particular brand or generic name includes all dosage forms of that drug.

CL - Prior Authorization / Class Criteria apply

QL - Quantity/Duration Limit

AL- Age Limit NR - Product was not reviewed - New Drug criteria will apply

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

ANTI-ALLERGENS, ORAL

Preferred Agents GRASTEK (timothy grass pollen allergen p. N-Co. ODACTRA (Dermatophagoides farinae and Dermatophagoides pteronyssinus)-k-Co. ORALAIR (sweet vernal/orchard/rye/ timothy/kentucky blue grass mixed pollen allergen extract) ^{Co.} PALFORZIA (peanut allergen powder drip) A-Co. RAGWITEK (weed pollen-short ragweed) ^{AL,OL} RAGWITEK (weed pollen-short ragweed) ^{AL,OL} ROBALAIR Brown of the provided particular and provided by positive skin test or in vitro testing for pollen specific lgE antibodies to Timothy grass or cross-reaction of the provided by positive skin test or in vitro testing for pollen specific lgE antibodies to Dermatophagoides farinae and Dermatophagoides farinae and Dermatophagoides premyssinus house dust mite allergen extracts or in vitro testing for pollen specific lgE antibodies to Dermatophagoides farinae and Dermatophagoides premyssinus house dust mite allergen extracts or in vitro testing for pollen specific lgE antibodies for Sweet Pollens. • For use in persons 12 through 65 years of age. PALFORZIA • Confirmed by positive skin test or in vitro testing for pollen specific lgE antibodies for Sweet Pollens. • For use in patients 5 through 65 years of age. PALFORZIA • Confirmed diagnosis of peanut allergy by allergist • For use in patients 5 through 65 years of age. PALFORZIA • Confirmed diagnosis of peanut allergy by allergist • 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 • Initial dose and increase titration doses should be given in a healthcare setting. • Should not be used in patients with uncontrolled asthma or concurrently on a NSAIU RAGWITEK • Confirmed by positive skin test or in vitro testing for pollen specific lgE antibodies for short tragweed pollen. • For use in patients 5 through 65 years of age.	ANTI-ALLERGENS, ORAL		
allergen) AL-OL ODACTRA (Dermatophagoides farinae and Dermatophagoides pieronyssinus) AL-OL ORAL-JAIR (sweet vernal/orchard/rye/timothy/kentucky blue grass mixed pollen allergen extract) Cl. PALFORZIA (peanut allergen powderdrip) AL-OL RAGWITEK (weed pollen-short ragweed) AL-OL RAGWITEK (weed pollen-short ragweed pollen. Por use in persons 12 through 65 years of age ORALAIR 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. Por use in patients 5 through 65 years of age. PALFORIA Confirmed diagnosis of peanut allergy by allergist Por use in patients 3 through 65 years of age. PALFORIA Solomitation of previous use within the past 90 days Initial dose and increase titration doses should be given in a healthcare setting	Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
		allergen) AL,QL ODACTRA (Dermatophagoides farinae and Dermatophagoides pteronyssinus) AL,QL ORALAIR (sweet vernal/orchard/rye/timothy/kentucky blue grass mixed pollen allergen extract) CL PALFORZIA (peanut allergen powderdnfp) AL,CL RAGWITEK (weed pollen-short	Drug-specific criteria: GRASTEK Confirmed by positive skin test or in vitro testing for pollen specific IgE antibodies for Timothy grass or cross-reactive grass pollens. For use in persons 5 through 65 years of age. ODACTRA Confirmed by positive skin test to licensed house dust mite allergen extracts or in vitro testing for IgE antibodies to Dermatophagoides farinae and Dermatophagoides farinae and Dermatophagoides pteronyssinus house dust mite For use in persons 12 through 65 years of age ORALAIR 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. For use in patients 5 through 65 years of age. PALFORZIA Confirmed diagnosis of peanut allergy by allergist 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 Initial dose and increase titration doses should be given in a healthcare setting Should not be used in patients with uncontrolled asthma or concurrently on a NSAID RAGWITEK Confirmed by positive skin test or in vitro testing for pollen specific IgE antibodies for short ragweed pollen. For use in patients 5 through 65

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

ANTIBIOTICS, GASTROINTESTINAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
metronidazole TABLET neomycin tinidazole (generic Tindamax) ^{CL} metronidazole (generic Tindamax) ^{CL} metronidazole (generic Tindamax) ^{CL} metronidazole (generic Tindamax) ^{CL} vand Vov	CID (fidaxomicin) CL TAB, SUSP MEZ (metronidazole) SUSP ronidazole CL CAPS zoxanide generic Alinia) TABAL, CL, QL pmomycin LOSEC (secnidazole) comycin CAPS (generic ancocin) CL comycin (generic Firvanq) QL VST (fecal microbiota spores) AL, QL AXAN (rifaximin) CL	 Note: Although azithromycin, ciprofloxacin, and trimethoprim/ sulfmethoxazole are not included in this review, they are available without prior authorization Drug-specific criteria: Alinia /nitazoxanide tablet: Trial and failure with metronidazole is required for a diagnosis of giardiasis Dificid®: 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. Flagyl®/Metronidazole 375mg capsules and / Metronidazole 750mg ER tabs: Clinical reason why the generic regular release cannot be used tinidazole: Approvable diagnoses include: Giardia Amebiasis intestinal or liver abscess Bacterial vaginosis or trichomoniasis vancomycin capsules: Requires patient specific documentation of why the Firvanq/vancomycin solution is not appropriate for patient Xifaxan®: Approvable diagnoses include: Travelers's diarrhea resistant to quinolones Hepatic encephalopathy with treatment failure of lactulose or neomycin Diarrhea-Predominant IBS (IBS-D) 550mg strength only with treatment failure of Lomotil® AND Imodium®

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

ANTIBIOTICS, INHALED CL

Preferred Agents ^{CL}	Non-Preferred Agents	Prior Authorization/Class Criteria
BETHKIS (tobramycin) KITABIS PAK (tobramycin) tobramycin (generic Tobi) TOBI-PODHALER (tobramycin) QL	ARIKAYCE (amikacin liposomal inh) SUSP CAYSTON (aztreonam lysine) ^{QL} tobramycin (generic Bethkis)	 Diagnosis of Cystic Fibrosis is required for all agents ICD10 Group = E84, ICD9 = 277.00, 277.01, 277.02, 277.03, 277.09 Drug-specific criteria: Arikayce: 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 Cayston®: Trial of tobramycin via nebulizer and demonstration of TOBI® compliance required Tobi Podhaler®: Requires trial of tobramycin via nebulizer or documentation why nebulized tobramycin cannot be used

ANTIBIOTICS, TOPICAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
bacitracin OINT bacitracin OINT OTC bacitracin/polymyxin (generic Polysporin) mupirocin OINT (generic Bactroban) neomycin/polymyxin/bacitracin (generic Neosporin, Triple AB) neomycin/polymyxin/pramoxine neomycin/polymyxin/bacitracin/ pramoxine	CENTANY (mupirocin) gentamicin OINT, CREAM mupirocin CREAM (generic Bactroban) ^{CL}	 Non-preferred agents will be approved for patients who have failed ALL preferred agents within this drug class within the last 12 months Drug-specific criteria: Mupirocin® Cream: Clinical reason the ointment cannot be used

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

ANTIBIOTICS, VAGINAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
CLEOCIN OVULES (clindamycin) clindamycin CREAM (generic Cleocin) metronidazole, vaginal NUVESSA (metronidazole)	CLEOCIN CREAM (clindamycin) CLINDESSE (clindamycin) metronidazole (generic Nuvessa) ^{NR} VANDAZOLE (metronidazole) XACIATO (clindamycin phosphate) GEL AL	 Non-preferred agents will be approved for patients who have failed a therapeutic trial (duration = 3 days) with ONE 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)	fondaparinux (generic Arixtra) FRAGMIN (dalteparin) PRADAXA (dabigatran) CAPS, PELLETS SAVAYSA (edoxaban)CL,QL XARELTO (rivaroxaban)CLSUSP	 Non-preferred agents will be approved for patients who have failed ONE preferred agent within this drug class within the last 12 months Coumadin®: Clinical reason generic warfarin cannot be used Savaysa®: Approved diagnoses include: Stroke and systemic embolism (SE) risk reduction in nonvalvular atrial fibrillation (NVAF) OR

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

ANTIEMETICS/ANTIVERTIGO AGENTS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
CANNABINOIDS		Non-preferred agents will be
dronabinol (generic Marinol) ^{AL}	CESAMET (nabilone)	approved for patients who have failed ONE preferred agent within this drug class within the same
5HT3 RECEPT	OR BLOCKERS	group
ondansetron (generic Zofran/Zofran ODT) ^{QL}	ANZEMET (dolasetron) granisetron (generic Kytril) SANCUSO (granisetron) ^{CL} ZUPLENZ (ondansetron)	 Drug-specific criteria: Akynzeo®/Varubi®: Approved for Moderately/Highly emetogenic chemotherapy with dexamethasone and a 5-HT3 antagonist
NK-1 RECEPTO	R ANTAGONIST	Regimens include: AC combination (Doxorubicin or Epirubicin with
aprepitant (generic Emend) CAPS QL	AKYNZEO (netupitant/palonosetron) ^{CL} aprepitant (generic Emend) PACK EMEND (aprepitant) CAPS, PACK, POWDER ^{QL} VARUBI (rolapitant) TAB ^{CL}	Cyclophosphamide), Aldesleukin, Amifostine, Arsenic trioxide, Azacitidine, Bendamustine, Busulfan, Carmustine, Carbplatin, Cisplatin, Clofarabine, Cyclophosphamide, Cytarabine, Dacarbazine, Dactinomycin, Daunorubicin,
TRADITIONAL	ANTIEMETICS	Epirubicin, Etoposide,
DICLEGIS (doxylamine/pyridoxine) ^{CL,QL} dimenhydrinate (generic Dramamine) OTC meclizine (generic Antivert) metoclopramide (generic Reglan) phosphoric acid/dextrose/fructose (generic Emetrol) SOLN prochlorperazine(generic Compazine) promethazine (generic Phenergan) SYRUP , TAB promethazine 12.5mg, 25mg SUPPOSITORY scopolamine TRANSDERMAL	BONJESTA (doxylamine/pyridoxine),CL,QL COMPRO (prochlorperazine) doxylamine/pyridoxine (generic Diclegis)CL,QL metoclopramide ODT (generic Metozolv ODT) prochlorperazine SUPPOSITORY (generic Compazine) promethazine SUPPOSITORY 50mg TRANSDERM-SCOP (scopolamine) trimethobenzamide TAB (generic Tigan)	 Hexamethylmelamine, Idarubicin, Ifosfamide, Imatinib, Interferon α, Irinotecan, Mechlorethamine, Melphalan, Methotrexate, Oxaliplatin, Procarbazine, Streptozotocin, Temozolomide Diclegis/doxylamine-pyridoxine)/ Bonjesta: Approved only for treatment of nausea and vomiting of pregnancy Metozolv (metoclopramide) ODT®: Documentation of inability to swallow or Clinical reason oral liquid cannot be used Sancuso®/Zuplenz®: Documentation of oral dosage form intolerance

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

ANTIFUNGALS, ORAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
lotrimazole (mucous membrane, roche) luconazole SUSP, TAB (generic Diflucan) griseofulvin SUSP griseofulvin microsized TAB aystatin SUSP, TAB erbinafine (generic Lamisil)	BREXAFEMME (ibrexafungerp) ^{QL} CRESEMBA (isavuconazonium) ^{CL} flucytosine (generic Ancobon) ^{CL} griseofulvin ultramicrosize (generic GRIS-PEG) itraconazole (generic Sporanox) ^{CL} ketoconazole (generic Nizoral) NOXAFIL (posaconazole) AL SUSP, TAB NOXAFIL (posaconazole) AL,CL POWDERMIX nystatin POWDER posaconazole (generic Noxafil) AL,CL TOLSURA (itraconazole) CAPS voriconazole (generic VFEND) ^{CL}	 Non-preferred agents will be approved for patients who have failed a trial of TWO diagnosis-appropriate preferred agents within this drug class Drug-specific criteria: Cresemba®: Approved for diagnosis invasive aspergillosis or invasive mucormycosis Flucytosine: Approved for diagnosis of: Candida: Septicemia, endocarditis UTIs Cryptococus: Meningitis, pulmonary infections Noxafil/ posaconazole DR tablets, oral suspension, PowderMix® for delayed oral suspension:: No trial for diagnosis of Neutropenia Myelodysplastic Syndrome (MDS), Neutropenic Acute Myeloid Leukemia (AML), Neutropenic hematologic malignancies, Graft vs. Host disease (GVHD), Immunosuppression secondary to hematopoietic stem cell transplant Noxafil® Powdermix: pediatric patients 2 years of age and older who weigh 40 kg or less Noxafil/ posaconazole Suspension Oropharyngeal/esophageal candidias refractory to itraconazole and/or fluconazole Sporanox®/Itraconazole: Approved for diagnosis of Aspergillosis, Blastomycosis, Histoplasmosis, Onychomycosis due to terbinafine-resistant dermatophytes, Oropharyngeal/ esophageal candidiasis refractory to fluconazole Sporanox® Liquid: Clinical reason solid oral cannot be used Tolsura: Approved for diagnosis of Aspergillosis, Blastomycosis, and Histoplasmosis and requires a trial an failure of generic itraconazole Vfend/voriconazole:: No trial for diagnosis of Myelodysplastic Syndrome (MDS), Neutropenic Acute Myeloid Leukemia (AML), Graft vs. Host disease (GVHD), Candidemia (candida krusei), Esophageal Candidiasis, Blastomycosis, S. apiospermum and Fusarium spp., Oropharyngeal/esophageal candidias refractory to fluconazole

Unless otherwise specified, the listing of a particular brand or generic name includes all dosage forms of that drug.

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

ANTIFUNGALS, TOPICAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
ANTIF	UNGAL	Non-preferred agents will be
clotrimazole CREAM (generic Lotrimin) RX, OTC clotrimazole SOLN OTC ketoconazole CREAM, SHAMPOO (generic Nizoral) miconazole CREAM, POWDER OTC nystatin terbinafine OTC (generic Lamisil AT) tolnaftate AERO-POWDER OTC, CREAM-OTC, SOLN-OTC (generic Tinactin)	ALEVAZOL (clotrimazole) OTC ciclopirox CREAM, GEL, SUSP (generic Ciclodan, Loprox) ciclopirox NAIL LACQUERCL (generic Penlac) ciclopirox SHAMPOO (generic Loprox) clotrimazole SOLN RX (generic Lotrimin) DESENEX POWDER OTC (miconazole) econazole (generic Spectazole) ERTACZO (sertaconazole) EXELDERM (sulconazole) FUNGOID (miconazole) OTC JUBLIA (efinaconazole) CL ketoconazole FOAMCL (generic Extina, Ketodan) LOPROX (ciclopirox) SUSP, SHAMPOO, CREAM LOTRIMIN AF CREAM OTC (clotrimazole) LOTRIMIN ULTRA (butenafine) luliconazole (generic Luzu) MENTAX (butenafine) miconazole OTC OINT, SPRAY, SOLN miconazole/zinc oxide/petrolatum (generic Vusion) naftifine CREAM, GEL (generic Naftin) oxiconazole (generic Densal HP) tavaborole SOLNCL (generic Kerydin) tolnaftate POWDER OTC TRIPENICOL (undecylenic acid) NR CREAM OTC VOTRIZA-AL (clotrimazole) LOTION OTC	approved for patients who have failed a trial of TWO preferred agents within this drug class within the last 6 months Drug-specific criteria: Extina/ Ketodan/ ketoconazole foam: Requires trial and failure or contraindication to other ketoconazole forms Jublia and tavaborole: Approved diagnoses include Onychomycosis of the toenails due to T.rubrum OR T. Mentagrophytes ciclopirox nail lacquer: No trial required in diabetes, peripheral vascular disease (PVD), immunocompromised OR contraindication to oral terbinafine
ANTIFUNGAL/STER	OID COMBINATIONS	
	clotrimazole/betamethasone LOTION	
(generic Lotrisone) nystatin/triamcinolone (generic Mycolog) CREAM, OINT	(generic Lotrisone)	

Unless otherwise specified, the listing of a particular brand or generic name includes all dosage forms of that drug.

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

ANTIHISTAMINES, MINIMALLY SEDATING

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
cetirizine TAB (generic Zyrtec) cetirizine SOLN (OTC) (generic Zyrtec) loratadine TAB, SOLN (generic Claritin) levocetirizine TAB (generic Xyzal)	cetirizine CHEWABLE (generic Zyrtec) cetirizine SOLN (Rx) (generic Zyrtec) desloratadine (generic Clarinex) desloratadine ODT (generic Clarinex Reditabs) fexofenadine (generic Allegra) fexofenadine 180mg (generic Allegra 180mg) ^{QL} levocetirizine (generic Xyzal) SOLN loratadine CAPS, CHEWABLE, ODT (generic Claritin Reditabs)	 Non-preferred agents will be approved for patients who have failed TWO preferred agents within this drug class Combination products not covered – individual products may be covered

ANTIHYPERTENSIVES, SYMPATHOLYTICS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
clonidine TAB (generic Catapres) clonidine TRANSDERMAL guanfacine (generic Tenex) methyldopa	methyldopa/hydrochlorothiazide	Non-preferred agents will be approved for patients who have failed a 30-day trial with ONE preferred agent within this drug class

ANTIHYPERURICEMICS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
allopurinol (generic Zyloprim) colchicine TAB (generic Colcrys) probenecid probenecid/colchicine (generic Col- Probenecid)	allopurinol 200mg colchicine CAPS (generic Mitigare) febuxostat (generic Uloric) ^{CL} GLOPERBA SOLN (colchicine) ^{CL,QL} MITIGARE (colchicine)	 Non-preferred agents will be approved for patients who have failed a trial with ONE preferred agent within this drug class Gloperba: Approved for documented swallowing disorder Uloric/febuxostat: Clinical reason why allopurinol cannot be used

Unless otherwise specified, the listing of a particular brand or generic name includes all dosage forms of that drug.

CL - Prior Authorization / Class Criteria apply

QL - Quantity/Duration Limit

AL- Age Limit NR - Product was not reviewed - New Drug criteria will apply

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

ANTIMIGRAINE AGENTS, OTHER

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
IMOVIG (erenumab-aooe) CL,QL JOVY (fremanezumab-vfrm) CL, QL PEN, Autoinjector JOVY (fremanezumab-vfrm) Autoinjector 3-pack ^{CL,QL} MGALITY 120 mg/mL (galcanezumab-gnlm) CL, QL PEN, SYRINGE IURTEC ODT (rimegepant) AL,CL,QL BRELVY (ubrogepant) AL,CL, QL TAB	diclofenac (generic Cambia) POWDER dihydroergotamine mesylate NASAL ELYXYB (celecoxib)AL,QL SOLN EMGALITY 100 mg (galcanezumabgnlm) CL,QL SYR MIGERGOT (ergotamine/caffeine) RECTAL MIGRANAL (dihydroergotamine) NASAL REYVOW (lasmiditan)AL, CL,QL TAB TRUDHESA (dihydroergotamine mesylate)AL,QL NASAL ZAVZPRET (zavegepant)AL,QL NASAL	 In addition, all non-preferred agents will require a failed trial or contraindication of a preferred agent of the same indication For Acute Treatment: agents will be approved for patients who have a failed trial or a contraindication that triptan. For Prophylactic Treatment: Required 4 migraines per month for ≥ 3 months and has tried and failed a ≥ 1 month trial of two medications listed in the 2012 American Academy of Neurology/American Headache Society guidelines (examples include: antidepressants (amitriptyline, venlafaxine), Beta blockers (propranolol, metroprolol, atenolol), anti-epileptics (valproate, topiramate), ACE (lisinopril) Drug-specific criteria: Emgaility 100mg will only be approved for treatment of Episodic Cluster Headache Nurtec ODT: for use in acute treatment, will be approved for patients who have a failed trial or contraindication to a triptan. For use in preventative treatment, will be approved for patients who have a failed trial of ONE preferred injectable CGRP. Qulipta: May be approved for patients who have a failed trial of ONE preferred injectable CGRP

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

ANTIMIGRAINE AGENTS, TRIPTANS QL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
rizatriptan (generic Maxalt) rizatriptan ODT (generic Maxalt MLT) sumatriptan	almotriptan (generic Axert) eletriptan (generic Relpax) frovatriptan (generic Frova) IMITREX (sumatriptan) naratriptan (generic Amerge) RELPAX (eletriptan) ^{QL} sumatriptan/naproxen (generic Treximet) zolmitriptan (generic Zomig)	 Non-preferred agents will be approved for patients who have failed ALL preferred agents within this drug class Drug-specific criteria: Sumavel® Dosepro: Requires clinical reason sumatriptan injection cannot be used Onzetra, Zembrace: approved for patients who have failed ALL preferred agents
IMITREX (sumatriptan) sumatriptan (generic Imitrex Nasal)	SAL ONZETRA XSAIL (sumatriptan) TOSYMRA (sumatriptan) zolmitriptan (generic Zomig) ZOMIG (zolmitriptan)	
sumatriptan KIT, SYRINGE, VIAL	IMITREX (sumatriptan) INJECTION SUMAVEL DOSEPRO (sumatriptan) ZEMBRACE SYMTOUCH (sumatriptan)	

ANTIPARASITICS, TOPICAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
NATROBA (spinosad) permethrin 1% OTC (generic Nix) permethrin 5% RX (generic Elimite) pyrethrin/piperonyl butoxide (generic RID, A-200)	CROTAN (crotamiton) LOTION EURAX (crotamiton) CREAM, LOTION ivermectin (generic Sklice) LOTION lindane malathion (generic Ovide) spinosad (generic Natroba) VANALICE (piperonyl butoxide/pyrethrins)	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

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

ANTIPARKINSON'S AGENTS, ORAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
ANTICHOL	INERGICS	Non-preferred agents will be
benztropine (generic Cogentin) trihexyphenidyl (generic Artane)		approved for patients who have failed ONE preferred agent within
7. 7. 10	HIBITORS	this drug class
pramipexole (generic Mirapex) ropinirole (generic Requip)	entacapone (generic Comtan) ONGENTYS (opicapone) tolcapone (generic Tasmar) AGONISTS bromocriptine (generic Parlodel) ropinirole ER (generic Requip ER) ^{CL} NEUPRO (rotigotine) ^{CL} pramipexole ER (generic Mirapex ER) ^{CL} ropinirole ER (generic Requip XL) ^{CL}	Carbidopa/Levodopa ODT: Approved for documented swallowing disorder COMT Inhibitors: Approved if using as add-on therapy with levodopacontaining drug Gocovri: 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 Inbrija: Approval upon diagnosis of Parkinson's disease and concurrent treatment with carbidopa/levodopa agent Neupro®:
MAO-B IN selegiline CAPS, TABLET (generic Eldepryl)	rasagiline (generic Azilect) QL XADAGO (safinamide) ZELAPAR (selegiline)CL	For Parkinsons: Clinical reason required why preferred agent cannot be used For Restless Leg (RLS): Requires trial OR Contraindication to ropinirole AND pramipexole
OTUED ANTIDAD	(VI) 20112 DD1120	• Nourianz: Approval upon diagnosis of
amantadine CAPS, SYRUP TABLET (generic Symmetrel) carbidopa/levodopa (generic Sinemet) carbidopa/levodopa ER (generic Sinemet CR) levodopa/carbidopa/entacapone (generic Stalevo)	APOKYN (apomorphine) SUB-Q apomorphine (generic Apokyn)SUB-Q carbidopa (generic Lodosyn) carbidopa/levodopa ODT (generic Parcopa) DHIVY (carbidopa/levodopa) GOCOVRI (amantadine) INBRIJA (levodopa) CL,QL INHALER KYNMOBI (apomorphine)QL KIT, SUBLINGUAL NOURIANZ (istradefylline)CL,QL OSMOLEX ER (amantadine)QL RYTARY (carbidopa/levodopa) STALEVO (ledopa/carbidopa/entacapone)	

Unless otherwise specified, the listing of a particular brand or generic name includes all dosage forms of that drug.

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

ANTIPSORIATICS, ORAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
acitretin (generic Soriatane)	methoxsalen (generic Oxsoralen- Ultra)	 Non-preferred agents will be approved for patients who have failed a trial with THE preferred agent within this drug class Trial of acitretin (Pregnancy category X) not required in pregnancy or while attempting or planning pregnancy

ANTIPSORIATICS, TOPICAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
calcipotriene CREAM, OINT, SOLN	calcitriol (generic Vectical) ^{AL} OINT calcipotriene/betamethasone OINT (generic Taclonex) calcipotriene/betamethasone SUSP (generic Taclonex Scalp) CALCITRENE (calcipotriene) DOVONEX CREAM (calcipotriene) DUOBRII (halobetasol prop/tazarotene ENSTILAR (calcipotriene/betamethasone) SORILUX (calcipotriene) VTAMA (tapinarof) ^{AL} CREAM ZORYVE (roflumilast) ^{AL} CREAM	Non-preferred agents will be approved for patients who have failed a trial with ONE preferred agent within this drug class

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

ANTIVIRALS, ORAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
acyclovir (generic Zovirax) famciclovir (generic Famvir) valacyclovir (generic Valtrex)	acyclovir (generic for Zovirax) ^{CL} SUSP SITAVIG (acyclovir buccal) ^{CL}	 Non-preferred agents will be approved for patients who have failed a 10-day trial of ONE preferred agent within the same group
ANTI-INFLUE oseltamivir (generic Tamiflu) ^{QL} CAPS, SUSP	rimantadine (generic Flumadine) RELENZA (zanamivir) ^{QL} TAMIFLU (oseltamivir) ^{QL} CAPS, SUSP XOFLUZA (baloxavir marboxil) ^{AL,CL,QL}	 Acyclovir Susp: Prior authorization NOT required for children ≤ 12 years old Sitavig®: Approved for recurrent herpes labialis (cold sores) in immunocompetent adults Xofluza: Requires clinical, patient specific reason that a preferred agent cannot be used

ANTIVIRALS, TOPICAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
acyclovir OINT docosanol OTC	acyclovir CREAM, (generic Zovirax) DENAVIR (penciclovir) penciclovir (generic Denavir) XERESE (acyclovir/hydrocortisone)	 Non-preferred agents will be approved for patients who have failed a trial with ONE preferred ORAL Antiviral agent

ANXIOLYTICS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
alprazolam TABLET (generic for Xanax) buspirone (generic for Buspar) chlordiazepoxide diazepam TABLET , SOLN (generic for Valium) lorazepam INTENSOL , TABLET (generic for Ativan)	alprazolam ER (generic for Xanax XR) alprazolam ODT alprazolam INTENSOL ^{CL} clorazepate (generic for Tranxene-T) diazepam INTENSOL ^{CL} lorazepam ORAL SYRINGE LOREEV XR (lorazepam) ^{AL} meprobamate oxazepam	 Non-preferred agents will be approved for patients who have failed a trial with TWO preferred agents within this drug class Drug-specific criteria: Diazepam Intensol®: Requires clinical reason why diazepam solution cannot be used Alprazolam Intensol®: Requires trial of diazepam solution OR lorazepam Intensol®

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

BETA BLOCKERS, ORAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
atenolol (generic Tenormin) atenolol/chlorthalidone (generic Tenoretic) bisoprolol (generic Zebeta) bisoprolol/HCTZ (generic Ziac) BYSTOLIC (nebivolol) HEMANGEOL (propranolol) ^{AL} SOLN metoprolol (generic Lopressor) metoprolol ER (generic Toprol XL) nebivolol (generic Bystolic) propranolol (generic Inderal) propranolol ER (generic Inderal LA)	acebutolol (generic Sectral) betaxolol (generic Kerlone) INDERAL/INNOPRAN XL (propranolol ER) KAPSPARGO SPRINKLE (metoprolol ER) metoprolol/HCTZ (generic Lopressor HCT) nadolol (generic Corgard) nadolol/bendroflumethiazide pindolol (generic Viskin) propranolol/HCTZ (generic Inderide) timolol (generic Blocadren) TOPROL XL (metoprolol ER)	 Non-preferred agents will be approved for patients who have failed TWO diagnosis-appropriate preferred agents within this drug class Drug-specific criteria: Coreg CR/carvedilol: Requires clinical reason generic IR product cannot be used Hemangeol®: Covered for diagnosis of Proliferating Infantile Hemangioma Sotylize®: Covered for diagnosis of life –threatening ventricular arrhythmias OR maintenance of normal sinus rhythm in highly symptomatic atrial fibrillation/flutter (AFIB/AFL) Requires clinical reason generic sotalol cannot be used
BETA- AND ALF	PHA-BLOCKERS	
carvedilol (generic Coreg) labetalol (generic Trandate)	carvedilol ER ^{CL} (generic Coreg CR)	
ANTIARR	HYTHMIC	
sotalol (generic Betapace)	SOTYLIZE (sotalol)	

BILE SALTS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
ursodiol CAPSULE 300 mg (generic Actigall) ursodiol 250 mg TABLET (generic URSO) ursodiol 500 mg TABLET (generic URSO FORTE)	BYLVAY (odevixibat) CAP, PELLET CHENODAL (chenodiol) CHOLBAM (cholic acid) IQIRVO (elafibranor) ^{NR,QL} TAB LIVMARLI (maralixibat) SOLN ^{AL} OCALIVA (obeticholic acid) RELTONE (ursodiol 200mg,400mg) CAP	Non-preferred agents will be approved for patients who have failed a trial with ONE preferred agent within this drug class

Unless otherwise specified, the listing of a particular brand or generic name includes all dosage forms of that drug.

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

BLADDER RELAXANT PREPARATIONS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
fesoterodine (generic Toviaz) MYRBETRIQ (mirabegron) ^{AL} TAB oxybutynin IR, ER (generic Ditropan/Ditropan XL)	darifenacin ER (generic Enablex) flavoxate HCL GELNIQUE (oxybutynin) GEMTESA (vibegron)AL,QL mirabegron ER TAB (generic Myrbetriq)NR MYRBETRIQ (mirabegron) SUSPAL,CL,QL oxybutynin 2.5mg OXYTROL (oxybutynin) solifenacin (generic Vesicare) tolterodine IR, ER (generic Detrol/ Detrol LA) TOVIAZ (fesoterodine ER) trospium IR, ER (generic Sanctura/ Sanctura XR) VESICARE (solifenacin) VESICARE LS SUSP (solifenacin) AL	 Non-preferred agents will be approved for patients who have failed a trial with ONE preferred agent within this drug class Drug-specific criteria: Myrbetriq suspension: Covered for pediatric patients ≥ 3 years old with a diagnosis of Neurogenic Detrusor Overactivity (NDO)

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

BONE RESORPTION SUPPRESSION AND RELATED DRUGS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
BISPHOS	PHONATES	Non-preferred agents will be
ellendronate (generic Fosamax) TAB bandronate (generic Boniva) ^{QL}	alendronate SOLN (generic Fosamax) ^{QL} ATELVIA DR (risedronate)	approved for patients who have failed a trial of ONE preferred agent within the same group
	·	Drug-specific criteria:
	etidronate disodium (generic Didronel) FOSAMAX PLUS DQL	
	risedronate (generic Actonel) ^{QL}	 Atelvia DR[®]: Requires clinical reason alendronate cannot be
		 taken on an empty stomach Binosto[®]: Requires clinical reaso
	PRESSION AND RELATED DRUGS	 Binosto[®]: Requires clinical reaso why alendronate tablets OR
alcitonin-salmon NASAL	EVISTA (raloxifene)	Fosamax® solution cannot be use
FORTEO (teriparatide) ^{CL,QL}		 Etidronate disodium: Trial not required for diagnosis of
aloxifene (generic Evista)	TYMLOS (abaloparatide)	hetertrophic ossification
		 Forteo/ teriparatide: Covered for high risk of fracture
		High risk of fracture:
		BMD -3 or worse
		 Postmenopausal women with history of non-traumatic fractures
		 Postmenopausal women with or more clinical risk factors
		 Family history of non- traumatic fractures
		 DXA BMD T-score ≤ -2.5 any site
		 Glucocorticoid use ≥ 6 months at 7.5 dose of prednisolone equivalent
		 Rheumatoid Arthritis
		 Postmenopausal women with BMD T-score ≤ -2.5 at any sitwith any clinical risk factors
		 More than 2 units of alcohol per day
		Current smoker
		 Men with primary or hypogonadal osteoporosis
		 Osteoporosis associated with sustained systemic glucocorticoid therapy
		 Trial of calcitonin-salmon not required
		 Maximum of 24 months treatment per lifetime

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

BPH (BENIGN PROSTATIC HYPERPLASIA) TREATMENTS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
ALPHA B	ALPHA BLOCKERS	
alfuzosin (generic Uroxatral) doxazosin (generic Cardura) tamsulosin (generic Flomax) terazosin (generic Hytrin)	CARDURA XL (doxazosin) silodosin (generic Rapaflo)	approved for patients who have failed a trial of ONE preferred agent within this drug class Drug-specific criteria:
5-ALPHA-REDUCTAS	SE (5AR) INHIBITORS	Alfuzosin/dutasteride/finasteride
dutasteride (generic Avodart) finasteride (generic Proscar)	dutasteride/tamsulosin (generic Jalyn) ENTADFI (finasteride/tadalafil)	 Covered for males only Cardura XL®: Requires clinical reason generic IR form cannot be used Flomax/ tamsulosin: Females covered for a 7 day supply with diagnosis of acute kidney stones Jalyn/ dutasteride-tamsulosin: Requires clinical reason why individual agents cannot be used

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

BRONCHODILATORS, BETA AGONIST

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class
albuterol HFA (generic Proventil HFA) PROVENTIL HFA (albuterol) VENTOLIN HFA (albuterol) XOPENEX HFA (levalbuterol HFA)	albuterol HFA (generic ProAir HFA and Ventolin HFA) levalbuterol HFA (generic Xopenex HFA) PROAIR DIGIHALER (albuterol) PROAIR RESPICLICK (albuterol) ERS – Long Acting STRIVERDI RESPIMAT (olodaterol)	 Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class Drug-specific criteria: Xopenex/levalbuterol solution: Covered for cardiac diagnoses or side effect of tachycardia with albuterol product
INHAL	ATION SOLUTION	
albuterol (2.5mg/3ml premix or 2.5mg/0.5ml) albuterol 100 mg/20 mL albuterol low dose (0.63mg/3ml & 1.25mg/3ml)	arformoterol tartrate (generic Brovana) BROVANA (arformoterol) formoterol fumarate (generic Perforomist) levalbuterol (generic for Xopenex) PERFOROMIST (formoterol)	
	ORAL	
albuterol SYRUP	albuterol TAB albuterol ER (generic for Vospire ER) metaproterenol (formerly generic for Alupent) terbutaline (generic for Brethine)	

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

CALCIUM CHANNEL BLOCKERS, ORAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
SHORT-ACTING Dihydropyridines		Non-preferred agents will be approved for patients who have
	isradipine (generic Dynacirc) nicardipine (generic Cardene) nifedipine (generic Procardia) nimodipine (generic Nimotop) NYMALIZE (nimodipine) SOLN	failed a trial of ONE preferred agent within this drug class Drug-specific criteria: Nifedipine: May be approved without trial for diagnosis of Preterm Labor or Pregnancy
Non-dihydi	opyridines	Induced Hypertension (PIH)Nimodipine: Covered without trial
diltiazem (generic Cardizem) verapamil (generic Calan/Isoptin)		for diagnosis of subarachnoid hemorrhage Katerzia/ Norligva: May be
LONG-	ACTING	approved with documented
Dihydrog	pyridines	swallowing difficulty
amlodipine (generic Norvasc) nifedipine ER (generic Procardia XL/ Adalat CC)	felodipine ER (generic Plendil) KATERZIA (amlodipine) ^{QL} SUSP levamlodipine (generic Conjupri) nisoldipine (generic Sular) NORLIQVA (amolidipine) ^{AL,CL,QL} SOLN	
Non-dihydi	opyridines	
diltiazem ER (generic Cardizem CD) verapamil ER TAB	CALAN SR (verapamil) diltiazem ER (generic Cardizem LA) MATZIM LA (diltiazem ER) TIAZAC (diltiazem) verapamil ER CAPS verapamil 360mg CAPS verapamil ER (generic Verelan PM)	

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

CEPHALOSPORINS AND RELATED ANTIBIOTICS, ORAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
BETA LACTAM/BETA-LACTAMASE INHIBITOR COMBINATIONS		Non-preferred agents will be
amoxicillin/clavulanate TAB, SUSP	amoxicillin/clavulanate CHEWABLE amoxicillin/clavulanate ER (generic Augmentin XR) AUGMENTIN (amoxicillin/clavulanate)	approved for patients who have failed a 3-day trial of ONE preferred agent within the same group
	SUSP, TAB	Drug Specific Criteria • Cefixime- May be approved
CEPHALOSPORINS	S – First Generation	for a diagnosis of gonorrhea, with an appropriate ICD-10 diagnosis
cefadroxil CAPS, SUSP (generic Duricef) cephalexin CAPS, SUSP (generic Keflex)	cefadroxil TAB (generic Duricef) cephalexin TAB	code without a 3-day trial of a preferred agent Cefpodoxime- May be approved for a diagnosis of pyelonephritis, with an appropriate
CEPHALOSPORINS -	Second Generation	ICD-10 diagnosis code without a
cefprozil (generic Cefzil) cefuroxime TAB (generic Ceftin)	cefaclor (generic Ceclor) CEFTIN (cefuroxime) TAB, SUSP	3-day trial of a preferred agent
CEPHALOSPORINS -	- Third Generation	
cefdinir (generic Omnicef)	cefixime (generic Suprax) CAPS, SUSP cefpodoxime (generic Vantin) SUPRAX (cefixime) CAPS, CHEWABLE TAB, SUSP, TAB	

COLONY STIMULATING FACTORS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
FYLNETRA (pegfilgrastim-pbbk) NEUPOGEN DISP SYR NEUPOGEN (filgrastim) VIAL	FULPHILA (pegfilgrastim-jmdb) SUB-Q GRANIX (tbo-filgrastim) LEUKINE (sargramostim) NEULASTA (pegfilgrastim) SYR NIVESTYM (filgrastim-aafi) SYR,VIAL NYVEPRIA (pegfilgrastim-apgf) RELEUKO (filgrastim-ayow) SYR,VIAL STIMUFEND (pegfilgrastim-fpgk) UDENYCA (pegfilgrastim-cbqv) AUTOINJ UDENYCA (pegfilgrastim-cbqv) SUB-Q ZARXIO (filgrastim-sndz) ZIEXTENZO SYR (pegfilgrastim-bmez)	Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class

Unless otherwise specified, the listing of a particular brand or generic name includes all dosage forms of that drug.

CL - Prior Authorization / Class Criteria apply

QL - Quantity/Duration Limit

AL- Age Limit

NR - Product was not reviewed - New Drug criteria will apply

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

CONTRACEPTIVES, ORAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class
All reviewed agents are recommended preferred at this time Only those products for review are listed. Brand name products may be subject to Maximum Allowable Cost (MAC) pricing or require substitution with a generic equivalent Specific agents can be looked up using the Drug Look-up Tool at: https://druglookup.fhsc.com/druglookupweb/?client=nestate	Non-Preferred Agents EMZAHH (norethindrone) ^{NR}	Prior Authorization/Class

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

COPD (CHRONIC OBSTRUCTIVE PULMONARY DISEASE) AGENTS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
ANORO ELLIPTA (umeclidinium/vilanterol) ATROVENT HFA (ipratropium) COMBIVENT RESPIMAT (albuterol/ipratropium) SPIRIVA (tiotropium) STIOLTO RESPIMAT (tiotropium/olodaterol)	BEVESPI AEROSPHERE (glycopyrolate/formoterol) DUAKLIR PRESSAIR (aclidinium br and formoterol fum) INCRUSE ELIPTA (umeclidnium) SPIRIVA RESPIMAT (tiotropium)	 Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class OR Patient specific documentation of inability to use traditional inhaler device. Drug-specific criteria: Daliresp/roflumilast: Covered for diagnosis of severe COPD associated with chronic bronchitis Requires trial of a bronchodilator Requires documentation of one
INHALATIO	N SOLUTION	 exacerbation in last year upon initial review
albuterol/ipratropium (generic Duoneb) ipratropium SOLN (generic Atrovent)	LONHALA (glycopyrrolate inhalation soln) YUPELRI (revefenacin)	
ORAL	AGENT	
roflumilast (generic Daliresp) ^{CL,QL}	DALIRESP (roflumilast) ^{CL, QL}	

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

COUGH AND COLD, OPIATE COMBINATION

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
	guaifenesin/codeine LIQUID hydrocodone/homatropine SYRUP promethazine/codeine SYRUP promethazine/phenylephrine/codeine SYRUP	 Non-preferred agents will be approved for patients who have failed a trial of ONE dextromethorphan product All codeine or hydrocodone containing cough and cold combinations are limited to ≥ 18 years of age

CYSTIC FIBROSIS, ORAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
	BRONCHITOL (mannitol) AL,CL,QL KALYDECO PACKET, TAB (ivacaftor)QL, AL ORKAMBI (lumacaftor/ivacaftor) PACKET, TAB QL, AL SYMDEKO (tezacaftor/ivacaftor)QL, AL TRIKAFTA(elexacaftor, tezacaftor, ivacaftor)AL, CL PACKETCL, TAB	 Bronchitol: Approved for diagnosis of CF and documentation that the patient has passed the BRONCHITOL Tolerance Test Kalydeco®: Diagnosis of CF and documentation of the drug-specific FDA-approved mutation of CFTR gene Orkambi®: Diagnosis of CF and documentation of presence of two copies of the F580del mutation (homozygous) of CFTR gene Symdeko: Diagnosis of CF and documentation of the drug specific FDA approved mutation of CFTR gene. Trikafta: Diagnosis of CF and documentation of at least one F508del mutation in the CFTR gene

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

CYTOKINE & CAM ANTAGONISTS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
COSENTYX (secukinumab) ^{AL} PEN, SYRINGE ENBREL (etanercept) KIT, MINI CART, PEN, SYRINGE, VIAL QL HUMIRA (adalimumab) ^{QL} DTEZLA (apremilast) TAB ^{CL,QL}	ABRILADA KIT (adalimumabafzb)AL,NR (CF) ABRILADA PEN KIT (adalimumabafzb)AL,NR (CF) ACTEMRA (tocilizumab) SUB-Q ADALIMUMAB-AACF (CF)AL,NR PEN KIT ADALIMUMAB-AATY (CF)AL,NR PEN KIT ADALIMUMAB-ADAZ(CF)(biosim for Hyrimoz)AL PEN,SYRINGE ADALIMUMAB-ADBM(CF) PEN CROHNSAL,NR ADALIMUMAB-ADBM(CF) KIT, PEN ALIMUMAB-ADBM(CF) KIT, PEN PS-UV ADALIMUMAB-ADBM(CF) KIT, PENAL,NR ADALIMUMAB-FKJP (biosim for Hulio)AL PEN, SYRINGE ADALIMUMAB-RYVKAL,NR (biosim for Simlandi) KIT AMJEVITA (adalimumab-atto)AL AUTOINJ, SYR AMJEVITA(adalimumab-atto)AL,NR FEN KIT ARCALYST (nilonacept) BIMZELX (bimekizumab-bkzx)AL,NR PEN, SYR CIBINQO (abrocitinib)AL,QL CIMZIA (certolizumab pegol)QL CYLTEZO (adalimumab-adbm)AL KIT, PEN-PSORIASIS ENSPRYNG (satralizumab-mwge) SUB-Q ENTYVIO (vedolizumab)AL,NR PEN	 Preferred agents will be approved with FDA-approved indication – ICD-10 diagnosis code is required. Non-preferred agents will be approved for FDA-approved indications in patients who have failed a trial of TWO preferred agents within this drug class, or upon diagnosis for non-preferred agent with FDA-approved indication if no preferred agent has FDA approval for diagnosis. JAK-Inhibitors: For FDA approved indications that require a patient to have had an inadequate response to a TNF blocker, documentation of an inadequate response is required. Drug-specific criteria: Cosentyx: Requires treatment failure of Enbrel OR Humira with the same FDA approved indications and age limits. Otezla: Requires a trial of Humira

Unless otherwise specified, the listing of a particular brand or generic name includes all dosage forms of that drug.

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

CYTOKINE & CAM ANTAGONISTS, continued

Non-Preferred Agents HADLIMA (adalimumab- bwwd) ^{AL} PUSHTOUCH, SYRINGE	 Prior Authorization/Class Criteria Preferred agents will be approved with FDA-approved indication —
HADLIMA (CF) (adalimumab- bwwd) ^{AL} PUSHTOUCH, SYRINGE HULIO (adalimumab-fkjp) ^{AL} PEN, SYRINGE HYRIMOZ(CF) (adalimumab-adaz) ^{AL} PEN, SYRINGE IDACIO (adalimumab-aacf) ^{AL} PEN, SYRINGE ILUMYA (tildrakizumab) SUB-Q KEVZARA (sarilumab) SUB-Q, PEN, SYRINGE KINERET (anakinra) LITFULO (ritlecitinib) ^{AL,NR} CAPS OLUMIANT (baricitinib) TAB ^{CL,QL} OMVOH (mirikizumab-mrkz) ^{AL,NR} PEN SYRINGE ^{NR} ORENCIA (abatacept) SUB-Q RINVOQ ER (upadacitinib) ^{CL,QL} RINVOQ (upadacitinib) ^{AL,NR,QL} LQ	ICD-10 diagnosis code is required
	SYRINGE HYRIMOZ(CF) (adalimumab-adaz)AL PEN, SYRINGE IDACIO (adalimumab-aacf)ALPEN, SYRINGE ILUMYA (tildrakizumab) SUB-Q KEVZARA (sarilumab) SUB-Q, PEN, SYRINGE KINERET (anakinra) LITFULO (ritlecitinib)AL,NR CAPS OLUMIANT (baricitinib) TABCL,QL OMVOH (mirikizumab-mrkz)AL,NR PEN SYRINGENR ORENCIA (abatacept) SUB-Q RINVOQ ER (upadacitinib)CL,QL RINVOQ (upadacitinib)AL,NR,QL LQ SOLN SILIQ (brodalumab) SIMLANDI (CF) (adalimumab-ryvk)AL,NR KIT SIMPONI (golimumab) SKYRIZI (risankizamab-rzaa) SYR SKYRIZI ON-BODY (risankizamab-rzaa)QL SKYRIZI PEN (risankizamab-rzaa)QL SOTYKTU (deucravacitinib) TAB SPEVIGO (spesolimab-sbzo)AL,NR SYR STELARA (ustekinumab) SUB-Q TALTZ (ixekizumab)AL TREMFYA (guselkumab)QL TYENNE (tocilizumab-aazg)AL,NR AUTOINJ TYENNE (tocilizumab-aazg)AL,NR

Unless otherwise specified, the listing of a particular brand or generic name includes all dosage forms of that drug.

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
	VELSIPITY (etrasimod) ^{NR,QL} TAB XELJANZ (tofacitinib) TAB, SOLN ^{CL,QL} XELJANZ XR (tofacitinib) TAB ^{CL,QL} YUFLYMA 100mg/mL (CF) (adalimumab- aaty) ^{AL} KIT,PEN KIT YUFLYMA 80mg/mL (CF) (adalimumab- aaty) ^{AL,NR} AUTOINJ, PEN, KIT YUSIMRY (CF) (adalimumab- aqvh) ^{AL} PEN KIT ZYMFENTRA PEN, SYR (infliximab-dyyb) ^{NR}	 Preferred agents will be approved with FDA-approved indication – ICD-10 diagnosis code is required. Non-preferred agents will be approved for FDA-approved indications in patients who have failed a trial of TWO preferred agents within this drug class, or upon diagnosis for non-preferred agent with FDA-approved indication for preferred agent has FDA approval for diagnosis. JAK-Inhibitors: For FDA approved indications that require a patient to have had an inadequate response to a TNF blocker, documentation of an inadequate response is required. Drug-specific criteria: Cosentyx: Requires treatment failure Enbrel OR Humira with the same FDA approved indications and age limits. Otezla: Requires a trial of Humira

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

DIURETICS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
amiloride TAB bumetanide TAB chlorthalidone TAB (generic Diuril) furosemide SOLN, TAB (generic Lasix) hydrochlorothiazide CAPS, TAB	CAROSPIR (spironolactone) SUSP eplerenone TAB (generic Inspra)CL ethacrynic acid CAPS (generic Edecrin) KERENDIA (finerenone) TAB CL,QL spironolactone (generic Carospir) SUSP THALITONE (chlorthalidone) TAB triamterene (generic Dyrenium)	 Non-preferred agents will be approved for patients who have failed a trial of TWO preferred agents within this drug class Eplerenone: Will be approved with a failed trial or intolerance to spironolactone, a trial with two preferred agents is not required. Kerendia: For diagnosis of chronic kidney disease associated with Type-II diabetes in adults, trial of a preferred agent not required. spironolactone suspension: May be approved without trial of a
COMBINATIO	N PRODUCTS	preferred agent if there is a clinical
amiloride/HCTZ TAB spironolactone/HCTZ TAB (generic Aldactazide) triamterene/HCTZ CAPS , TAB (generic Dyazide, Maxzide)		reason why preferred spironolactone solid dosage form cannot be used.

ENZYME REPLACEMENT, GAUCHER'S DISEASE

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
ZAVESCA (miglustat) ^{CL}	CERDELGA (eliglustat) miglustat (generic Zavesca)	 Non-preferred agents require clinical documentation why the preferred product within this drug class is not appropriate Drug-specific criteria: Zavesca/miglustat: Approved for mild to moderate type 1 Gaucher disease for whom enzyme replacement therapy is not a therapeutic option

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

EPINEPHRINE, SELF-INJECTED QL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
AUVI-Q 0.1mg (epinephrine) epinephrine (AUTHORIZED GENERIC Epipen/ Epipen Jr.) AUTOINJ EPIPEN (epinephrine) AUTOINJ EPIPEN JR. (epinephrine) AUTOINJ	AUVI-Q 0.15mg (epinephrine) AUVI-Q 0.3mg (epinephrine) epinephrine (generic for Adrenaclick) epinephrine (generic for Epipen/ Epipen Jr.) AUTOINJ SYMJEPI (epinephrine) PFS	Non-preferred agents require clinical documentation why the preferred product within this drug class is not appropriate

ERYTHROPOIESIS STIMULATING PROTEINS

Preferred Agents	Non-Preferred Agents		Prior Authorization/Class Criteria
ARANESP (darbopoetine alfa) DISP SYR, VIAL EPOGEN (rHuEPO) RETACRIT (epoetin alfa-epbx) <i>Pfizer manufacturer only</i>	JESDUVROQ (daprodustat) ^{NR} TAB PROCRIT (rHuEPO) RETACRIT (epoetin alfa-epbx) <i>Vifor</i> manufacturer only	•	Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class

FLUOROQUINOLONES, ORAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
ciprofloxacin TAB (generic Cipro) levofloxacin TAB (generic Levaquin)	BAXDELA (delafloxacin) ciprofloxacin ER ciprofloxacin SUSP (generic Cipro) levofloxacin SOLN moxifloxacin (generic Avelox) ofloxacin	 Non-preferred agents will be approved for patients who have failed a 3-day trial of ONE preferred agent within this drug class Drug-specific criteria: Baxdela: Coverable with documented intolerance or failure of preferred MRSA agents (clindamycin, doxycycline, linezolid, sulfamethoxazole/trimethoprim) Ciprofloxacin/Levofloxacin Suspension: Coverable with documented swallowing disorders Ofloxacin: Trial of preferred not required for diagnoses of Pelvic Inflammatory Disease OR Acute Epididymitis (nongonorrhea)

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

GI MOTILITY, CHRONIC

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
AMITIZA (lubiprostone)AL, QL LINZESS (linaclotide)AL,QL MOVANTIK (naloxegol oxalate)QL RELISTOR (methylnaltrexone) SYR TRULANCE (plecanatide)QL	alosetron (generic Lotronex) IBSRELA (tenapanor) ^{AL,QL} lubiprostone (generic Amitiza) ^{AL,QL} MOTEGRITY (prucalopride succinate) RELISTOR (methylnaltrexone) ^{QL} TAB, VIAL SYMPROIC (naldemedine) VIBERZI (eluxodoline)	 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 Drug-specific criteria: Ibsrela: May be approved for diagnosis of IBS with constipation after trial of at least TWO OTC laxatives (senna, bisacodyl, etc.) Lotronex/ alosetron: Covered for diagnosis of IBS Diarrhea Predominant type with trial and failure of loperamide AND diphenoxylate Relistor® TAB: Covered for diagnosis of opioid-induced constipation in adults with chronic, non-cancer pain after trial of at least TWO OTC laxatives (senna, bisacodyl, etc.) and failure of Movantik Symproic: Covered for diagnosis of opioid-induced constipation in adult patients with chronic non-cancer pain after trial on at least TWO OTC laxatives and failure of Movantik Viberzi®: Covered for diagnosis of IBS Diarrhea Predominant type with trial and failure of loperamide AND diphenoxylate

GLUCAGON AGENTS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
BAQSIMI (glucagon) ^{AL,QL} NASAL GLUCAGON EMERGENCY (glucagon) ^{QL} INJ KIT (Lilly) glucagon ^{QL} INJ PROGLYCEM (diazoxide) SUSP ZEGALOGUE (dasiglucagon) ^{AL, QL} AUTO-INJ	diazoxide SUSP (generic Proglycem) GLUCAGON EMERGENCY (glucagon) ^{QL} INJ KIT (Fresenius) GVOKE (glucagon) ^{AL,QL} KIT , PEN , SYR , VIAL ZEGALOGUE (dasiglucagon) ^{AL,QL} SYR	 Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class

Unless otherwise specified, the listing of a particular brand or generic name includes all dosage forms of that drug.

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

GLUCOCORTICOIDS, INHALED

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
GLUCOCO ARNUITY ELLIPTA (fluticasone) ASMANEX (mometasone) ^{QL,AL} ASMANEX HFA (mometasone) FLOVENT HFA (fluticasone) PULMICORT FLEXHALER (budesonide)	ALVESCO (ciclesonide)AL,CL ARMONAIR DIGIHALER (fluticasone)AL,QL FLOVENT DISKUS (fluticasone) fluticasone (generic Flovent Diskus)NR fluticasone HFA (generic Flovent HFA)CL QVAR Redihaler (beclomethasone)	 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 this drug class within the last 6 months budesonide respules: 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.
ADVAIR DISKUS (fluticasone/salmeterol) ^{QL} ADVAIR HFA (fluticasone/salmeterol) ^{QL} DULERA (mometasone/formoterol) SYMBICORT (budesonide/ formoterol) TRELEGY ELLIPTA (fluticasone/umeclidinium/vilanterol)	AIRDUO DIGIHALER (fluticasone/salmeterol)AL,QL AIRSUPRA HFA (albuterol and budesonide)AL BREO ELLIPTA (fluticasone/vilanterol) BREZTRI (budesonide/formoterol/glycopyrrolate)QL budesonide/formoterol (generic for Symbicort) fluticasone/salmeterol (generic for Advair Diskus)QL fluticasone/salmeterol (generic for Advair HFA)QL fluticasone/salmeterol (generic for Airduo Respiclick) fluticasone/vilanterol (Breo Ellipta) WIXELA INHUB (generic for Advair Diskus)QL	• fluticasone HFA: Covered without PA for age ≤ 8 years
INHALATION	budesonide RESPULES (generic for Pulmicort)	

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

GLUCOCORTICOIDS, ORAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
budesonide EC CAPS (generic Entocort EC) dexamethasone ELIXIR, SOLN dexamethasone TAB hydrocortisone TAB methylprednisolone tablet (generic Medrol) prednisolone SOLN prednisolone sodium phosphate prednisone DOSE PAK prednisone TAB	ALKINDI (hydrocortisone) GRANULES ^{AL} CORTEF (hydrocortisone) cortisone TAB dexamethasone INTENSOL ENTOCORT EC (budesonide) EOHILIA (budesonide) ^{AL,NR,QL} SUSP HEMADY (dexamethasone) methylprednisolone 8mg, 16mg, 32mg ORTIKOS ER (budesonide) ^{AL,QL} prednisolone sodium phosphate (generic Millipred/Veripred) prednisolone sodium phosphate ODT prednisone SOLN prednisone INTENSOL RAYOS DR (prednisone) TAB TARPEYO (budesonide) CAPS	 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 Drug-specific criteria: Intensol Products: Patient specific documentation of why the less concentrated solution is not appropriate for the patient Tarpeyo: Indicated for the treatment of primary immunoglobulin A nephropathy (IgAN)

GROWTH HORMONES

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
GENOTROPIN (somatropin)	HUMATROPE (somatropin)	Growth Hormone PA Form
NORDITROPIN (somatropin)	NGENLA (somatrogon-ghla) ^{AL}	Growth Hormone Criteria
	NUTROPIN AQ (somatropin)	
	OMNITROPE (somatropin)	
	SAIZEN (somatropin)	
	SEROSTIM (somatropin)	
	SKYTROFA (lonapegsomatropin-tcgd)	
	SOGROYA (somapacitan-beco)	
	ZOMACTON (somatropin)	
	ZORBTIVE (somatropin)	

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

H. PYLORI TREATMENTS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
PYLERA (bismuth, metronidazole, tetracycline) ^{QL}	lansoprazole/amoxicillin/clarithromycin (generic Prevpac) ^{QL} OMECLAMOX-PAK (omeprazole, clarithromycin, amoxicillin) ^{QL} bismuth,metronidazole,tetracycline (generic Pylera) ^{QL} TALICIA (omeprazole/amoxicillin/rifabutin) VOQUEZNA (vonoprazan) ^{QL}	Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class

HAE TREATMENTS CL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
Preferred Agents BERINERT (C1 esterase inhibitor, human) INTRAVENOUS HAEGARDA (C1 esterase inhibitor, human) AL.CL SUB-Q icatibant acetate (generic for FIRAZYR) AL SUB-Q	CINRYZE (C1 esterase inhibitor, human) ^{AL,CL} INTRAVENOUS FIRAZYR (icatibant acetate) ^{AL} SUB-Q ORLADEYO (berotralstat) CAP ^{AL,QL} RUCONEST (recombinant human C1 inhibitor) ^{AL} INTRAVENOUS TAKHZYRO (lanadelumab-flyo) ^{AL,CL} VIAL	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 estrogencontaining products is contraindicated Non-preferred agents will be
	TAKHZYRO (lanadelumab-flyo) ^{AL,CL} SYRINGE	
		 Drug-Specific Criteria Cinryze, Haegarda, Orladeyo, and Takhzyro, require a history of two or more HAE attacks monthly

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

HEMOPHILIA TREATMENTS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
F/	ACTOR VIII	 Non-preferred agents will be
ALPHANATE HUMATE-P KOVALTRY NOVOEIGHT NUWIQ XYNTHA KIT, SOLOFUSE	ADVATE ADYNOVATE AFSTYLA ALTUVIIIO ELOCTATE ESPEROCT HEMOFIL-M JIVIAL KOATE-DVI KIT KOATE-DVI VIAL KOGENATE FS OBIZUR RECOMBINATE	approved for patients who have failed a trial of ONE preferred agent within this drug class
F	ACTOR IX	
ALPROLIX	ALPHANINE SD	
BENEFIX	IDELVION IXINITY MONONINE PROFILNINE SD REBINYN RIXUBIS	
FACTOR VIIa AND PROTHRO	OMBIN COMPLEX-PLASMA DERIVED	
NOVOSEVEN RT	FEIBA NF SEVENFACT ^{AL}	
FACTOR X	AND XIII PRODUCTS	
COAGADEX CORIFACT	TRETTEN	
VON WILLE	BRAND PRODUCTS	
WILATE	VONVENDI	
BISPEC	CIFIC FACTORS	
HEMLIBRA		

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

HEPATITIS B TREATMENTS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
entecavir TAB	adefovir dipivoxil BARACLUDE (entecavir) SOLN, TAB EPIVIR HBV (lamivudine) TAB, SOLN lamivudine hbv TAB VEMLIDY (tenofovir alafenamide fumarate)	 Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class Drug Specific Criteria tenofovir disoproxil fumarate (generic Viread) tablet: Diagnosis for use required. May be indicated for chronic hepatitis B or HIV-1 infection. See HIV/AIDS class for drug listing and placement

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

HEPATITIS C TREATMENTS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
DIRECT ACTING ANTI-VIRAL		Hepatitis C Treatments PA Form
sofosbuvir/velpatasvir (generic Epclusa) ^{CL} MAVYRET (glecaprevir/pibrentasvir) TAB ^{CL} , PELLET ^{AL,CL} VOSEVI (sofosbuvir/velpatasvir/ voxilaprevir) ^{CL}	HARVONI 200/45MG, TAB (ledipasvir/sofosbuvir) ^{CL} HARVONI (ledipasvir/sofosbuvir) ^{CL} PELLET ledipasvir/sofosbuvir (generic Harvoni) ^{CL} SOVALDI (sofosbuvir) ^{CL} PELLET SOVALDI TAB (sofosbuvir) ^{CL} VIEKIRA PAK (ombitasvir/paritaprevir/ritonavir/dasabuvir) ^{CL} ZEPATIER (elbasvir/grazoprevir) ^{CL}	Hepatitis C Criteria
RIBA	VIRIN	genotype 1 or 4
ribavirin 200mg CAPSULE, TAB		Vosevi: Requires documentation of non-response after previous treatment course of Direct Acting
INTERFERON		Anti-viral agent (DAA) for genotype
PEGASYS (pegylated interferon alfa- 2a) ^{CL}		1-6 without cirrhosis or with compensated cirrhosis

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

HISTAMINE II RECEPTOR BLOCKERS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
famotidine TAB (generic for Pepcid) famotidine SUSP	cimetidine TAB , SOLN ^{CL} (generic Tagamet) famotidine ^{NR} CHEW-TAB nizatidine CAPS (generic for Axid)	 Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class
		Drug-specific criteria: Cimetidine: Approved for viral M. contagiosum or common wart V. Vulgaris treatment

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

HIV / AIDS CL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
CAPSID	INHIBITOR	All agents require:
	SUNLENCA (lenacapavir) ^{QL}	 Diagnosis of HIV/AIDS required, OR
CCR5 AN	TAGONISTS	 Diagnosis of Pre and Post
SELZENTRY SOLN, TAB (maraviroc)	maraviroc (generic Selzentry)	 Exposure Prophylaxis Non-preferred agents will be approved for patients who have a
FUSION	INHIBITORS	diagnosis of HIV/AIDS and patient
FUZEON SUB-Q (enfuvirtide) ^{QL}		specific documentation of why the preferred products within this drug
HIV-1 ATTACH	MENT INHIBITOR	class are not appropriate for patient, including, but not limited
	RUKOBIA ER (fostemsavir) ^{AL,QL}	 to, drug resistance or concomitant conditions not recommended with preferred agents
INTEGRASE STRAND TRA	NSFER INHIBITORS (INSTIs)	 Patients undergoing treatment at
ISENTRESS (raltegravir) ^{QL} ISENTRESS HD (raltegravir) TIVICAY (dolutegravir)	TIVICAY PD (dolutegravir)	the time of any preferred status change will be allowed to continue therapy
NON-NUCLEOSIDE REVERSE TRA	NSCRIPTASE INHIBITORS (NNRTIs)	-
efavirenz CAPS , TABLET (generic Sustiva) INTELENCE (etravirine) ^{QL} PIFELTRO (doravirine) ^{QL}	etravirine (generic Intelence) ^{QL} nevirapine IR, ER (generic Viramune/Viramune XR) basglaSUSTIVA CAPS, TABLET (efavirenz) VIRAMUNE (nevirapine) SUSP	
NUCLEOSIDE REVERSE TRAN	SCRIPTASE INHIBITORS (NRTIs)	
abacavir SOLN, TABLET (generic Ziagen) EMTRIVA CAPS, SOLN (emtricitabine) lamivudine SOLN, TABLET (generic Epivir) zidovudine CAPS, SYRUP, TABLET (generic Retrovir)	didanosine DR (generic Videx EC) emtricitabine CAPS (generic for Emtriva) EPIVIR (lamivudine) RETROVIR (zidovudine) stavudine CAPS (generic Zerit) ZIAGEN (abacavir)	
NUCLEOTIDE REVERSE TRAN	ISCRIPTASE INHIBITORS (NRTIs)	
tenofovir TABLET (generic Viread)	VIREAD (tenofovir) POWDER	
PHARMACOKIN	IETIC ENHANCER	
	TYBOST (cobicistat) ^{QL}	

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

HIV / AIDS CL (Continued)

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
PROTEASE INHIBITORS		All agents require:
atazanavir CAPS (generic Reyataz) NORVIR (ritonavir) TAB PREZISTA (darunavir) TAB ritonavir TAB (generic Norvir)	APTIVUS CAPS, SOLN (tipranavir) CRIXIVAN (indinavir) DARUNAVIR PROPYLENE GLYCOLATEAL TAB darunavir ethanolate (generic Prezista)AL TAB fosamprenavir TAB (generic Lexiva) LEXIVA SUSP (fosamprenavir) LEXIVA TAB (fosamprenavir) NORVIR POWDER, SOLN (ritonavir) PREZISTA (darunavir) SUSP REYATAZ POWDER (atazanavir) VIRACEPT (nelfinavir)	 Diagnosis of HIV/AIDS required, OR Diagnosis of Pre and Post Exposure Prophylaxis 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 patient, including, but not limited to drug resistance or concomitant conditions not recommended with preferred agents Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy
	KALETRA SOLN (lopinavir/ritonavir) KALETRA TAB (lopinavir/citonavir) PREZCOBIX (darunavir/cobicistat) ALETRA TAB (lopinavir/cobicistat)	 All agents require: Diagnosis of HIV/AIDS required; OR Diagnosis of Pre and Post Exposure Prophylaxis 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 patient, including, but not limited to drug resistance or concomitant conditions not recommended with preferred agents Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy
abacavir/lamivudine (generic Epzicom) CIMDUO (lamivudine/tenofovir)QL DESCOVY (emtricitabine/tenofovir)QL emtricitabine/tenofovir (generic Truvada) lamivudine/zidovudine (generic Combivir)	abacavir/lamivudine/zidovudine (generic Trizivir) COMBIVIR (lamivudine/zidovudine) EPZICOM (abacavir sulfate/lamivudine) TRIZIVIR (abacavir/lamivudine/zidovudine) TRUVADA (emtricitabine/tenofovir)	

Unless otherwise specified, the listing of a particular brand or generic name includes all dosage forms of that drug.

CL - Prior Authorization / Class Criteria apply

QL - Quantity/Duration Limit

AL- Age Limit

NR - Product was not reviewed - New Drug criteria will apply

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

HIV / AIDS CL (Continued)

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
COMBINATION PRODU	CTS – MULTIPLE CLASSES	 All agents require:
BIKTARVY (bictegravir/emtricitabine/ tenofovir) ^{QL} COMPLERA (rilpivirine/emtricitabine/tenofovir) DELSTRIGO (doravirine/lamivudine/tenofovir) ^{QL} DOVATO (dolutegravir/lamivudine) ^{QL} efavirenz/emtricitabine/tenofovir (generic Atripla) ^{CL} GENVOYA (elvitegravier/cobicistat/ emtricitabine/tenofovir) ^{QL, AL} JULUCA (dolutegravir/rilpivirine) ^{QL} ODEFSEY (emtricitabine/rilpivirine/ tenofovir) ^{QL} STRIBILD (elvitegravir/cobicistat/ emtricitabine/tenofovir) ^{QL} SYMFI (efavirenz/lamivudine/ tenofovir) ^{QL} SYMFI LO (efavirenz/lamivudine/ tenofovir) ^{QL} SYMTUZA (darunavir/cobicistat/ emtricitabine/tenofovir) ^{QL} TRIUMEQ (dolutegravir/abacavir/ lamivudine)	ATRIPLA (efavirenz/emtricitabine/tenofovir) efavirenz/lamivudine/tenofovir (generic for Symfi) ^{QL} efavirenz/lamivudine/tenofovir (generic for Symfi Lo) ^{QL} TRIUMEQ PD (abacavir, dolutegravir, and lamivudine) SUSP	 Diagnosis of HIV/AIDS required, OR Diagnosis of Pre and Post Exposure Prophylaxis 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 patient, including, but not limited to, drug resistance or concomitant conditions not recommended with preferred agents Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy

HYPOGLYCEMICS, ALPHA-GLUCOSIDASE INHIBITORS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
acarbose (generic for Precose)	miglitol (generic for Glyset) GLYSET (miglitol)	 Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

HYPOGLYCEMICS, INCRETIN MIMETICS/ENHANCERS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
GLUCAGON-LIKE PEPTIDE-1 RE	CEPTOR AGONIST (GLP-1 RA)CL	GLP-1 RA Criteria
OZEMPIC (semaglutide) ^{QL} TRULICITY (dulaglutide) VICTOZA (liraglutide) subcutaneous	ADLYXIN (lixisenatide) BYDUREON BCISE PEN (exenatide) ^{QL} BYETTA (exenatide) subcutaneous MOUNJARO (tirzepatide) PEN RYBELSUS (semaglutide)	Preferred agents require a diagnosis of Type II diabetes AND a trial and failure or intolerance to metformin OR A diagnosis of ASCVD associated with a diagnosis of Type II diabetes (no metformin trial required) Non-preferred agents will be approved for patients who have: • Failed a trial of TWO preferred agents within GLP-1 RA AND
INSULIN/GLP-1 RA	A COMBINATIONS	Diagnosis of diabetes with HbA1C AND
	SOLIQUA (insulin glargine/lixisenatide) XULTOPHY (insulin degludec/liraglutide)	 ≥ 7 AND Trial of metformin, or contraindication or intolerance to metformin
AMYLIN	ANALOG	Amylin Analog Criteria
	SYMLIN (pramlintide) subcutaneous -4 (DPP-4) INHIBITORAL,QL alogliptin (generic Nesina) alogliptin/metformin (generic Kazano) alogliptin/pioglitazone (generic Oseni) GLYXAMBI (empagliflozin/linagliptin) JENTADUETO XR (linagliptin/metformin) KOMBIGLYZE XR (saxagliptin/metformin) ONGLYZA (saxagliptin) QTERN (dapagliflozin/saxagliptin) saxagliptin (generic Onglyza) saxagliptin/metformin ER (generic Kombiglyze ER) sitagliptin (generic Zituvio)NR sitagliptin (generic Zituvio)NR STEGLUJAN (ertugliflozin/sitagliptin) TRIJARDY XR (empagliflozin/linagliptin/metformin) ZITUVIO (sitagliptin)	ALL criteria must be met Concurrent use of short-acting mealtime insulin Current therapy compliance No diagnosis of gastroparesis HbA1C ≤ 9% within last 90 days Monitoring of glucose during initiation of therapy DPP-4 Inhibitor Criteria Preferred agents require a diagnosis of Type II diabetes AND a trial and failure or intolerance to metformin. Non-preferred DPP-4s will be approved for patients who have failed a trial of ONE preferred agent within the DPP-4 inhibitor class

Unless otherwise specified, the listing of a particular brand or generic name includes all dosage forms of that drug.

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

HYPOGLYCEMICS, INSULIN AND RELATED DRUGS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
VIAL HUMALOG (insulin lispro) U-100 CARTRIDGE, PEN, VIAL HUMALOG JR. (insulin lispro) U-100 KWIKPEN HUMALOG MIX VIAL (insulin lispro/lispro protamine) HUMALOG MIX KWIKPEN (insulin lispro/lispro protamine) HUMALOG MIX KWIKPEN (insulin lispro/lispro protamine) HUMULIN (insulin) VIAL HUMULIN 70/30 VIAL HUMULIN U-500 VIAL HUMULIN 500 U/M PENCL	ADMELOG (insulin lispro) PEN, VIAL AFREZZA (regular insulin) INHALATION BASAGLAR (insulin glargine, rec) PEN, TEMPO PEN FIASP (insulin aspart) CARTRIDGE, PEN, VIAL HUMALOG U-100 TEMPO PEN HUMALOG (insulin lispro) ^{CL} U-200 KWIKPEN insulin degludec (generic Tresiba) 100U/mL PEN, VIAL insulin degludec (generic Tresiba) 200U/mL PEN insulin glargine (Toujeo) insulin glargine max (Toujeo Max) insulin glargine-YFGN PEN, VIAL (generic for Semglee-YFGN) insulin lispro/lispro protamine KWIKPEN (Humalog Mix Kwikpen) LYUMJEV KWIKPEN, VIAL(insulin lispro-aabc) LYUMJEV (insulin lispro-aabc) TEMPO PEN NOVOLIN (insulin) NOVOLIN 70/30 VIAL(insulin) NOVOLOG MIX (insulin aspart/aspart protamine) VIAL REZVOGLAR (insulin glargine-aglr) KWIKPEN SEMGLEE (insulin glargine) PEN, VIAL TOUJEO SOLOSTAR (insulin glargine) TRESIBA (insulin degludec)	 Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class Drug-specific criteria: Afrezza®: Approved for T1DM on long-acting insulin with no current history of smoking or chronic lung disease Humulin® R U-500 Kwikpen: May be approved for patients who require >200 units/day Humalog U-200 Pen: May be approved for patients who require > 100 units/day AND using an insulin pump

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

HYPOGLYCEMICS, MEGLITINIDES

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
repaglinide (generic for Prandin)	nateglinide (generic for Starlix) ^{CL}	 Non-preferred agents will be approved for patients with: Failure of a trial of ONE preferred agent in another Hypoglycemic class OR T2DM and inadequate glycemic control

HYPOGLYCEMICS. METFORMINS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
metformin IR & ER (generic Glucophage/Glucophage XR)	metformin ER (generic Fortamet/Glumetza) metformin SOLN (generic Riomet) RIOMET ER (metformin ER) ^{AL}	 Metformin ER (generic Fortamet®)/Glumetza®: Requires clinical reason why generic Glucophage XR® cannot be used Metformin solution: Prior authorization not required for age <7 years

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

HYPOGLYCEMICS, SGLT2

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

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

HYPOGLYCEMICS, SULFONYLUREAS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
glimepiride (generic Amaryl) glipizide IR & ER (generic Glucotrol/ Glucotrol XL) glyburide (generic Diabeta/Glynase)	chlorpropamide tolazamide 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 glyburide/metformin (generic Glucovance)		

HYPOGLYCEMICS, TZD

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
THIAZOLIDINEDIONES (TZDs)		 Non-preferred agents will be
pioglitazone (generic for Actos)		approved for patients who have failed a trial of THE preferred agent
TZD COMBINATIONS		within this drug class
	pioglitazone/glimepiride (generic for Duetact) pioglitazone/metformin (generic for Actoplus Met)	 Combination products: Require clinical reason why individual ingredients cannot be used

IDIOPATHIC PULMONARY FIBROSIS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
OFEV (nintedanib esylate) ^{CL} pirfenidone (generic Esbriet) ^{QL}	ESBRIET (pirfenidone) ^{QL}	 Non-preferred agent requires trial of preferred agent within this drug class FDA approved indication required – ICD-10 diagnosis code

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

IMMUNOMODULATORS, ASTHMA CL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
FASENRA (benralizumab) ^{AL} PEN XOLAIR (omalizumab)	NUCALA (mepolizumab) ^{AL} AUTO-INJ , SYR	Immunomodulators Self-Injectable PA Form
AUTO-INJAL,NR,QL, SYRAL,QL	TEZSPIRE (tezepelumab-ekko) ^{AL} PEN	 All agents require prior authorization AND an FDA-approved diagnosis for approval Non-preferred agents require a trial of a preferred agent within this drug class with the same indication For asthma indications: All agents must be prescribed by or in consultation with an allergist, immunologist, or pulmonologist Agents listed may have other FDA approved indications, and will be subject to prior authorization Drug Specific Criteria: Dupixent: (For other indications, see Immunomodulators, Atopic Dermatitis therapeutic class) For Eosinophilic Asthma or Corticosteroid Dependent Asthma: Patients must be ages 6 and older. Documentation of moderate to severe asthma with either eosinophils >/= 150 + 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

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

IMMUNOMODULATORS, ATOPIC DERMATITIS AL

ADBRY (tralokinumab-ldrm) AL,CL,QL SUB-Q OPZELURA (ruxolitinib phosphate) CREAM AL,CL,QL pimecrolimus (generic for Elidel) PROTOPIC (tacrolimus)	Immunomodulators Self-Injectable PA Form (For Adbry and Dupixent only)
EUCRISA (crisaborole) ^{CL,QL} tacrolimus (generic for Protopic) ZORYVE (roflumilast) ^{AL,NR} FOAM	 Non-preferred agents require: Trial of a topical steroid AND Trial of one preferred product within this drug class Drug-specific criteria: ADBRY: May be approved after a trial or failure of a topical corticosteroid AND a topical calcineurin inhibitor Dupixent: Atopic Dermatitis: May be approved after a maximum of a 90-day trial or failure of a topical corticosteroid AND a topical calcineurin inhibitor Eosinophilic Esophagitis: Trial, failure, or technique difficulty to a swallowed topical corticosteroid or treatment failure of a proton pump inhibitor. Prescribed by, or in consultation with an allergist, gastroenterologist, or immunologist. Documentation that the Patient has a confirmed diagnosis of eosinophilic esophagitis with ≥ 15 eosinophils/high-power field. Nasal Polyps: May be approved with documentation of treatment failure or contraindication within the previous year to an intranasal corticosteroid OR systemic corticosteroid therapy OR prior nasal surgery. Prescribed by, or in consultation with an allergist, pulmonologist, or otolaryngologist [ENT]. Prurigo Nodularis: Patient must have a diagnosis of Prurigo Nodularis with provider attestation of > 20 nodular lesions. Trial and failure of a topical corticosteroid. Prescribed by, or in consultation with an allergist, dermatologist, or immunologist. Eucrisa: 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 Opzelura: May be approved for a diagnosis of Atopic Dermatitis and after a trial/failure of a topical steroid and trial of a

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

IMMUNOMODULATORS, TOPICAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
imiquimod (generic for Aldara)	HYFTOR (sirolimus) ^{AL} GEL imiquimod (generic Zyclara) podofilox (generic Condylox) GEL ^{NR} , SOLN VEREGEN (sinecatechins) ZYCLARA (imiquimod)	Non-preferred agents require clinical reason why preferred agent within this drug class cannot be used

IMMUNOSUPPRESSIVES, ORAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
azathioprine (generic Imuran) azathioprine (generic Azasan) ^{NR} cyclosporine, modified (generic Neoral) CAPS everolimus (generic for Zortress) ^{AL} mycophenolate (generic Cellcept) CAPS, TAB RAPAMUNE (sirolimus) SOLN RAPAMUNE (sirolimus) TAB sirolimus (generic Rapamune) SOLN, TAB tacrolimus	ASTAGRAF XL (tacrolimus) AZASAN (azathioprine) cyclosporine CAP, SOFTGEL cyclosporine, modified (generic Neoral) SOLN ENVARSUS XR (tacrolimus) GENGRAF (cyclosporine, modified) CAP, SOLN mycophenolate (generic Cellcept) SUSP mycophenolic acid MYFORTIC (mycophenolate sodium) MYHIBBIN (mycophenolate) PROGRAF (tacrolimus) CAPS, PACKET REZUROCK (belumosudil) SANDIMMUNE (cyclosporine) CAPS, SOLN TAVNEOS (avacopan) CAPS ZORTRESS (everolimus) AL	 Patients established on existing therapy will be allowed to continue Drug Specific Criteria Tavneos (avacopan) No trial of a preferred agent required with appropriate FDA

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

INTRANASAL RHINITIS DRUGS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
ANTICHOLINERGICS		Non-preferred agents will be
ipratropium (generic for Atrovent)		approved for patients who have failed a 30-day trial of ONE preferred
ANTIHIS	TAMINES	agent within this drug class
azelastine 0.1% (generic for Astelin)	azelastine 0.15% (generic for Astepro) azelastine/fluticasone (generic for Dymista) olopatadine (generic for Patanase) RYALTRIS (olopatadine/mometasone) ^{AL}	 Drug-specific criteria: mometasone: Prior authorization NOT required for children ≤ 12 years budesonide: Approved for use in Pregnancy (Pregnancy Category B) Xhance: Indicated for treatment of
CORTICO	STEROIDS	
fluticasone Rx (generic Flonase)	BECONASE AQ (beclomethasone) budesonide Rx (generic for Rhinocort) flunisolide (generic for Nasalide) fluticasone OTC (generic Flonase OTC) mometasone (generic for Nasonex) RX, OTC ^{NR} OMNARIS (ciclesonide) QNASL 40 & 80 (beclomethasone) XHANCE (fluticasone) ZETONNA (ciclesonide)	

LEUKOTRIENE MODIFIERS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
montelukast (generic for Singulair) TABQL/CHEWABLEAL	montelukast GRANULES (generic Singulair) ^{CL, AL} zafirlukast (generic Accolate) zileuton ER (generic Zyflo CR) ZYFLO (zileuton)	 Non-preferred agents will be approved for patients who have failed a 30-day trial of THE preferred agent within this drug class Drug-specific criteria: montelukast granules: PA not required for age < 2 years

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

LINCOSAMIDES / OXAZOLIDINONES / STREPTOGRAMINS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
clindamycin CAPS clindamycin palmitate SOLN linezolid TAB	CLEOCIN (clindamycin) CAPS CLEOCIN PALMITATE (clindamycin) linezolid SUSP SIVEXTRO (tedizolid phosphate) ZYVOX (linezolid) SUSP, TAB	 Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class

LIPOTROPICS, OTHER

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
BILE ACID SE	QUESTRANTS	Non-preferred agents will be
cholestyramine (generic Questran) colestipol TAB (generic Colestid)	colesevelam (generic Welchol) TAB, PACKET colestipol GRANULES (generic Colestid) QUESTRAN LIGHT (cholestyramine)	approved for patients who have failed a trial of ONE preferred agent within this drug class Drug-specific criteria: Colesevelam: Trial not required for diabetes control and monotherapy with metformin,
TREATMENT OF HOMOZYGOUS FA	MILIAL HYPERCHOLESTEROLEMIA	sulfonylurea, or insulin has been
	JUXTAPID (lomitapide) ^{CL}	inadequate
	KYNAMRO (mipomersen) ^{CL}	Juxtapid®/ Kynamro®:
FIRRIC ACID		 Approved for diagnosis of homozygous familial
fenofibrate (generic Tricor)	DERIVATIVES fenofibric acid (generic Fibricor/Trilipix)	hypercholesterolemia (HoFH)
fenofibrate (generic Lofibra)	fenofibrate (generic Antara/Fenoglide/	OR
gemfibrozil (generic Lopid)	Lipofen/Triglide)	 Treatment failure/maximized dosing/contraindication to ALL
gennibrozii (generic Lopid)	Lipototii Tiigiido)	the following: statins,
NIA	CIN	ezetimibe, niacin, fibric acid
niacin ER (generic Niaspan)	NIACOR (niacin IR)	derivatives, omega-3 agents, bile acid sequestrants
		 Require faxed copy of REMS
		PA form
OMEGA-3 F	ATTY ACIDS	
omega-3 fatty acids (generic Lovaza)	icosapent (generic Vascepa) ^{CL}	
VASCEPA (icosapent)	omega-3 OTC	
CHOLESTEROL ABSORPTION INHIBITORS		
ezetimibe (generic Zetia)	NEXLETOL (bempedoic acid)	
	NEXLIZET (bempedoic acid/	
	ezetimibe) ^{QL}	

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

LIPOTROPICS, OTHER (Continued)

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
PROPROTEIN CONVERTASE SUBTILISIN/KEXIN TYPE 9 (PCSK9) INHIBITORS		Praluent®: Approved for diagnoses of:
PRALUENT (alorocumab) ^{CL}	REPATHA (evolocumab) ^{CL}	 atherosclerotic cardiovascula disease (ASCVD) heterozygous familial hypercholesterolemia (HeFH) Homozygous familial hypercholesterolemia (HoFH) as an adjunct to other LDL-C lowering therapies AND Trial and failure or intolerance to a statin for 8 continuous weeks Failure to reach target LDL-C levels: ASCVD - < 70 mg/dL, HeFH - < 100 mg/dL Repatha®: May be approved for: adult diagnoses of atherosclerotic cardiovascula disease (ASCVD) heterozygous familial hypercholesterolemia (HeFH) in adults and pediatric patient aged 10 years and older homozygous familial hypercholesterolemia (HoFH) in adults and pediatric patient aged 10 years and older Maximized high-intensity stat WITH ezetimibe for 3+ continuous months Failure to reach target LDL-C levels: ASCVD - < 70 mg/dL, HeFH - < 100 mg/dL Concurrent use of maximally-tolerated statin must continue except for statin-induced rhabdomyolysis or a contraindication to a statin

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

LIPOTROPICS, STATINS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
atorvastatin (generic Lipitor) ^{QL} lovastatin (generic Mevacor) pravastatin (generic Pravachol) rosuvastatin (generic Crestor)	ALTOPREV (lovastatin ER) ^{CL} ATORVALIQ (atorvastatin) ^{QL} SUSP EZALLOR SPRINKLE (rosuvastatin) ^{QL} fluvastatin IR/ER (generic Lescol/	 Non-preferred agents will be approved for patients who have failed a trial of TWO preferred agent within this drug class, within the last 12 months
simvastatin (generic Zocor)	Lescol XL) LIVALO (pitavastatin) ^{AL,QL} pitavastatin (generic Livalo) ^{AL,NR,QL} ZYPITAMAG (pitavastatin)	 Drug-specific criteria: Altoprev®: One of the TWO trials must be IR lovastatin Combination products: Require clinical reason why individual ingredients cannot be used
STATIN COM	atorvastatin/amlodipine (generic Caduet) simvastatin/ezetimibe (generic Vytorin)	 fluvastatin ER: Requires trial of TWO preferred agents AND trial of IR fluvastatin OR clinical reason IR cannot be used simvastatin/ezetimibe: Approved for 3-month continuous trial of ONE standard dose statin

MACROLIDES AND KETOLIDES, ORAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
MACRO	MACROLIDES	
azithromycin (generic Zithromax) clarithromycin TAB , SUSP (generic Biaxin) E.E.S. SUSP (erythromycin ethylsuccinate)	clarithromycin ER (generic Biaxin XL) E.E.S. TAB (erythromycin ethylsuccinate) ERY-TAB (erythromycin) erythromycin ethylsuccinate SUSP ERYPED SUSP (erythromycin) ERYTHROCIN (erythromycin) erythromycin base TAB , CAPS	clinical reason why preferred products within this drug class cannot be used AND ≥ 3-day trial on a preferred product

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

METHOTREXATE

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
methotrexate PF VIAL, TABLET, VIAL	JYLAMVO (methotrexate) ^{NR} SOLN OTREXUP (methotrexate) SUB-Q RASUVO (methotrexate) SUB-Q REDITREX (methotrexate) SUB-Q TREXALL (methotrexate) TABLET XATMEP (methotrexate) SOLN	Non-preferred agents require a trial of the preferred agent AND will be approved for an FDA-approved indication Drug-specific criteria: • Xatmep™:Indicated for pediatric patients only

MOVEMENT DISORDERS

FPreferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
AUSTEDO (deutetrabenazine) ^{CL} AUSTEDO XR (deutetrabenazine) ^{CL} AUSTEDO XR Titration Pack	INGREZZA (valbenazine) ^{AL,CL} INITIATION PACK, SPRINKLES ^{NR} XENAZINE (tetrabenazine) ^{CL}	All drugs require an FDA approved indication – ICD-10 diagnosis code required.
(deutetrabenazine) ^{CL} INGREZZA (valbenazine) ^{AL,CLQL} CAPS tetrabenazine (generic for		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.
Xenazine) ^{CL}		 Drug-specific criteria: Austedo/Austedo XR/Ingrezza: 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 tetrabenazine: Diagnosis of chorea with Huntington's Disease

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

MULTIPLE SCLEROSIS DRUGS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
AVONEX (interferon beta-1a) ^{QL} BETASERON (interferon beta-1b) ^{QL} COPAXONE 20mg (glatiramer) ^{QL} dimethyl fumarate (generic for Tecfidera) fingolimod (generic Gilenya) ^{QL} KESIMPTA (Ofatumumab) ^{CL,QL} teriflunomide (generic Aubagio) ^{QL}	AUBAGIO (teriflunomide) ^{QL} BAFIERTAM (monomethyl fumarate) ^{QL} dalfampridine (generic Ampyra) ^{QL} EXTAVIA (interferon beta-1b) ^{QL} GILENYA (fingolimod) ^{QL} glatiramer (generic Copaxone) ^{QL} MAVENCLAD (cladribine) MAYZENT (siponimod) ^{QL} PLEGRIDY (peginterferon beta-1a) ^{QL} PONVORY (ponesimod) REBIF (interferon beta-1a) ^{QL} TASCENSO ODT (fingolimod) TAB ^{AL} TECFIDERA (dimethyl fumarate) VUMERITY (diroximel) ^{QL} ZEPOSIA (ozanimod) ^{AL,CL,QL}	 Non-preferred agents will be approved for patients who have failed a trial of TWO preferred agent within this drug class Drug-specific criteria: Ampyra/ dalfampridine: Approved for diagnosis of gait disorder associated with MS AND EDSS score ≤ 7 Plegridy: Approved for diagnosis of relapsing MS Kesimpta: Approved for patients who have failed a trial of a preferred injectable agent within this class Zeposia: Approved for a diagnosis of relapsing forms of multiple sclerosis (MS) with trial of ONE preferred agent OR a diagnosis of moderately to severely active ulcerative colitis and treatment failure of Humira.

NITROFURAN DERIVATIVES

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
itrofurantoin macrocrystals CAPSULE (generic Macrodantin) itrofurantoin monohydrate- macrocrystals CAPS (generic Macrobid)	nitrofurantoin SUSPENSION (genericFuradantin)	 Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

NSAIDs, ORAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
COX-I SE	LECTIVE	Non-preferred agents within COX-
diclofenac sodium (generic Voltaren) ibuprofen OTC, Rx (generic Advil, Motrin) CHEW, DROPS, SUSP, TAB ibuprofen OTC (generic Advil, Motrin) CAPS indomethacin (generic Indocin) CAPS ketorolac (generic Toradol) meloxicam (generic Mobic) TAB nabumetone (generic Relafen) naproxen Rx, OTC (generic Naprosyn) naproxen enteric coated sulindac (generic Clinoril)	diclofenac potassium (generic Cataflam, Zipsor) diclofenac SR (generic Voltaren-XR) diflunisal (generic Dolobid) etodolac & SR (generic Lodine/XL) fenoprofen (generic Nalfon) flurbiprofen (generic Ansaid) ibuprofen/famotidine (generic Duexis) ^{CL} indomethacin ER (generic Indocin) ketoprofen & ER (generic Orudis) meclofenamate (generic Meclomen) mefenamic acid (generic Ponstel) meloxicam (generic Vivlodex) ^{CL, QL} CAP meloxicam (generic Naprelan) naproxen CR (generic Naprelan) naproxen (generic Naprosyn) SUSP naproxen sodium (generic Anaprox) naproxen-esomeprazole (generic Vimovo) oxaprozin (generic Daypro) piroxicam (generic Tolectin) ketorolac (generic Sprix Nasal) QL NASAL	1 SELECTIVE group will be approved for patients who have failed no less than 30-day trial of TWO preferred agents within this drug class Drug-specific criteria: meclofenamate: Approvable without trial of preferred agents for menorrhagia Sprix/ketoralac Nasal: Approved for patients unable to tolerate, swallow OR absorb oral NSAIDs OR contraindication OR trial of TWO preferred oral NSAIDs

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

NSAIDs, ORAL (Continued)

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
COX-I SELECTIVE (continued)		All combination agents require a
	ALL BRAND NAME NSAIDs including: DUEXIS (ibuprofen/famotidine)CL NALFON (fenoprofen) RELAFEN DS (nabumetone)	clinical reason why individual agents can't be used separately
NSAID/GI PROTECTA	ANT COMBINATIONS	-
	diclofenac/misoprostol (generic Arthrotec)	
COX-II SE	LECTIVE	
celecoxib (generic Celebrex)		

NSAIDs, TOPICAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
diclofenac sodium GEL (OTC only) PENNSAID PUMP (diclofenac)	diclofenac PUMP (generic Pennsaid) ^{CL} diclofenac SOLN (generic Pennsaid) FLECTOR PATCH (diclofenac) ^{CL} LICART PATCH (diclofenac) ^{CL} PENNSAID PACKET (diclofenac) ^{CL} VOLTAREN GEL (diclofenac) ^{CL}	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.

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

NOTE: Other oral oncology agents not listed here may also be available. See https://nebraska.fhsc.com/default.asp for coverage information and prior authorization status for products not listed.

ONCOLOGY AGENTS, ORAL, BREAST

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
CDK 4/6 INHIBITOR		Non-preferred agents DO NOT
	IBRANCE (palbociclib) KISQALI (ribociclib) KISQALI FEMARA CO-PACK VERZENIO (abemaciclib)	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 • Patients undergoing treatment at the time of any preferred status
CHEMO	THERAPY	change will be allowed to continue
capecitabine (generic Xeloda) cyclophosphamide	XELODA (capecitabine)	therapy
		Drug-specific critera
HORMONE	BLOCKADE	 anastrozole: May be approved for malignant neoplasm of male breast
anastrozole (generic Arimidex) exemestane (generic Aromasin) letrozole (generic Femara) tamoxifen citrate (generic Nolvadex)	ORSERDU (elacestrant) SOLTAMOX SOLN (tamoxifen) ^{CL} toremifene (generic Fareston) ^{CL}	 (male breast cancer) Fareston/toremifene: Require clinical reason why tamoxifen cannot be used letrozole: Approved for diagnosis of breast cancer with day supply
ОТ	HER	greater than 12 – NOT approved for short term use
	NERLYNX (neratinib) PIQRAY (alpelisib) lapatinib (generic Tykerb) TALZENNA (talazoparib tosylate) QL TUKYSA(tucatinib)QL TRUQAP (capivasertib)NR	 Soltamox: May be approved with documented swallowing difficulty

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

NOTE: Other oral oncology agents not listed here may also be available. See https://nebraska.fhsc.com/default.asp for coverage information and prior authorization status for products not listed.

ONCOLOGY AGENTS, ORAL, HEMATOLOGIC

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
mercaptopurine	PURIXAN (mercaptopurine) ^{AL}	 Non-preferred agents DO NOT require a trial of a preferred agent, but DO require an FDA-approved
	AML	 indication OR documentation submitted supporting off-label use
	DAURISMO (glasdegib maleate) ^{QL} IDHIFA (enasidenib) REZLIDHIA (olutasidenib) ^{QL} RYDAPT (midostaurin) TIBSOVO (ivosidenib) ^{QL} VANFLYTA (quizartinib) XOSPATA (gilteritinib) ^{QL}	from current treatment guidelines Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy Drug-specific critera Hydrea®: Requires clinical reason
	CLL	why generic cannot be used
LEUKERAN (chlorambucil)	COPIKTRA (duvelisib) QL IMBRUVICA (ibrutinib) VENCLEXTA (venetoclax) ZYDELIG (idelalisib)	 Purixan: Prior authorization not required for age ≤12 or for documented swallowing disorder Tabloid: Prior authorization not required for age <19
	CML	 Xpovio: Indicated for relapsed or
hydroxyurea (generic for Hydrea) imatinib (generic for Gleevec) MYLERAN (busulfan)	BOSULIF (bosutinib) TAB GLEEVEC (imatinib) HYDREA (hydroxyurea) ICLUSIG (ponatinib) SCEMBLIX (asciminib) SPRYCEL (dasatinib) TASIGNA (nilotinib) ^{CL}	refractory multiple myeloma. Requires concomitant therapy with dexamethasone
	MPN	-
	JAKAFI (ruxolitinib)	-
M	YELOMA	
melphalan (generic Alkeran) REVLIMID ^{QL} (lenalidomide)	lenalidomide ^{QL} (generic Revlimid) NINLARO (ixazomib) POMALYST (pomalidomide) THALOMID (thalidomide) XPOVIO (selinexor) ^{CL}	
(OTHER	
MATULANE (procarbazine) TABLOID (thioguanine) tretinoin (generic for Vesanoid) ^{AL}	BRUKINSA (zanubrutinib ^{QL} CALQUENCE (acalabrutinib) ^{QL} INREBIC (fedratinib dihydrochloride) ^{QL} INQOVI (decitabine/cedazuridine) OJJAARA (momelotinib) ^{NR} VONJO (pacritinib) ^{QL} ZOLINZA (vorinostat)	

Unless otherwise specified, the listing of a particular brand or generic name includes all dosage forms of that drug.

CL - Prior Authorization / Class Criteria apply

QL - Quantity/Duration Limit

AL- Age Limit

NR - Product was not reviewed - New Drug criteria will apply

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

NOTE: Other oral oncology agents not listed here may also be available. See https://nebraska.fhsc.com/default.asp for coverage information and prior authorization status for products not listed.

ONCOLOGY AGENTS, ORAL, LUNG

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
	ALK ALECENSA (alectinib) ALUNBRIG (brigatinib) LORBRENA (lorlatinib) ZYKADIA (ceritinib) CAPS, TAB	 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 Patients undergoing treatment at the time of any preferred status change will be allowed to continue
ALK / R	OS1 / NTRK	therapy
	AUGTYRO (repotrectinib) ^{NR} ROZLYTREK (entrectinib) ^{QL} CAPS, PELLETS ^{NR} XALKORI (crizotinib) CAPS, PELLETS ^{NR}	
1	EGFR	
erlotinib (generic for Tarceva)	EXKIVITY (mobocertinib) ^{QL} gefitinib (generic Iressa) GILOTRIF (afatinib) IRESSA (gefitinib) TAGRISSO (osimertinib) TARCEVA (erlotinib) VIZIMPRO (dacomitinib) ^{QL}	
C	OTHER	
	GAVRETO (pralsetinib) ^{QL} HYCAMTIN (topotecan) KRAZATI (adagrasib) LUMAKRAS (sotrasib) ^{QL} RETEVMO (selpercatinib) ^{AL} TABRECTA (capmatinib) ^{QL} TEPMETKO (tepotinib) ^{QL}	

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

NOTE: Other oral oncology agents not listed here may also be available. See https://nebraska.fhsc.com/default.asp for coverage information and prior authorization status for products not listed.

ONCOLOGY AGENTS, ORAL, OTHER

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
temozolomide (generic Temodar)	AYVAKIT (avapritinib) ^{AL,QL} BALVERSA (erdafitinib) CAPRELSA (vandetanib) COMETRIQ (cabozantinib) FRUZAQLA (fruquintinib) ^{NR} CAPS HEXALEN (altretamine) IWILFIN (eflornithine) ^{NR} JAYPIRCA (pirtobrutinib) KOSELUGO (selumetinib) ^{AL} LONSURF (trifluridine/tipiracil) LYNPARZA (olaparib) LYTGOBI (futibatinib) OGSIVEO (nirogacestat) ^{NR} TAB PEMAZYRE (pemigatinib) ^{QL} QINLOCK (ripretinib) RUBRACA (rucaparib) STIVARGA (regorafenib) TAZVERIK (tazemetostat) ^{AL} TURALIO (pexidartinib) ^{QL} TRUSELTIQ (infigratinib) CAPS VITRAKVI (larotrectinib) CAPS, SOLN ZEJULA (niraparib) CAPS, TABS	 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 Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy

ONCOLOGY AGENTS, ORAL, PROSTATE

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
abiraterone (generic Zytiga) ^{AL,QL} bicalutamide (generic Casodex) flutamide XTANDI (enzalutamide) ^{AL,QL}	AKEEGA (niraparib/abiraterone) EMCYT (estramustine) ERLEADA (apalutamide) ^{QL} nilutamide (generic Nilandron) NUBEQA (darolutamide) ^{QL} ORGOVYX (relugolix) ^{AL} YONSA (abiraterone acetonide, submicronized) ZYTIGA (abiraterone) ^{AL,QL}	 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 Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy

Unless otherwise specified, the listing of a particular brand or generic name includes all dosage forms of that drug.

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

NOTE: Other oral oncology agents not listed here may also be available. See https://nebraska.fhsc.com/default.asp for coverage information and prior authorization status for products not listed.

ONCOLOGY AGENTS, ORAL, RENAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
SUTENT (sunitinib) VOTRIENT (pazopanib)	AFINITOR DISPERZ (everolimus) ^{CL} CABOMETYX (cabozantinib) everolimus (generic Afinitor) everolimus SUSP (generic Afinitor Disperz) FOTIVDA (tivozanib) INLYTA (axitinib) LENVIMA (lenvatinib) NEXAVAR (sorafenib) PAZOPANIB (generic Votrient) ^{NR} TAB sorafenib (generic Nexavar) sunitinib malate (generic Sutent) TORPENZ (generic everolimus) ^{NR} TAB WELIREG (belzutifan) ^{QL}	 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 Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy

ONCOLOGY AGENTS, ORAL, SKIN

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
ERIVEDGE (vismodegib)	ODOMZO (sonidegib) ^{CL}	 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
MEKINIST (trametinib) TAFINLAR (dabrafenib)	BRAFTOVI (encorafenib) COTELLIC (cobimetinib) MEKINIST (trametinib) SOLN MEKTOVI (binimetinib) OJEMDA (tovorafenib) ^{NR} SUSP ^{AL} , TAB TAFINLAR (dabrafenib) SUSP ZELBORAF (vemurafenib)	 Patients undergoing treatment at the time of any preferred status change will be allowed to continue therapy

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

OPHTHALMICS, ALLERGIC CONJUNCTIVITIS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
ALREX (loteprednol 0.2%) cromolyn (generic Opticrom) ketotifen OTC (generic Zaditor) olopatadine OTC (Pataday once daily) olopatadine OTC (Pataday twice daily)	ALOCRIL (nedocromil) ALOMIDE (lodoxamide) azelastine (generic Optivar) BEPREVE (bepotastine besilate) bepotastine besilate (generic Bepreve) epinastine (generic Elestat) LASTACAFT (alcaftadine) LASTACAFT (alcaftadine) OTC loteprednol ^{NR} 0.2% (generic Alrex) olopatadine DROPS (generic Pataday) olopatadine 0.1% (generic Patanol) PATADAY XS (olopatadine 0.7%) PATADAY OTC (olopatadine 0.2%) ZERVIATE (certirizine) ^{AL}	 Non-preferred agents will be approved for patients who have failed a trial of TWO preferred agents within this drug class

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

OPHTHALMICS, ANTIBIOTICS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
FLUOROQUINOLONES		Non-preferred agents will be
ciprofloxacin SOLN (generic Ciloxan) ofloxacin (generic Ocuflox)	BESIVANCE (besifloxacin) CILOXAN (ciprofloxacin) gatifloxacin 0.5% (generic Zymaxid) levofloxacin MOXEZA (moxifloxacin) moxifloxacin (generic Vigamox) moxifloxacin (generic Moxeza) VIGAMOX (moxifloxacin)	 approved for patients who have failed a one-month trial of TWO preferred agent within this drug class Azasite®: Approval only requires trial of erythromycin Drug-specific criteria: Natacyn®: Approved for documented fungal infection
MACRO	OLIDES	
erythromycin	AZASITE (azithromycin) ^{CL}	
AMINOGLYCOSIDES		
tobramycin (generic Tobrex drops)		
OTHER OPHTH	ALMIC AGENTS	
bacitracin/polymyxin B (generic Polysporin) polymyxin B/trimethoprim (generic Polytrim)	bacitracin NATACYN (natamycin) ^{CL} neomycin/bacitracin/polymyxin B OINT neomycin/polymyxin B/gramicidin NEOSPORIN (neomycin/polymyxin B/gramcidin) sulfacetamide SOLN (generic Bleph-10) sulfacetamide OINT	

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

OPHTHALMICS, ANTIBIOTIC-STEROID COMBINATIONS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
neomycin/polymyxin/dexamethasone (generic Maxitrol) sulfacetamide/prednisolone TOBRADEX SUSP, OINT (tobramycin and dexamethasone) tobramycin/dexamethasone SUSP (generic TobraDex) all other manufacturers only	BLEPHAMIDE (prednisolone and sulfacetamide) BLEPHAMIDE S.O.P. neomycin/polymyxin/HC neomycin/bacitracin/poly/HC PRED-G SUSP, OINT (prednisolone/gentamicin) tobramycin/dexamethasone SUSP (generic TobraDex) Falcon manufacturer TOBRADEX S.T. (tobramycin and dexamethasone) ZYLET (loteprednol, tobramycin)	 Non-preferred agents will be approved for patients who have failed a trial of TWO preferred agents within this drug class

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

OPHTHALMICS, ANTI-INFLAMMATORIES

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
CORTICOSTEROIDS		ALL sub-classes unless listed
fluorometholone 0.1% (generic FML) OINT LOTEMAX SOLN (loteprednol 0.5%) MAXIDEX (dexamethasone) PRED MILD (prednisolone 0.12%)	dexamethasone (generic Maxidex) difluprednate (generic Durezol) DUREZOL (difluprednate) FLAREX (fluorometholone) FML (fluorometholone 0.1% SOLN) FML FORTE (fluorometholone 0.25%) FML S.O.P. (fluorometholone 0.1%) INVELTYS (loteprednol etabonate) LOTEMAX OINT, GEL (loteprednol) loteprednol GEL (generic Lotemax Gel) loteprednol 0.5% SOLN (generic Lotemax SOLN) prednisolone acetate 1% (generic Omnipred, Pred Forte) prednisolone sodium phosphate prednisolone sodium phosphate 1%	below: Non-preferred agents will be approved for patients who have failed a trial of TWO preferred agents within this drug class NSAID class: Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within the same sub-class
NS	AID	
diclofenac (generic Voltaren) ketorolac 0.5% (generic Acular)	ACUVAIL (ketorolac 0.45%) bromfenac 0.09% (generic Bromday) bromfenac (generic Bromsite) ^{NR} bromfenac 0.07% (generic Prolensa) ^{NR} BROMSITE (bromfenac) flurbiprofen (generic Ocufen) ILEVRO (nepafenac 0.3%) ketorolac LS 0.4% (generic Acular LS) NEVANAC (nepafenac) PROLENSA (bromfenac 0.07%)	

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

OPHTHALMICS, ANTI-INFLAMMATORY / IMMUNOMODULATORS

Preferred Agents	Non-Preferred Agents		Prior Authorization/Class Criteria
RESTASIS (cyclosporine) RESTASIS MULTIDOSE (cyclosporine) XIIDRA (lifitegrast)	CEQUA (cyclosporine) QL cyclosporine (generic Restasis) EYSUVIS (loteprednol etabonate) QL MIEBO (perfluorohexyloctane) TYRVAYA (varenicline tartrate) QL VERKAZIA (cyclosporine emulsion) VEVYE (cyclosporine) NR	•	Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

OPHTHALMICS, GLAUCOMA

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
MIO	TICS	 Non-preferred agents will be
pilocarpine	PHOSPHOLINE IODIDE (echothiophate iodide) VUITY (pilocarpine)	approved for patients who have failed a trial of ONE preferred agen within this drug class
SYMPATHO		_Drug-specific criteria:
ALPHAGAN P (brimonidine 0.15%) brimonidine 0.2% (generic for Alphagan) BETA BLO levobunolol (generic for Betagan) timolol (generic for Timoptic)	ALPHAGAN P (brimonidine 0.1%) apraclonidine (generic lopidine) brimonidine P 0.15% (generic Alphagan P 0.15%) brimonidine 0.1% (generic Alphagan P 0.1%)	Rhopressa and Rocklatan: Electronically approved for patients who have a trial of ONE generic agent, within ophthalmics - glaucoma within 60 days
	solution)	_
CARBONIC ANHYD	RASE INHIBITORS	
dorzolamide (generic for Trusopt)	AZOPT (brinzolamide) brinzolamide (generic Azopt)	
PROSTAGLANI	DIN ANALOGS	
latanoprost (generic for Xalatan) TRAVATAN Z (travoprost)	bimatoprost (generic Lumigan) IYUZEH (latanoprost) tafluprost (generic Zioptan) travoprost (generic Travatan Z) VYZULTA (latanoprostene) XALATAN (latanoprost) ZIOPTAN (tafluprost)	
COMBINATION	ON DRUGS	
COMBIGAN (brimonidine/timolol) dorzolamide/timolol (generic Cosopt)	brimonidine/timolol (generic Combigan) COSOPT (dorzolamide/timolol) dorzolamide/timolol PF (generic Cosopt PF) SIMBRINZA (brinzolamide/brimonidine)	

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

OPHTHALMICS, GLAUCOMA (Continued)

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
ОТН	IER	
RHOPRESSA (netarsudil) ^{CL} ROCKLATAN (netarsudil and latanoprost) ^{CL}		 Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class Drug-specific criteria: Rhopressa and Rocklatan: Electronically approved for patients
		who have a trial of ONE generic agent, within ophthalmics - glaucoma within 60 days

OPIOID DEPENDENCE TREATMENTS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
buprenorphine SL buprenorphine/naloxone TAB (SL) SUBOXONE FILM (buprenorphine/naloxone)	buprenorphine/naloxone FILM LUCEMYRA (lofexidine) ^{CL,QL} ZUBSOLV (buprenorphine/naloxone)	Opioid Dependence Treatment PA Form Opioid Dependence Treatment Informed Consent 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. Drug-specific criteria: Lucemyra: Approved for FDA approved indication and dosing per label. Trial of preferred product not required.

OPIOID-REVERSAL TREATMENTS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
naloxone NASAL(Rx), SYR, VIAL naltrexone TAB	KLOXXADO (naloxone) NASAL naloxone (generic Narcan) OTC NASAL NARCAN (naloxone) NASAL NARCAN (naloxone) NASAL OTC OPVEE (nalmefene) ^{AL} NASAL REXTOVY (naloxone) NR NASAL ZIMHI (naloxone) SYR	 Non-preferred agents will be approved with documentation of why preferred products within this drug class are not appropriate for the patient

Unless otherwise specified, the listing of a particular brand or generic name includes all dosage forms of that drug.

CL - Prior Authorization / Class Criteria apply

QL - Quantity/Duration Limit

AL- Age Limit

NR - Product was not reviewed - New Drug criteria will apply

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

OTIC ANTI-INFECTIVES & ANESTHETICS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
acetic acid (generic for Vosol)	acetic acid/hydrocortisone (generic for Vosol HC)	 Non-preferred agents will be approved for patients who have failed a trial of the preferred agent within this drug class

OTIC ANTIBIOTICS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
CIPRO HC (ciprofloxacin/hydrocortisone) CIPRODEX (ciprofloxacin/dexamethasone) ciprofloxacin/dexamethasone (generic CIPRODEX) neomycin/polymyxin/hydrocortisone (generic Cortisporin) ofloxacin (generic Floxin)	ciprofloxacin ciprofloxacin/fluocinolone (generic Otovel) CORTISPORIN TC (colistin/neomycin thonzonium/hydrocortisone OTOVEL (ciprofloxacin/fluocinolone)	 Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

PAH (PULMONARY ARTERIAL HYPERTENSION AGENTS), ORAL AND INHALED

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
REVATIO (sildenafil) ^{QL} TAB sildenafil (generic Revatio) ^{CL} SUSP tadalafil (generic for Adcirca) ^{CL} TRACLEER (bosentan) TAB TYVASO (treprostinil) INHALATION VENTAVIS (iloprost) INHALATION CORR SI TT	ADEMPAS (riociguat) ^{CL} ADCIRCA (tadalafil) ^{CL} cosentan (generic Tracleer) TAB LETAIRIS (ambrisentan) LIQREV (sildenafil) SUSP DPSUMIT (macitentan) DPSYNVI (macitentan and tadalafil) ^{NR} TAB DRENITRAM ER (treprostinil) REVATIO (sildenafil) ^{CL} SUSP sildenafil (generic Revatio) ^{CL} TAB TADLIQ (tadalafil) SUSP TRACLEER (bosentan) TAB FOR SUSPENSION TYVASO DPI (treprostinil) INHALATION POWDER JPTRAVI (selexipag)	 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 Drug-specific criteria: Adcirca/Liqrev/Revatio/sildenafil tablets and suspension/tadalafil: Approved for diagnosis of Pulmonary Arterial Hypertension (PAH) Adempas®:

PANCREATIC ENZYMES

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
CREON ZENPEP (pancrelipase)	PERTZYE (pancrelipase) VIOKACE (pancrelipase)	 Non-preferred agents will be approved for patients who have failed a trial of TWO preferred agents within this drug class

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

PEDIATRIC VITAMIN PREPARATIONS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
ped mvi no. 17) OTC CHEW	DEKAs PLUS ^{AL} DAVIMET W/ FLUORIDE (ped mvi no.247/ fluoride) ^{NR} CHEW OTC	 Non-preferred agents will be approved for patients who have failed a trial of TWO preferred agents within this drug class
(mand mand man OA (income frame)	FLORIVA (ped mvi no.85/fluoride) CHEW	Drug specific criteria: DEKAs Plus: Approved for
CHILDREN'S CHEWABLES OTC (ped mvi no. 25/FA, ped mvi no. 31 /iron/FA, ped mvi no.17/iron sulf)	FLORIVA PLUS (ped mvi no.161/fluoride) OTC DROP	diagnosis of Cystic Fibrosis and does not require a trial of a preferred agent
CHILDREN'S VITAMINS W/ IRON CHEW OTC (mvi with iron)	MULTI-VIT-FLOR (ped mvi no.205/fluoride) CHEW	
FLUORIDE/VITAMINS A,C,AND D DROPS (ped mvi A,C,D3 no.21/	PEDI MVI NO.242/FLUORIDE CHEW OTC	
	POLY-VI-FLOR (ped mvi no.217/fluoride, ped mvi no. 205/fluoride) CHEW	
MVI NO.2 W-FLUORIDE) DROPS	POLY-VI-FLOR (ped mvi no.213 w/fluoride) DROPS	
MULTIVITS W/ IRON & FLUORIDE DROPS (ped mvi no. 45/fluoride/iron)	POLY-VI-FLOR W/ IRON (ped mvi no. 205/fluoride/iron) CHEW	
	POLY-VI-FLOR W/ IRON (ped mvi no. 214/fluoride/iron) DROP	
POLY-VITAMIN (ped mvi no. 212) DROPS OTC	QUFLORA (ped mvi no.84/fluoride, ped	
POLY-VITAMIN W/ IRON (ped mvi no. 207 w/ferrous sulf) DROPS OTC	mvi no. 63/fluoride, ped mvi no. 83/fluoride)	
TRI-VITAMIN W/ FLUORIDE (ped mvi A,C, D3 no. 21/fluoride)	QUFLORA FE (ped mvi 142/iron/fluoride, ped mvi no. 151/iron/fluoride) CHEW	
	QUFLORA (ped mvi no.157/ fluoride) OTC	
	TRI-VI-FLOR (ped mvi A,C,D3 no.38/fluoride) DROPS	

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

PENICILLINS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
amoxicillin CAPS, CHEWABLE TAB, SUSP, TAB ampicillin CAPS dicloxacillin penicillin VK		 Non-preferred agents will be approved for patients who have failed a 3-day trial of ONE preferred agent within this drug class

PHOSPHATE BINDERS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
calcium acetate TAB CALPHRON OTC (calcium acetate) RENVELA (sevelamer carbonate) PWD PACK, TAB	AURYXIA (ferric citrate) calcium acetate CAPS lanthanum (generic FOSRENOL) PHOSLYRA (calcium acetate) RENAGEL (sevelamer HCI) RENVELA (sevelamer carbonate) PWD PACK sevelamer HCI (generic Renagel) sevelamer carbonate (generic Renvela) VELPHORO (sucroferric oxyhydroxide) XPHOZAH (tenapanor) TAB	 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

PLATELET AGGREGATION INHIBITORS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
aspirin BRILINTA (ticagrelor) clopidogrel (generic Plavix) dipyridamole (generic Persantine) prasugrel (generic Effient)	aspirin/dipyridamole (generic Aggrenox) ticlopidine (generic Ticlid) YOSPRALA (aspirin/omeprazole)	 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

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

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

https://druglookup.fhsc.com/druglookupweb/?client=nestate

PRENATAL VITAMINS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
FE C/FA PNV 11-IRON FUM-FOLIC ACID-OM3 PNV 2/IRON B-G SUC-P/FA/OMEGA-3 PNV NO.118/IRON FUM& PS CMP/FA PNV WITH CA,NO.72/IRON/FA PNV WITH CA,NO.74/IRON/FA OTC PNV#16/IRON FUM & PS/FA/OM-3 PNV119/IRON FUMARATE/FA/DSS PRENATAL MULTI OTC PRENATAL VIT/FE FUMARATE/FA OTC SELECT-OB + DHA STUART ONE OTC TENDERA-OB OTC TRICARE TRINATAL RX 1 VITAFOL CHEW TAB VITAFOL ULTRA VITAFOL-OB VITAFOL-ONE	CITRANATAL B-CALM COMPLETENATE CHEW TAB DERMACINRX PRENATRIX OTC DERMACINRX PRETRATE TAB ENBRACE HR MARNATAL-F MULTI-MAC OTC NATAL PNV (pnv no.164/iron/folate no.6) NESTABS NESTABS ABC NESTABS ONE OB COMPLETE ONE OB COMPLETE PETITE OB COMPLETE PREMIER OB COMPLETE WITH DHA OTC PNV COMBO#47/IRON/FA #1/DHA PNV W-CA NO.40/IRON FUM/FA CMB NO.1 PNV WITH CA NO.68/IRON/FA NO.1/DHA PRENATAL + DHA OTC PRENATE AM PRENATE CHEW TAB PRENATE ENHANCE PRENATE ESSENTIAL PRENATE ESSENTIAL PRENATE FIXIE PRENATE RESTORE PRENATE STAR PRIMACARE SELECT-OB CHEW TAB TRISTART DHA VITAFOL NANO WESTGEL DHA	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

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

PROTON PUMP INHIBITORS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
Nexium) RX ^{QL} omeprazole (generic Prilosec) RX pantoprazole (generic Protonix) ^{QL} PROTONIX SUSP (pantoprazole) rabeprazole (generic Aciphex)	dexlansoprazole (generic Dexilant) esomeprazole magnesium (generic Nexium) OTCQL esomeprazole strontium KONVOMEP (omeprazole/sodium bicarb) SUSP lansoprazole (generic Prevacid)QL NEXIUM SUSP (esomeprazole) omeprazole/sodium bicarbonate (generic Zegerid RX) pantoprazole GRANULES QL	 Non-preferred agents will be approved for patients who have failed an 8-week trial of THREE preferred agents. Pediatric Patients: Patients ≤ 4 years of age – No PA required for Prevacid 30mg/ lansoprazole 30mg capsules (used to compound suspensions). Drug-specific criteria: Prilosec®OTC/Omeprazole OTC: EXCLUDED from coverage Acceptable as trial instead of Omeprazole 20mg Prevacid (lansoprazole) Solutab: may be approved after trial of compounded suspension.

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

SEDATIVE HYPNOTICS

temazepam 15 mg, 30 mg (generic for Restoril) Restoril) stemazepam 15 mg, 30 mg (generic for Restoril) Restoril) stemazepam (generic for Restoril) 7.5 mg, 22.5 mg triazolam (generic for Halcion) stemazepam (generic for Restoril) 7.5 mg, 22.5 mg triazolam (generic for Halcion) stemazepam 7.5/22.5 mg: Requires clinical reason why 15 mg/30 mg cannot be used Others Criteria Non-preferred agents require a trial of TWO preferred agents in the OTHERS sub-category	Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
DAYVIGO (lemborexant) AL.QL doxepin (generic for Silenor) EDLUAR (zolpidem sublingual) eszopiclone (generic for Lunesta) HETLIOZ (tasimelteon) ^{CL} HETLIOZ LQ (tasimelteon) SUSP AL.QL QUVIVIQ (daridorexant) ^{QL} ramelteon (generic for Rozerem) tasimelteon (generic for Hetlioz) CL zolpidem GL CAP zolpidem SL (generic for Intermezzo) ZOLDIDEM SL (generic for Intermezzo) Silenor/doxepin Tablet: Must meet ONE of the following: Contraindication to all of the preferred oral sedative hypnotics agents in the OTHERS sub-category Medical necessity for doxepin dose < 10 mg Age greater than 65 years old or hepatic impairment (3 mg dose will be approved if this criterion is met) zolpidem SL (generic for Ambien CR) zolpidem SL (generic for Intermezzo) zolpidem SL: Requires clinical reason why half of zolpidem tablet cannot be used or documented	temazepam 15 mg, 30 mg (generic for Restoril) OTH zaleplon (generic for Sonata)	estazolam (generic for ProSom) temazepam (generic for Restoril) 7.5 mg, 22.5 mg triazolam (generic for Halcion) ERS BELSOMRA (suvorexant) AL,QL DAYVIGO (lemborexant) AL,QL doxepin (generic for Silenor) EDLUAR (zolpidem sublingual) eszopiclone (generic for Lunesta) HETLIOZ (tasimelteon) HETLIOZ LQ (tasimelteon) SUSP AL,QL QUVIVIQ (daridorexant) QL ramelteon (generic for Rozerem) tasimelteon (generic for Hetlioz) CL zolpidem QL CAP zolpidem ER (generic for Ambien CR)	 Benzodiazepines Criteria Non-preferred agents require a trial of the preferred benzodiazepine agent temazepam 7.5/22.5 mg: Requires clinical reason why 15 mg/30 mg cannot be used Others Criteria Non-preferred agents require a trial of TWO preferred agents in the OTHERS sub-category Silenor/doxepin Tablet: Must meet ONE of the following: Contraindication to all of the preferred oral sedative hypnotics agents in the OTHERS sub-category Medical necessity for doxepin dose < 10 mg Age greater than 65 years old or hepatic impairment (3 mg dose will be approved if this criterion is met) zolpidem/zolpidem ER: Maximum daily dose for females: zolpidem 5 mg; zolpidem ER 6.25 mg zolpidem SL: Requires clinical reason why half of zolpidem tablet

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

SICKLE CELL ANEMIA TREATMENT AL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
DROXIA (hydroxyurea) ENDARI (L-glutamine) ^{CL}	OXBRYTA (voxelotor) ^{CL} SIKLOS (hydroxyurea)	 Endari: Patient must have documented two or more hospital admissions per year due to sickle cell crisis despite maximum hydroxyurea dosage. Oxbryta: Not inidcated 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 tranfusion therapy Siklos: May be approved for use in patients ages 2 to 17 years old without a trial of Droxia

SINUS NODE INHIBITORS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
	CORLANOR SOLN , TAB (ivabradine)	 Diagnosis of Chronic Heart Failure with left ventricular ejection fraction less than or equal to 35%, AND Sinus rhythm with resting heart rate greater than or equal to 70 beats per minute, AND On maximally tolerated doses of beta-blockers OR have a contraindication to beta-blocker use

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

SKELETAL MUSCLE RELAXANTS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
baclofen (generic Lioresal) chlorzoxazone (generic Parafon Forte) cyclobenzaprine (generic Flexeril) ^{QL} methocarbamol (generic Robaxin) tizanidine TAB (generic Zanaflex)	baclofen (generic Fleqsuvy) ^{QL} SUSP baclofen (generic Ozobax) ^{QL} SOLN baclofen (generic Ozobax DS) SUSP carisoprodol (generic Soma) ^{CL,QL} carisoprodol compound cyclobenzaprine ER (generic Amrix) ^{CL} dantrolene (generic Dantrium) FEXMID (cyclobenzaprine ER) FLEQSUVY (baclofen) ^{QL} SUSP LORZONE (chlorzoxazone) ^{CL} LYVISPAH (baclofen) ^{QL} GRANULES metaxalone (generic Skelaxin) NORGESIC FORTE (orphenadrine/ASA/caffeine) orphenadrine ER PARAFON FORTE (chlorzoxazone) tizanidine CAPS ZANAFLEX (tizanidine) CAPS, TAB	 Non-preferred agents will be approved for patients who have failed a 1-week trial of TWO preferred agents within this drug class Drug-specific criteria: cyclobenzaprine ER: Requires clinical reason why IR cannot be used Approved only for acute muscle spasms NOT approved for chronic use carisoprodol: Approved for Acute, musculoskeletal pain - NOT for chronic pain Use is limited to no more than 30 days Additional authorizations will not be granted for at least 6 months following the last dayy of previous course of therapy Dantrolene: Trial NOT required for treatment of spasticity from spinal cord injury Lorzone®: Requires clinical reason why chlorzoxazone cannot be used Soma® 250 mg: Requires clinical reason why 350 mg generic strength cannot be used Zanaflex® Capsules: Requires clinical reason generic cannot be used

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

STEROIDS, TOPICAL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
LOW P	OTENCY	Low Potency Non-preferred agents will be approved for potients who
DERMA-SMOOTHE FS (fluocinolone) hydrocortisone OTC & RX CREAM, LOTION, OINT (Rx only) hydrocortisone/aloe OINT	alclometasone dipropionate (generic for Aclovate) DESONATE (desonide) GEL desonide LOTION (generic for Desowen) desonide CREAM, OINT (generic Desowen, Tridesilon) fluocinolone 0.01% OIL (generic DERMA-SMOOTHE-FS) hydrocortisone/aloe CREAM hydrocortisone OTC OINT HYDROXYM (hydrocortisone) GEL TEXACORT (hydrocortisone)	will be approved for patients who have failed a trial of ONE preferred agent within this drug class
fluticasone propionate CREAM, OINT (generic for Cutivate) mometasone furoate CREAM, OINT, SOLN (generic for Elocon)	betamethasone valerate (generic for Luxiq) clocortolone (generic for Cloderm) fluocinolone acetonide (generic for Synalar) flurandrenolide (generic for Cordran) fluticasone propionate LOTION (generic for Cutivate) hydrocortisone butyrate (generic for Locoid) hydrocortisone butyrate/emoll (generic for Locoid Lipocream) hydrocortisone valerate (generic for Westcort) PANDEL (hydrocortisone probutate 0.1%) prednicarbate (generic for Dermatop)	Medium Potency Non-preferred agents will be approved for patients who have failed a trial of TWO preferred agents within this drug class

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

STEROIDS, TOPICAL (Continued)

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
HIGH P	OTENCY	 High Potency Non-preferred
triamcinolone acetonide OINTMENT, CREAM triamcinolone LOTION	amcinonide CREAM, LOTION, OINTMENT betamethasone dipropionate betamethasone / propylene glycol betamethasone valerate desoximetasone diflorasone diacetate fluocinonide SOLN fluocinonide CREAM, GEL, OINT fluocinonide emollient halcinonide CREAM (generic for Halog) HALOG (halcinonide) CREAM, OINT, SOLN KENALOG AEROSOL (triamcinolone) SERNIVO (betamethasone dipropionate) triamcinolone SPRAY (generic for Kenalog spray) VANOS (fluocinonide)	agents will be approved for patients who have failed a trial of TWO preferred agents within this drug class
VERY HIG	H POTENCY	 Very High Potency Non-preferre
clobetasol emollient (generic Temovate-E) clobetasol propionate CREAM, OINT, SOLN halobetasol propionate (generic Ultravate)	APEXICON-E (diflorasone) BRYHALI (halobetasol prop) LOTION clobetasol SHAMPOO, LOTION clobetasol propionate GEL, FOAM, SPRAY halobetasol propionate FOAM (generic for Lexette) AL,QL IMPEKLO (clobetasol) LOTIONAL LEXETTE(halobetasol propionate) AL,QL OLUX-E /OLUX/OLUX-E CP (clobetasol)	agents will be approved for patients who have failed a trial o TWO preferred agents within this drug class

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

STIMULANTS AND RELATED AGENTS AL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
CNS STIMULANTS		 Non-preferred agents will be
Ampheta	mine type	approved for patients who have failed a trial of ONE preferred
ADDERALL XR (amphetamine salt combo) amphetamine salt combination IR DYANAVEL XR (amphetamine) ^{QL} VYVANSE (lisdexamfetamine) ^{QL} CAPS, CHEWABLE	ADZENYS XR (amphetamine) amphetamine ER (generic Adzenys ER) SUSP amphetamine salt combination ER (generic for Adderall XR) amphetamine salt combination ER (generic Mydayis) ^{AL, NR} CAP amphetamine sulfate (generic for Evekeo) dextroamphetamine (generic for Dexedrine) dextroamphetamine ER (generic for Procentra) dextroamphetamine ER (generic for Dexedrine ER) EVEKEO ODT (amphetamine sulfate) lisdexamfetamine (generic Vyvanse Chew) ^{AL,QL} CHEW lisdexamfetamine (generic Vyvanse) ^{AL,QL} CAP methamphetamine (generic for Desoxyn) MYDAYIS (amphetamine salt combo) ^{QL} XELSTRYM (detroamphetamine) AL,QL PATCH ZENZEDI (dextroamphetamine)	agent within this drug class Drug-specific criteria: Procentra/ dextroamphetamine soln: May be approved with documentation of swallowing disorder Zenzedi®: Requires clinical reason generic dextroamphetamine IR cannot be used

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

STIMULANTS AND RELATED ADHD DRUGS (Continued)^{AL}

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
18 mg, 27 mg, 36 mg, 54 mg DAYTRANA PATCH (methylphenidate) ^{QL} dexmethylphenidate (generic for Focalin IR) dexmethylphenidate (generic Focalin XR) METHYLIN SOLN (methylphenidate) methylphenidate (generic Ritalin) methylphenidate SOLN (generic Methylin) QUILLICHEW ER CHEWTAB (methylphenidate) QUILLIVANT XR (methylphenidate)SUSP m m m m m m m m m m m m m	ADHANSIA XR (methylphenidate) QL APTENSIO XR (methylphenidate) AZSTARYS (serdexmethylphenidate and dexmethylphenidate)QL OTEMPLA XR-ODT (methylphenidate)QL	 Non-preferred agents will be approved for patients who have failed a trial of TWO preferred agents within this drug class Maximum accumulated dose of 108mg per day for ages < 18 Maximum accumulated dose of 72mg per day for ages > 19 Drug-specific criteria: Daytrana/methylphenidate patch: May be approved in history of substance use disorder by parent, caregiver, or patient. May be approved with documentation of difficulty swallowing QuilliChew ER: May be approved for children ≤ 12 years of age OR with documentation of difficulty swallowing

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

STIMULANTS AND RELATED ADHD DRUGS (Continued)^{AL}

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
atomoxetine (generic Strattera) ^{QL} guanfacine ER (generic Intuniv) ^{QL} QELBREE (viloxazine) ^{QL}	clonidine ER (generic Kapvay) ^{QL} STRATTERA (atomoxetine) EPTICS armodafinil (generic Nuvigil) ^{CL} modafanil (generic Provigil) ^{CL}	Note: generic guanfacine IR and clonidine IR are available without prior authorization Non-preferred agents will be approve for patients who have failed a trial of ONE preferred agent within this class Drug-specific criteria: armodafinil and Sunosi: Require trial of modafinil armodafinil and modafinil:
	SUNOSI (solriamfetol) CL,QL WAKIX (pitolisant) CL,QL	approved only for: Sleep Apnea with documentation via sleep study and documentation that C-PAP has been maxed Narcolepsy with documentation of diagnosis via sleep study 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 Sunosi approved only for: Sleep Apnea with documentation via sleep study and documentation that C-PAP has been maxed Narcolepsy with documentation of diagnosis via sleep study Wakix: approved only for excessive daytime sleepiness in adults with narcolepsy with documentation of narcolepsy diagnosis via sleep study

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

TETRACYCLINES

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
doxycycline hyclate IR (generic Vibramycin) doxycycline monohydrate 50MG, 100MG CAPS doxycycline monohydrate SUSP, TAB (generic Vibramycin) minocycline HCI CAPS (generic Dynacin/ Minocin/Myrac) tetracycline	demeclocycline (generic Declomycin) ^{CL} DORYX MPC DR (doxycycline pelletized) doxycycline hyclate DR (generic Doryx) doxycycline monohydrate 40MG, 75MG and 150MG CAP (generic Adoxa/Monodox/ Oracea) minocycline HCl TAB (generic Dynacin/Myrac) minocycline HCl ER (generic Solodyn) NUZYRA (omadacycline) VIBRAMYCIN SUSP (doxycycline) XIMINO (minocycline ER) ^{QL}	 Non-preferred agents will be approved for patients who have failed a sequential 3-day trial of TWO preferred agents within this drug class Drug-specific criteria: Demeclocycline: Approved for diagnosis of SIADH doxycycline suspension: May be approved with documented swallowing difficulty

THROMBOPOIESIS STIMULATING PROTEINS CL

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
PROMACTA (eltrombopag) TAB	ALVAIZ (eltrombopag choline) ^{AL,NR} DOPTELET (avatrombopag) MULPLETA (lusutrombopag) PROMACTA (eltrombopag) SUSP TAVALISSE (fostamatinib)	 All agents will be approved with FDA-approved indication, ICD-10 code is required. Non-preferred agents require a trial of a preferred agent with the same indication or a contraindication. Drug-Specific Criteria Doptelet/Mulpleta: 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

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

THYROID HORMONES

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
levothyroxine TAB (generic Synthroid) liothyronine TAB (generic Cytomel) thyroid, pork TAB UNITHROID (levothyroxine)	ADTHYZA (thyroid, pork) ERMEZA (levothyroxine) SOLN EUTHYROX (levothyroxine) LEVO-T (levothyroxine) levothyroxine CAPS (generic Tirosint) THYQUIDITY (levothyroxine) SOLN TIROSINT CAPS (levothyroxine) TIROSINT-SOL LIQUID (levothyroxine) ^{CL}	 Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class Drug-specific criteria: Tirosint-Sol: May be approved with documented swallowing difficulty

ULCERATIVE COLITIS

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
APRISO (mesalamine) LIALDA (mesalamine) PENTASA (mesalamine) Sulfasalazine IR, DR (generic	balsalazide (generic Colazal) budesonide DR (generic Uceris) DIPENTUM (olsalazine) mesalamine ER (generic Apriso)	 Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class Drug-specific criteria: Asacol HD®/Delzicol DR®:
Azulfidine)	mesalamine ER (generic Pentasa) mesalamine (generic Asacol HD/ Delzicol/Lialda)	Requires clinical reason why preferred mesalamine products cannot be used
mesalamine SUPPOSITORY	CANASA (mesalamine)	_
(generic Canasa)	mesalamine ENEMA (generic	
Sulfite-Free ROWASA (mesalamine)	Rowasa)	
	ROWASA (mesalamine)	
	UCERIS (budesonide)	

July PDL Highlighted in Red indicates May 2024 P&T changes that become effective July 19, 2024

https://nebraska.fhsc.com/Downloads/neclaimlimitations.pdf

UTERINE DISORDER TREATMENT

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
MYFEMBREE (relugolix/ estradiol/ norethindrone acetate) ^{AL, CL,QL} ORIAHNN (elagolix/ estradiol/ norethindrone) ^{AL,CL} ORILISSA (elagolix sodium) ^{QL,CL}		Myfembree, Orilissa, and Oriahnn: Requires an FDA approved indication, must follow FDA dosing guidelines, and have had a trial and failure of an NSAID and oral contraceptive

VASODILATORS, CORONARY

Preferred Agents	Non-Preferred Agents	Prior Authorization/Class Criteria
isosorbide dinitrate TAB isosorbide dinitrate/hydralazine (Bidil) ^{CL} isosorbide mono IR/SR TAB nitroglycerin SUBLINGUAL, TRANSDERMAL nitroglycerin ER TAB	BIDIL (isosorbide dinitrate/hydralazine) ^{CL} GONITRO (nitroglycerin) isosorbide dinitrate TAB (Oceanside Pharm MFR only) NITRO-BID OINT (nitroglycerin) NITRO-DUR (nitroglycerin) nitroglycerin TRANSLINGUAL (generic Nitrolingual) VERQUVO (vericiguat) ^{AL,CL,QL}	 Non-preferred agents will be approved for patients who have failed a trial of ONE preferred agent within this drug class Drug-specific criteria: BiDil/ isosorbide dinitrate-hydralazine: Approved for the treatment of heart failure as an adjunct therapy to standard therapy in self-identified black patients Verquvo: 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%